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# Sex biases in bird and mammal natural history collections

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# **Abstract**

Natural history specimens are widely used across ecology, evolutionary 14 biology, and conservation. Although biological sex may influence all of 15 these areas, it is often overlooked in large-scale studies using museum 16 specimens. If collections are biased towards one sex, studies may not be 17 representative of the species. Here, we investigate sex ratios in over two 18 million bird and mammal specimen records from five large international 19 museums. We found a slight bias towards males in birds (40% females) 20 and mammals (48% females), but this varied among orders. The 21 proportion of female specimens has not significantly changed in 130 years, 22 but has decreased in species with showy male traits like colourful 23 plumage and horns. Body size had little effect. Male bias was strongest in 24 name-bearing types; only 27% of bird and 39% of mammal types were 25 female. These results imply that previous studies may be impacted by 26 undetected male bias, and vigilance is required when using specimen 27 data, collecting new specimens, and designating types. 28

<sup>29</sup> Keywords: sex bias, museum specimens, natural history collections,

<sup>30</sup> birds, mammals

# **Introduction**

<sup>32</sup> Museum specimens are used extensively in studies of taxonomy,

<sup>33</sup> systematics, biogeography, genomics, comparative anatomy,

<sup>34</sup> morphological variability, development, parasitology, stable isotope

<sup>35</sup> ecology, toxicology, morphological evolution and more.<sup>1–3</sup> They are also

of vital importance for understanding how biodiversity responds to 36 anthropogenic impacts.<sup>4</sup> Large studies of species phenotypes using 37 museum specimens, especially in vertebrates, are becoming increasingly 38 common (e.g. evolutionary dynamics in birds,<sup>5</sup> ecomorphological 39 diversification in squamates<sup>6</sup>) and are revealing new insights into the 40 evolution of diversity. These studies require large amounts of data, which 41 can mean the focus is on collecting data from as many species as possible, 42 to the detriment of other sources of variation. Sex is an important factor 43 that influences many aspects of an individual's ecology and life-history 44 (Table 1), but it is often treated as a nuisance variable, overlooked entirely, 45 or data collection focuses on just one sex (e.g. only measuring female rates 46 of phenotypic evolution<sup>7</sup>) to avoid the issue. If natural history collections 47 have unbiased sex ratios (i.e. close to 50% males and females, or reflective 48 of the sex ratio for the species in the wild<sup>8</sup>) then this may not be a 49 problem; if there is a bias in the sex composition of collections, this has 50 implications for studies that assume their samples are representative of 51 the whole population or species (Table 1). No large-scale study of sex 52 ratios in bird and mammal museum collections exists, therefore 53 investigating this is of vital importance as the number of studies using 54 museum specimens continues to rise (e.g. this recent special issue on 55 using museum specimens to study biodiversity in the Anthropocene<sup>4</sup>). 56

Table 1: Prominent uses of natural history specimens and how research outcomes may be influenced by sex biases.

Use	Might sex biases in birds and mammals affect research outcomes?						
Taxonomy	Yes. Sexes often have external differences; if these are used in the taxonomy of						
	the group (e.g. male plumage colouration in birds <sup>9</sup> ) then it may be more difficult						
	to identify individuals to species-level in one sex than another. Consistent over-						
	representation of one sex in samples used in taxonomic studies, and in selection of						
	name-bearing types in particular, may mean that interspecific distinctions between						
	taxa are framed most often for that sex, making the less-represented sex harder to						
	identify and distinguish across species, even if important differences exist, a consid-						
	erable practical problem.						

- SystematicsMaybe. For standard molecular phylogenies, commonly used genes do not differ sub-<br/>stantially among sexes (i.e. not to the extent that they would form different branches).In phylogenomic studies, however, gene trees may vary across a genome if sex chro-<br/>mosomes are included in the sample.<sup>10</sup> Morphological phylogenies are likely to be<br/>most affected, as morphological characters can vary extensively between males and<br/>females. This also has implications for Total Evidence phylogenies that use both mor-<br/>phological and molecular data. mtDNA is often used to investigate species limits (e.g.<br/>gentes limits in cuckoos<sup>11</sup>) and this may differ across sexes.
- Biogeography Maybe. This depends on the scale at which you consider biogeography. In species where all reproduction is sexual, sexes (necessarily) do not differ in terms of largescale historical biogeography, i.e. colonisation of new regions will not succeed if only one sex colonises. However, locally sexes may be spatially segregated (e.g. bat roosting sites<sup>12</sup>), and have different dispersal rates<sup>13</sup> or patterns of habitat use, and differential migration of sexes is common in birds.<sup>14</sup>

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- Genomics
   Yes. Mammals and birds have chromosomal sex determination; in mammals XY male and XX female, in birds ZZ male and ZW female.<sup>15</sup> The X and Z chromosomes are larger and have more genes than W and Y, thus genome size differs among sexes. Many genes are also sex-linked, so genomes will differ between sexes.
- Comparative Yes. Males and females have internal and external anatomical differences, thus sex biases will influence comparative anatomy studies.
- Development **Maybe**. In most vertebrates, early developmental stages are almost identical in males and females, however later development and sexual maturation involve highly divergent growth to result in adult sex differences.<sup>16</sup> If research is focused on early development or juvenile life-history stages then sex biases are unlikely to pose a problem.
- MorphologicalMaybe. Perceived wisdom is that males are more variable than females. However,variabilitymany detailed morphometric studies do not find this (e.g. in mammalian dentition<br/>and skull variation17,18 and references within) in birds or mammals when a large<br/>sample is included. In some groups females are more variable than males (e.g. in<br/>
  Pyriglena antbirds19).

- Parasitology Yes. Males are commonly more susceptible to infection, have lower immune function, and higher parasite loads than females.<sup>20</sup> This is likely due to testosterone inhibiting the immune system.<sup>21</sup> However, this is not true for all species and all kinds of parasites, e.g. breeding female birds have more blood parasites than males.<sup>22</sup> Differences in either direction may cause parasite load and diversity to be misrepresented where collections are sex biased.
- StableisotopeYes. The demands of producing eggs, brooding, pregnancy, and lactation can alterecologystable isotope ratios.23 Many species also have sex segregated diets, e.g. leopards,24and foraging ranges, so stable isotope ratios may vary among sexes even in non-<br/>breeding individuals.
- Toxicology Yes. As above, sexes may differ in foraging ecology, which has consequences for contaminant burden. Furthermore, females may be able to eliminate some contaminants via eggs (e.g. mercury<sup>25</sup>), an option not available to males.

MorphologicalYes. There is extensive sexual dimorphism in many of the traits used in studies ofevolutionmorphological evolution, for example body size,26 thus tempo and mode of evolutionmay vary with sex.

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Here we investigate sex biases in natural history collections of birds and 57 mammals using over two million specimen records from five major 58 international museums. We find a slight bias towards males in both 59 groups. Curators and collections managers do not as a rule discard 60 specimens based on sex, thus any bias is likely related to what is collected 61 in the field. Male bias may be related to active selection for males by 62 collectors in the field, or active avoidance of females with young due to 63 legislation, ethical or conservation considerations. In particular a major 64 suspected source of male bias in collections for some species is deliberate 65 selection for large, "impressive" male specimens, especially where males 66 are larger or more colourful than females, or possess ornaments or 67 weaponry such as horns or antlers. Given the age of most major natural 68 history collections, some male bias may be related to the changes in 69 attitudes towards sex through time, therefore, we expect male bias to 70 decrease towards the present due to changes in collection methods and 71 motivations over the last century. Alternatively, male bias may be 72 accidental, for example due to trapping biases (i.e. trapping method, 73 season of collecting, conspicuous male behaviors or traits), difficulties 74 identifying females to species-level, or in some cases simply because there 75 were more males in a population. In some mammals, higher dispersal, 76 and broader habitat use, in males may result in them being more likely to 77 come into contact with hunters or traps; males may also exhibit lower 78 levels of neophobia increasing their likelihood of being captured, though 79 evidence for this is limited.<sup>27, 28</sup> In some birds, male territorial calls are 8c often used to bring individuals towards a trap, which may also bias 81 collections towards males. 82

To investigate these biases further, we tested whether male bias differed
among orders, with sexual size dimorphism, with the possession of
colourful plumage (birds) or ornamentation or weaponry (mammals) in
males, and through time. Note that these variables mostly test for
deliberate selection for males, because these data are easier to collate, but
our results are likely a combination of deliberate and accidental male
biased collecting.

# <sup>90</sup> Materials and Methods

## <sup>91</sup> Data collection and cleaning

#### 92 Specimen data

We obtained museum bird and mammal collection records from the 93 Global Biodiversity Information Facility (GBIF<sup>29</sup>). Specifically we collated 94 data from the American Museum of Natural History (AMNH; 95 n = 271,407 records<sup>30,31</sup>), Field Museum of Natural History (FMNH; n = 182,984 records<sup>32,33</sup>), Muséum National d'Histoire Naturelle 97 (MNHN; n = 86, 126 records<sup>34, 35</sup>), National Museum of Natural History, 98 Smithsonian Institution (NMNH; n = 496,735 records<sup>36</sup>), and Natural 99 History Museum, London (NHMUK;  $n = 251, 409^{37}$ ). These specimens 100 were obtained between 1751 and 2018, mostly through hunting or 101 trapping, and sexed based on internal or external genitalia or secondary 102 sexual characters, for example plumage colouration or antlers. All raw 103 data can be downloaded from GBIF.<sup>29</sup> 104

Prior to analyses we cleaned the data as follows. (i) Record type. To avoid 105 confusing specimens with archives describing specimens we selected only 106 preserved specimen records; (ii) Age. Juveniles can be harder to sex so we 107 excluded all juveniles, young and foetuses from the dataset; (iii) Year. We 108 removed collection years later than 2018 as these were clearly errors; (iv) 109 Taxonomy. We removed subspecies names and used species binomials 110 because we were interested in species-level sex ratios. To ensure our 111 results were not due to female specimens more frequently being identified 112 to the genus-level only, we also also created a dataset with all specimens 113 with valid generic names. We corrected bird taxonomy using the GBIF 114 backbone taxonomy,<sup>29</sup> and mammal taxonomy using Mammal Species of 115 the World;<sup>38</sup> (v) Type status. We split types into name bearing (Holotype, 116 Syntype, Lectotype, Neotype) and non-name bearing (all others) types. 117 Where the records did not specify the kind of type we define these as 118 ambiguous types; (vi) Sex. We standardized sex to either Female, Male or 119 non-sexed, and removed intersex or hermaphrodite individuals. Note that 120 we recognise that biological sex is a spectrum.<sup>39</sup> We focus here on 121 specimens identified as females and males for simplicity because there 122 were very few recorded intersex specimens in collections databases (only 123 five remained after other data cleaning), but we recognize the importance 12/ of these individuals. We also excluded non-sexed individuals from the 125 analyses. The final dataset contained 2,496,611 specimens (1,395,748 birds 126 and 1,100,863 mammals), 1,647,409 (708,355 birds and 939,054 mammals) 127 of which were sexed (Table A1). 128

#### <sup>129</sup> Sexual dimorphism, plumage colouration and ornamentation data.

We extracted median body masses (g) for males and females from
Lislevand et al.<sup>40</sup> for birds and Jones et al.<sup>41</sup> for mammals, then calculated
sexual size dimorphism by dividing mean male body mass by female
body mass. Note that the sample size for these variables is lower because
sex disaggregated body size data are rare (see Table A2).

To explore how "showiness" might influence sex bias, we included a 135 measure of plumage colouration for passerine birds taken from Dale et 136 al.<sup>42,43</sup> This measure is based on the mean RGB (red green blue) values for 137 400 randomly chosen pixels in six patches (nape, crown, forehead, throat, 138 upper breast, and lower breast) for each sex. We then calculated a 139 plumage dimorphism score by dividing male plumage score by female 140 score for each species. For mammals, we used the Handbook of Mammals 141 of the World to identify mammals where males have "ornamentation". We 142 defined ornamentation as a feature that might increase the likelihood of a 143 collector targeting an individual, specifically horns, antlers, tusks, 144 well-defined manes (i.e. in lions and some baboons), enlarged nasal 145 appendages (e.g. in elephant seals, proboscis monkeys and hammerhead 146 bats), facial colouration (e.g. in mandrills) or large cheek extensions (e.g. 147 orangutans). Where species had ornaments, we recorded whether both 148 sexes or only males routinely possess them. Note that the majority of 149 species with ornaments in our models were Artiodactyla (59 of 67 species). 150

<sup>151</sup> Bird species in several groups show reverse sexual dimorphism, where <sup>152</sup> females are larger or showier than the males. To see if we detect a different <sup>153</sup> pattern in these species we also divided the bird data into species where the female is generally the larger or showier sex (the families Accipitridae,
Falconidae, Scolopacidae, Charadriidae, Jacanidae, Stercorariidae, Sulidae,
Fregatidae, Cuculidae, Trochilidae, Pipridae, and the orders Strigiformes
and Struthioniformes - list taken from<sup>44</sup>), and species where the male is
generally the larger or showier sex (all other species).

<sup>159</sup> The final cleaned data are available on the NHM Data Portal.<sup>45</sup>

#### 160 Analyses

We analysed bird and mammal data separately and performed all
 analyses in R version 3.5.0.<sup>46</sup> Reproducible scripts are available on GitHub
 at https://github.com/nhcooper123/sex-bias-museums.<sup>47</sup>

We first summarised the overall proportion of female, male and unsexed
specimens, and calculated the median proportion of females across
species (using the main species-level dataset) and genera (using the
generic-level dataset). We then summarised differences in the proportion
of female specimens across orders and types.

Unsexed specimens may bias our results toward males if the majority of 169 unsexed specimens are female. We tested for this by fitting linear models 170 to compare the percentage of unsexed specimens and the percentage of 171 female specimens within species; if these unsexed specimens are generally 172 female we expect to see a negative relationship between unsexed 173 specimens and the proportion of females in a species. In addition, we 174 investigated variation in the numbers of unsexed specimens by order, 175 collection continent and collection decade (see Supplementary Materials). 176

Most species were represented by only a few specimens (Figure A1), with 177 large skews towards either males or females at low numbers (Figure A2). 178 To reduce problems this is likely to cause when fitting models, we used 179 only species with 100 or more specimens in our models (see 180 Supplementary Materials for more details), except in our change through 181 time models. In these models our response variable was the proportion of 182 males and females in each species for each year from 1880-2010 (before 187 1880 and after 2010 we did not have any species with sufficient specimens 184 to include). As there were only 55 bird species and 1,216 mammal species 185 with over 100 specimens in a year, change through time models instead 186 used all species with more than 50 specimens in a single year to increase 187 the sample size (see Table A2). 188

We fitted all models using generalised linear models (GLM) with 189 quasibinomial errors, with the proportion of female specimens (success) 190 and the proportion of male specimens (failure) for each species as the 191 response variable (i.e. a binomial response where the number of females 192 and the number of males for each species were jointly modeled). 193 Quasibinomial rather than binomial errors were used due to 194 overdispersion (all models have deviance/residual degrees of freedom far 195 greater than two; see output on GitHub for exact values), and we assessed 196 the significance of model terms using Type II sums of squares. We used 197 standard model checks for GLMs (Q-Q plot, histogram of residuals, 198 residuals vs. linear predictors, response vs. fitted values) to assess model 199 fit. We tested whether the proportion of female and male specimens 200 varied with (i) orders; (ii) collection years (1880-2010); (iii) male body 201 mass (log transformed); (iv) sexual size dimorphism (log transformed); (v) 202

<sup>203</sup> whether males were larger/more showy than females or vice versa
<sup>204</sup> (reverse sexual dimorphism; birds only); (vi) plumage dimorphism (log
<sup>205</sup> transformed; passerine birds only); and (vii) ornamentation (mammals
<sup>206</sup> only). The number of specimens and species in each model are shown in
<sup>207</sup> Table A2.

# **Results and Discussion**

Of the 2,496,328 specimen records (1,395,748 birds and 1,100,580 200 mammals) in our dataset, 20% of bird specimens were female, 31% were 210 male, and 49% were not sexed (Table A1). For mammals, the number of 211 non-sexed individuals was much lower at 15%, likely because it is often 212 easier to identify sex in mammals, with 41% female and 44% male 217 specimens. If we consider only sexed specimens, 40% of bird and 48% of 214 mammal specimens were female (Figure A<sub>3</sub>). In real terms this represents 215 143,905 more male than female specimens in birds and 40,468 more male 216 specimens in mammals. This male bias was not due to unsexed specimens 217 mostly being female (see Supplementary Materials; Figures A4-A8). 218 Results were also qualitatively similar using the generic-level data (Table 210 A1), so we focus only on species-level data below. 220

In the wild, adult sex ratios in many bird species are male skewed, though on average not as skewed as our results (n = 187 species, median 44.8% female;<sup>48</sup> see Supplementary Materials; Figure A9), however, 48% is not a large deviation from the 50% expected in many natural populations of mammals.<sup>8</sup> Well sampled species (i.e. those with at least 100 specimens) with the most extreme sex ratios in our data, i.e. species with fewer than <sup>227</sup> 25% female or 25% male specimens, are shown in Table A3.

#### <sup>228</sup> Variation among orders.

The proportion of female specimens varied across orders for both birds 229  $(F_{24,1721} = 29.81, p < 0.001;$  Figure 1; Figure A10; Table A4) and mammals 230  $(F_{24,1488} = 19.80, p < 0.001;$  Figure 2; Figure A11; Table A4). Most orders 231 had more males than females (Table A4). In birds, of the 25 orders with 232 sufficient data, only tinamous (Tinamiformes; 50.4%) had more females, 233 but these represented just four species in the dataset. The most 234 male-biased orders with more than 25 species were pigeons and doves 235 (Columbiformes; 36.8% female), hummingbirds and swifts (Apodiformes; 236 37.2%; but see Supplementary Materials; Figure A12), and passerines 237 (Passeriformes; 38.4%). Adult sex ratios in Columbiformes and 238 Passeriformes are generally male-skewed,<sup>48–50</sup> but hummingbirds are 239 often female-skewed in the wild.<sup>48,50</sup> This, along with evidence that, on 240 average, Passeriformes are not as male biased as our results (n = 54241 species, median 45.1% female<sup>48</sup>), suggests that greater availability of males 242 alone cannot account for our results. 243

Seven of the 25 mammalian orders with sufficient data had more females, the most extreme being anteaters and sloths (Pilosa; 71.1% female). Most mammal species have a sex ratio of 1:1 at birth,<sup>8</sup> though this can vary in adults. Several species of sloth have higher numbers of females (up to 68.8% females<sup>51</sup>) which may explain why we also found more females in collections, however, giant anteaters (*Myrmecophaga tridactyla*) show variable sex ratios in the field,<sup>52</sup> but strong female bias in collections

(71.3% female). Among the orders represented by more than 25 species in 251 our data, only bats have more females (Chiroptera; 52.2% female; Figure 252 2), despite reportedly balanced adult sex ratios in the wild.<sup>12</sup> This is likely 253 related to widespread sex segregation in bat roosting sites, with many 254 roosts containing individuals of only one sex.<sup>12</sup> In the the past, bats were 255 often trapped by collecting all individuals in a roost site, and female bats 256 may use fewer roost sites than males (e.g. in *Myotis daubentonii*<sup>53</sup>), so skew 257 towards females is not surprising. The most male-biased order of 258 mammals were the even-toed ungulates (Artiodactyla; 39.7% females), but 259 although they exhibit a great deal of variation in adult sex ratio, on 260 average, there are more females than males in wild populations<sup>54</sup> 261 suggesting strong selection for male specimens in this order derived from 262 the deliberate hunting of large males that was common in the 19th and 263 early 20th centuries. 264

#### <sup>265</sup> Changes through time.

- We found male bias increased for birds ( $F_{1,389} = 7.167$ , p = 0.008; Figure
- A13), but decreased for mammals ( $F_{1,3426} = 6.86$ , p = 0.009; Figure A13),
- <sup>268</sup> however the effect sizes were extremely small (birds:

 $_{269}$  slope  $\pm$  SE =  $-0.002 \pm < 0.001$ ; mammals: slope  $\pm$  SE =  $0.001 \pm < 0.001$ ),

- <sup>270</sup> indicating very little change in either class, i.e. there has been no
- <sup>271</sup> improvement in the sex balance of collecting over the last 130 years.

#### <sup>272</sup> Male body mass and sexual size dimorphism.

We found significant effects of male body size on the proportion of female 273 specimens in both birds and mammals (Table A5), however, the direction 274 and strength of the relationship varied among classes and orders (Figures 275 A14-A16; Table A5). Bird species with larger males tended to have more 276 female specimens, whereas the reverse was true for mammals. In 277 mammals this was likely driven by a few orders with large males that 278 have long been favored in collections (e.g. Artiodactyla, Carnivora) and 279 have low median percentages of female specimens (Figure 2; Table A4). 280 Interestingly however, selection for males in these groups did not increase 281 with increasing male body size (Figure A16), instead it appears male 282 carnivores and artiodactyls were preferred over females, regardless of 287 their body size. 284

Rather than selecting large males per se, collectors may favour males when 285 the difference in size between females and males, i.e. sexual size 286 dimorphism, is large. We found that as sexual size dimorphism increased, 287 i.e. as males became increasingly larger than females, there was more bias 288 towards male specimens (Table A5), however, this result was entirely 280 driven by differences among orders (Figures A17-A18); when order was 290 included in the models, sexual size dimorphism did not significantly 291 influence specimen sex ratios over the effects of order (Table A5). As with 292 body mass, this suggested certain orders were more likely to contain more 293 male specimens, regardless of their size with respect to females, 294 suggesting that other characteristics were driving their selection. 295

<sup>296</sup> In birds that show reverse sexual dimorphism we found that the median

<sup>297</sup> percentage of females for species where the male was the larger or <sup>298</sup> showier sex was 40%, the same as for the whole dataset. For species <sup>299</sup> where the female is the larger or showier sex the median percentage of <sup>300</sup> females was 44.6%, closer to the expected 50:50 ratio. There were <sup>301</sup> significantly more females in species where the female is the larger or <sup>302</sup> showier sex ( $F_{1,1744} = 167.9$ , p < 0.001; Figure A19).

#### <sup>303</sup> Plumage and ornaments.

In passerine birds, as males became increasingly more colourful than 304 females, the proportion of female specimens decreased (Figure 3; 305  $F_{1,828} = 58.95, p < 0.001; slope \pm SE = -0.416 \pm 0.054$ ). This relationship 306 was not strong, but fits with anecdotal evidence of collectors preferentially 307 selecting colourful male specimens, especially where plumage differences 308 are large, for example in birds of paradise. Although the numbers are too 309 small to drive the differences seen here, there is also bias towards 310 displaying male specimens in exhibits.55 311

For mammals, species with ornaments (horns, antlers, tusks, manes etc.) 312 had significantly fewer female specimens than those without ornaments 313 (Figure 4;  $F_{1,1510} = 46.98$ , p < 0.001;  $slope \pm SE = -0.328 \pm 0.048$ ). Note, 314 however, that only 67 species across four orders in our analyses had 315 ornamentation (Table A6). Most of the species with ornaments in our 316 models were artiodactyls, and most artiodactyls have horns, antlers or 317 tusks, suggesting that the strong male bias in Artiodactyla (39.7% female; 318 Figure 2) was due to selection for males with these features. Within 319 ornamented species there was no significant difference if both sexes or 320

only males possessed the ornament (Figure 4;  $F_{1,65} = 0.725$ , p = 0.398), suggesting that even where females are phenotypically different, preference is still given to collecting males. This is particularly concerning since most artiodactyl species are female skewed in the wild.<sup>54</sup> We note, however, that artiodactyls only make up 4% of our dataset, and just over 4% of all mammal species,<sup>38</sup> so while deliberate hunting of large males may be a driver in this order, it is unlikely to drive all male biases we see.

#### <sub>328</sub> Type specimens.

Perhaps our most notable finding focused on name bearing type 329 specimens (holotypes, syntypes, lectotypes, and neotypes). Here the bias 330 towards male specimens was extreme; only 25% of bird and 39% of 331 mammal types were female (Figure A3). Although in some instances, 332 males might be considered the appropriate sex for holotypes because male 333 characters such as plumage or bacula are diagnostic (e.g. in bats<sup>56</sup>), we see 334 no reason to not also designate a female paratype to represent the 335 phenotypic range of a species - with the exception that in rare species, or 336 species with few specimens for another reason, this may not be possible. 337 In mammals paratypes were almost 49% female, but bird paratypes were 338 38% female. Additionally, for newly discovered species, genetic 339 identification of species limits may remove the need for male diagnostic 340 characteristics, and thus male holotypes. Currently sex does not form any 341 part of the International Code for Zoological Nomenclature (ICZN) 342 recommendations for designating types, although some historical 343 instructions for collectors emphasize the importance of multiple types 344 (e.g. the classical description of the type<sup>57</sup>). Adding this to the ICZN is of 345

<sup>346</sup> vital importance moving forwards.

#### <sub>347</sub> Conclusions and recommendations.

Here we tested for deliberate selection of large, showy males, especially in 348 species with high levels of sexual dimorphism in these traits. Our 349 analyses suggest that some male bias in collections is the likely result of 350 historical active selection of males. In mammals, males are favoured in 351 species with larger males, and in species with ornaments. These results, 352 however, are driven by carnivores and artiodactyls, and do not account for 353 male biases in species which are not actively targeted by hunters, for 354 example most rodents. In birds, showier males appear to be favoured 355 within passerines, and species that exhibit reverse sexual size dimorphism 356 show less bias towards males. As passerines make up 58% of our dataset 357 (and around 60% of all bird species), active selection for males may be a 358 much larger problem in birds than in mammals. To reduce these 359 imbalances, collectors in the field should strive to avoid trapping biases 360 and biases in selecting individuals to collect. 361

Much bias towards males is probably non deliberate, and related to the 362 characteristics of individual species and how they are trapped. In small 363 mammals, for example, higher dispersal and broader habitat use in males 364 may result in males being more likely to come into contact with hunters 365 or traps.<sup>13</sup> In passerines, male calls are often used to draw birds towards a 366 mist net, resulting in higher numbers of males being trapped (a simple 367 solution to this is to also use playback of female calls, an active area of 368 current research in ornithology $5^8$ ). Some trapping is also opportunistic, so 369

characteristics that mean one sex is exposed to collectors more often may 370 play a key role. For example, the slight female bias in Tinamiformes 371 (50.4%) in this study may be because they are often collected on their 372 nests. Similar patterns are likely for other conspicuous ground-nesting 373 birds. Differential migration of sexes may also influence the numbers of 374 males and females caught at certain locations throughout the year.<sup>14</sup> More 375 ecological studies on species with strong biases towards males or females 376 are needed to help explain these patterns. On top of these factors, females 377 may be harder to identify than males - they may appear similar to 378 juveniles, or lack diagnostic features such as bacula that make 379 identification simpler. If this is the case, many of our unsexed specimens 380 may be female. Our supplemental analyses (Figures A4-A8) suggest this 381 does not cause the male bias, but until these 687,393 unsexed bird and 382 161,526 unsexed mammal specimens are sexed, we cannot determine it 383 would have no effect. 384

Museum professionals, and those using museum collections, should have 385 an awareness of the biases within their collections (not just in terms of sex 386 but also in terms of age, locality, and other factors), and attempt to acquire 387 material to best resolve those biases, whatever their cause. Natural history 388 collections play a critical role in informing multiple research disciplines 389 answering vital questions for the future of biodiversity<sup>4</sup> and are also key 390 resources for public engagement and interaction with biodiversity.55 391 Therefore it is paramount that we continue developing these resources 392 while using a more comprehensive and better informed approach. Finally, 393 researchers investigating broad-scale variation in species should account 394 for these biases when designing data collection protocols and/or in 395

downstream analyses and declare how they dealt with those biases in
resulting publications. Our analyses place particular pressure on
taxonomists to think more carefully about sex when defining
name-bearing types, and suggest more designation of opposite sex
paratypes would be desirable, particularly in birds.

# **Acknowledgments**

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# **Data accessibility**

- <sup>405</sup> Data are available from the NHM Data Portal<sup>45</sup> and GBIF.<sup>29</sup> R code is
- <sup>406</sup> available from GitHub
- <sup>407</sup> (https://github.com/nhcooper123/sex-bias-museums; Zenodo DOI:
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# **592** Author contributions

- <sup>593</sup> NC performed the analyses and wrote the first draft. All authors
- <sup>594</sup> contributed to study design, interpreted results, revised the manuscript,
- <sup>595</sup> and approved the submitted version.

# 596 Competing interests

<sup>597</sup> The authors declare no competing interests.

# 598 Figures

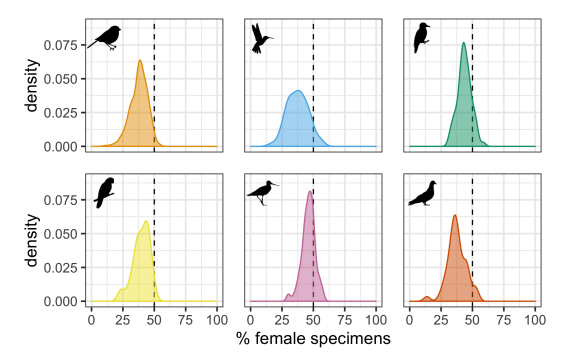


Figure 1: Kernel density plots showing the % female specimens in each species across the six largest orders of birds (from left to right, top to bottom: Passeriformes, Apodiformes, Piciformes, Psittaciformes, Charadriiformes, and Columbiformes). Only species with at least 100 specimens are included. The dashed line represents 50% female specimens. Silhouettes are from PhyloPic.org contributed by Ferran Sayol (parrot, hummingbird, tit), Steven Traver (woodpecker) and Alexandre Vong (shorebird).

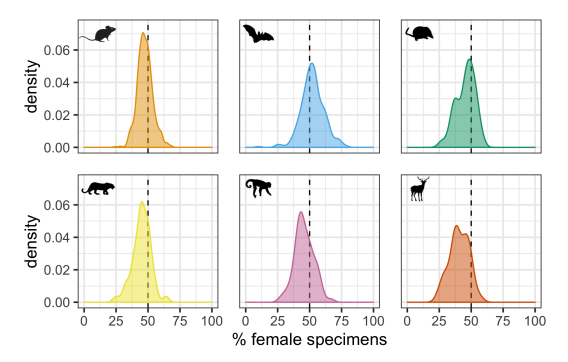


Figure 2: Kernel density plots showing the % female specimens in each species across the six largest orders of mammals (from left to right, top to bottom: Rodentia, Chiroptera, Soricomorpha, Carnivora, Primates, and Artiodactyla). Only species with at least 100 specimens are included. The dashed line represents 50% female specimens. Silhouettes are from PhyloPic.org contributed by Daniel Jaron (mouse), Yan Wong (bat), Becky Barnes (shrew), Lukasiniho (tiger), Sarah Werning (monkey), and Oscar Sanisidro (deer).

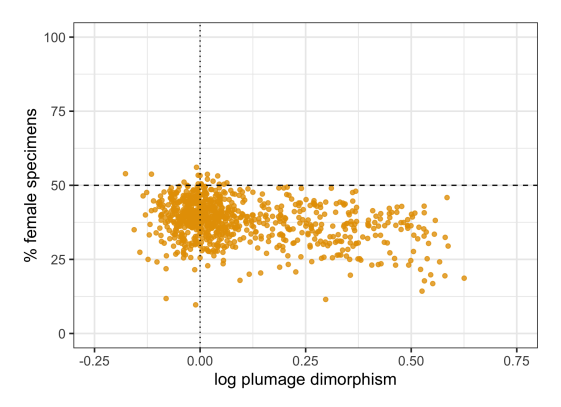


Figure 3: Relationship between the percentage of female specimens for each species and log plumage dimorphism scores in passerine birds. Only species with at least 100 specimens are included. The dashed line represents 50% female specimens; the dotted line is the point at which males and females have the same plumage colouration. Plumage dimorphism scores were calculated by dividing male plumage scores by female plumage scores (see Methods).

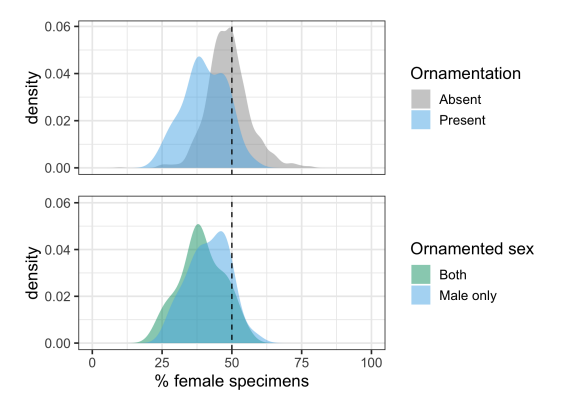


Figure 4: Kernel density plots comparing the % female specimens in each mammal species where ornaments, i.e. horns, tusks, antlers, manes etc., are present or absent (top panel), and when species have ornaments, whether these are found in both sexes or only males (bottom panel). Only species with at least 100 specimens are included. The dashed line represents 50% female specimens.

# <sup>599</sup> Table and figure legends

Table 1: Prominent uses of natural history specimens and how research
 outcomes may be influenced by sex biases.

Figure 1: Kernel density plots showing the % female specimens in each 602 species across the six largest orders of birds (from left to right, top to 603 bottom: Passeriformes, Apodiformes, Piciformes, Psittaciformes, 604 Charadriiformes, and Columbiformes). Only species with at least 100 605 specimens are included. The dashed line represents 50% female 606 specimens. Silhouettes are from PhyloPic.org contributed by Ferran Sayol 607 (parrot, hummingbird, tit), Steven Traver (woodpecker) and Alexandre 608 Vong (shorebird). 600

Figure 2: Kernel density plots showing the % female specimens in each 610 species across the six largest orders of mammals (from left to right, top to 611 bottom: Rodentia, Chiroptera, Soricomorpha, Carnivora, Primates, and 612 Artiodactyla). Only species with at least 100 specimens are included. The 613 dashed line represents 50% female specimens. Silhouettes are from 614 PhyloPic.org contributed by Daniel Jaron (mouse), Yan Wong (bat), Becky 615 Barnes (shrew), Lukasiniho (tiger), Sarah Werning (monkey), and Oscar 616 Sanisidro (deer). 617

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<sub>625</sub> Figure 4: Kernel density plots comparing the % female specimens in each

mammal species where ornaments, i.e. horns, tusks, antlers, manes etc.,

are present or absent (top panel), and when species have ornaments,

<sup>628</sup> whether these are found in both sexes or only males (bottom panel). Only

- <sup>629</sup> species with at least 100 specimens are included. The dashed line
- <sub>630</sub> represents 50% female specimens.