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1 Evolution of large males is associated with female-skewed adult 2 sex ratios in amniotes

3 4 5 **Abstract**

6 Body size often differs between the sexes (leading to sexual size dimorphism, SSD), as a
7 consequence of differential responses by males and females to selection pressures. Adult sex
8 ratio (the proportion of males in the adult population, ASR) should influence SSD because
9 ASR relates to both the number of competitors and available mates, which shape the intensity
10 of mating competition and thereby promotes SSD evolution. However, whether ASR
11 correlates with SSD variation among species has not been yet tested across a broad range of
12 taxa. Using phylogenetic comparative analyses of 462 amniotes (i.e. reptiles, birds and
13 mammals), we fill this knowledge gap by showing that male bias in SSD increases with
14 increasingly female-biased ASRs in both mammals and birds. This relationship is not
15 explained by the higher mortality of the larger sex because SSD is not associated with sex
16 differences in either juvenile or adult mortality. Phylogenetic path analysis indicates that
17 higher mortality in one sex leads to skewed ASR, which in turn may generate selection for
18 SSD biased towards the rare sex. Taken together, our findings provide evidence that skewed
19 ASRs in amniote populations can result in the rarer sex evolving large size to capitalise on
20 enhanced mating opportunities.

21
22 **Keywords:** sexual selection, mating competition, mating opportunity, sex-biased mortality,
23 comparative method

25 INTRODUCTION

26 Sexual size dimorphism (SSD, measured as the size of males relative to females) is
27 widespread in nature and is one of the most conspicuous phenotypic difference between the
28 sexes (Darwin 1871; Andersson 1994; Fairbairn et al. 2007). It is the consequence of different
29 optimal body size for the sexes resulting from opposing selection forces (some of which may
30 influence only one of the sexes) that equilibrate differently in males and females
31 (Blanckenhorn 2005).

32 A large volume of research has focused on how sex-specific behaviour (e.g. mating
33 system, parental care), ecological processes (e.g. abundance and quality of resources), and life
34 history traits (e.g. fecundity in indeterminate growers) can generate size differences between
35 the sexes (Andersson 1994; Blanckenhorn 2005). These studies have concluded that sexual
36 selection is often a major driver of SSD evolution by either intra-sexual competition for
37 access to mates or inter-sexual mate choice, although other evolutionary mechanisms (e.g.
38 fertility selection and competition for resources) may also be important (Jehl and Murray
39 1986; Andersson 1994; Blanckenhorn 2005; Fairbairn et al. 2007; Clutton-Brock 2016).
40 Strong sexual selection for large body size in one sex is particularly likely in species where
41 that sex competes for mates by physical contests or endurance rivalry, as observed in several
42 vertebrate taxa (e.g. reptiles, birds, and mammals; Jehl and Murray 1986; Andersson 1994;
43 Cox *et al.* 2007; Székely *et al.* 2007; Clutton-Brock 2016).

44 Adult sex ratio (ASR), best measured as the proportion of males in the adult
45 population (Ancona et al. 2017) is a key demographic property of populations that influences
46 both the number of competitors for mates and the number of mates available to an individual
47 (Murray 1984; Székely et al. 2014b; Jennions and Fromhage 2017; Schacht et al. 2017). For
48 example, a male-skewed ASR means potentially more competitors and fewer available
49 partners for males than for females. An increasing number of studies show that ASR covaries

50 with several reproductive traits such as mating system, parental sex roles, divorce rate, extra-
51 pair mating and cooperative breeding both in non-human animals and humans (Liker et al.
52 2013, 2014; Schacht et al. 2014; Kappeler 2017; Komdeur et al. 2017; Eberhart-Phillips et al.
53 2018; Grant and Grant 2019). However, whether and how ASR is related to the evolution of
54 SSD is still poorly understood.

55 Theories suggest that ASR can drive the evolution of SSD in at least two ways. First,
56 the intensity of sexual competition may increase with the number of competitors. As Darwin
57 wrote (1871, p. 217): “*That some relation exists between polygamy and development of*
58 *secondary sexual characters, appears nearly certain; and this supports the view that a*
59 *numerical preponderance of males would be eminently favourable to the action of sexual*
60 *selection*”. According to his idea, highly skewed ASRs may intensify selection for
61 competitive traits such as weapons and large body size in the more abundant sex. Thus this
62 ‘mating competition hypothesis’ predicts that the extent of male-bias in SSD should increase
63 with the degree of male skew in the ASR. Later work refined Darwin’s (1871) original idea
64 by suggesting that the operational sex ratio (OSR, the number of sexually active males per
65 receptive female at a given time) rather than the ASR determines the intensity of mating
66 competition in a population (Emlen and Oring 1977). Thus, according to this latter theory
67 ASR would predict SSD if ASR covaries with OSR, for example because OSR is in part
68 determined by ASR (together with sex differences in behaviour like parental care; Kokko et
69 al. 2012). Although the relationship between ASR and OSR is yet to be fully explored, their
70 positive association has been demonstrated both by theoretical models (Kokko and Jennions
71 2008: Fig. 4a; Fromhage and Jennions 2016: Fig. 3c,d) and comparative analyses (Mitani et
72 al. 1996, correlation between ASR and OSR in 18 primates: $r = 0.4$, $P = 0.002$; unpublished
73 result using data from their Table 1). Empirical studies commonly use ASR and OSR

74 interchangeably in testing their relationship with SSD (Poulin 1997) and other proxies of
75 sexual selection (Janicke and Morrow 2018).

76 Second, models of reproductive sex roles predict that ASR should influence the
77 evolution of SSD because individuals of a given sex may allocate less to parental care when
78 the sex ratio is skewed towards the opposite sex than when it is skewed towards their own sex
79 (Queller 1997; McNamara et al. 2000). According to these models, males in female-skewed
80 populations display a higher reproductive success due to increased probability of breeding
81 with multiple partners and therefore may evolve to reduce parental care (Queller 1997:
82 section 3., McNamara et al. 2000: section ‘Sex ratio’). This association between ASR and
83 parental sex roles can drive the evolution of SSD because more elaborate trait expression in
84 males is evolutionarily linked to female-biased care and stronger sexual selection on males
85 (the so called ‘sex-role syndrome’, Janicke et al. 2016: Fig 3.). Thus, this ‘mating
86 opportunity hypothesis’ predicts that the extent of male bias in mating competition, and hence
87 in SSD, should decrease with increasing male skew in the ASR. A demographic analysis of
88 mating systems by Murray (1984) also predicts that female-skewed ASRs should be
89 associated with both polygyny and male-biased SSD, whereas male-skewed ASRs should be
90 associated with polyandry and female-biased SSD.

91 Alternatively, SSD may drive changes in sex ratios through sex differences in
92 mortality resulting from sexual competition. According to this ‘mortality cost hypothesis’, the
93 skewed ASR is a consequence rather than a cause of intense sexual selection, because when
94 males allocate a lot to mating competition they may suffer increased mortality, which in turn
95 leads to female-skewed ASR (Trivers 1972; Clutton-Brock et al. 1985; Liker and Székely
96 2005; Kalmbach and Benito 2007). This hypothesis predicts that in species exhibiting SSD
97 (1) the larger sex should have higher mortality due to the costs of being large, including the

98 direct costs associated with competition (e.g. fights, displays); which leads to (2) decreasing
99 male skew in the ASR with increasing degree of male bias in the SSD.

100 Studies that have investigated the relationships between sex ratios, SSD and sex-
101 specific mortality have so far yielded inconsistent results. While some studies found a
102 positive link between SSD and ASR or OSR (i.e. an increasing male bias in SSD with
103 increasing male skew in the sex ratios; Mitani *et al.* 1996; Poulin 1997), others reported
104 negative associations (Clutton-Brock *et al.* 1977; Wittenberger 1978; Georgiadis 1985; Haro
105 *et al.* 1994; Johansson *et al.* 2005; Lovich *et al.* 2014), or found no consistent relationships
106 (Owen-Smith 1993; Hirst and Kjørboe 2014; Muralidhar and Johnson 2017). Similarly,
107 mortality costs paid by the larger sex in dimorphic species were reported in some studies
108 (Clutton-Brock *et al.* 1985; Promislow 1992; Promislow *et al.* 1992; Moore and Wilson 2002;
109 Benito and González-Solís 2007; Kalmbach and Benito 2007), whereas no consistent
110 relationship between SSD and sex differences in mortality was found by others (Owens and
111 Bennett 1994; Toïgo and Gaillard 2003; Lemaître and Gaillard 2013; Székely *et al.* 2014a;
112 Tidière *et al.* 2015). Many of these studies focused on a narrow range of taxonomic groups
113 and were based on a relatively small number of species (typically fewer than 50) in
114 comparative analyses. Furthermore, none of the studies tested explicitly whether statistical
115 models assuming that ASR drives variation in SSD (as proposed by the mating competition
116 and mating opportunity hypotheses) or alternative models (like the mortality costs hypothesis)
117 fit better to the data.

118 Here we investigate the strength and direction of the relationship between ASR and
119 SSD in populations of wild amniotes, using the largest existing comparative dataset on ASR
120 compiled to date (462 species). First, we investigate whether SSD increases or decreases with
121 ASR across species, as predicted by the mating competition and mating opportunity
122 hypotheses, respectively. We also test whether the relationship is consistent among three

123 major amniote taxa (reptiles, birds, and mammals) because these taxa differ in multiple
124 ecological, behavioural and life history traits. Since the extent and direction of SSD can be
125 influenced by ecological, life history and behavioural factors besides mating competition, we
126 also control for several potential confounding variables in the analyses. Second, we study
127 whether SSD drives ASR variation by generating sex-biased mortality as proposed by the
128 mortality cost hypothesis. We test this latter hypothesis by investigating whether SSD is
129 related to sex differences in juvenile or adult mortality, and by comparing path models
130 representing different structural relationships between SSD, ASR and sex-specific mortality.

131

132 **METHODS**

133 **Data collection**

134 Data were extracted from published sources (see Appendix S1 in Supporting Information).
135 The initial dataset was based on Pipoly *et al.* (2015) that contains ASR and SSD for 344
136 amniote species. We excluded amphibians included in Pipoly *et al.* (2015) because sex-
137 specific mortality data (see below) are very scarce for this taxon, especially in juveniles. The
138 initial dataset was augmented with additional reptile and mammal species, and with
139 information on sex-specific mortality. These additional data were taken from existing
140 comparative datasets (Berger and Gompper 1999 and Bókony *et al.* 2019 for ASR in
141 mammals and reptiles, respectively, and Székely *et al.* 2014a for mortality in birds) or from
142 primary publications. In the latter case we searched the literature through the search engines
143 Web of Science and Google Scholar, using the search terms ‘sex ratio’, ‘sex-specific
144 mortality OR survival’ or ‘male female mortality OR survival’ together with taxonomic
145 names. Data for different variables for the same species were often available only from
146 different populations or studies. The final dataset includes 462 species with both ASR and
147 SSD available (155 reptiles, 185 birds, 122 mammals).

148

149 **Body mass and SSD**

150 Sex-specific body mass (g) was available for all birds and mammals in our dataset. Since
151 body mass data were missing for many reptiles, we also collected body length data (mm) for
152 this taxon in the form of snout-vent length for squamates and crocodylians and plastron or
153 carapace length for turtles. We estimated body mass from body length using published
154 allometric equations (Appendix S2). We used estimated body mass for reptiles instead of
155 body length in the combined analyses of all species because (1) data on mass are more readily
156 available than data on body length in birds and mammals, which provided the majority of
157 species, and (2) body mass is measured in a standardized way in all taxa, whereas the
158 measurement of body length varies because different parts of the body are recorded as a proxy
159 for length in different taxa. If multiple mass or length data were available for a species, we
160 used the mean value. Average adult body mass was calculated as \log_{10} -transformed mean
161 mass of the sexes.

162 We calculated SSD as $\log_{10}(\text{male mass} / \text{female mass})$. **Earlier studies criticised**
163 **measures of SSD that are based on male/female (or female/male) ratios and suggested other**
164 **approaches, for example to analyse male size as response variable in models that also include**
165 **female size as a control variable (see Smith 1999 and Fairbairn 2007 for reviews). In his**
166 **seminal paper, however, Smith (1999, p. 444) convincingly demonstrated that ratios can be**
167 **safely used in the context of SSD analyses because "the risk of spurious correlation is**
168 **negligible to non-existent" due to the statistical properties of male and female size variables**
169 **(i.e. their high correlation and approximately equal coefficients of variation, leading to an**
170 **isometric relationship). We checked the assumption of isometry between male and female**
171 **body mass in our dataset and found that male and female body mass (on a \log_{10} - \log_{10} scale)**
172 **are strongly correlated ($r = \text{XX}$) with a slope very close and not different from 1**

173 (phylogenetic generalized least squares, slope \pm SE: 1.0096 ± 0.0102 , 95% CI: $0.989 \leq \beta \leq$
174 1.029 , $n= 462$ species). Furthermore, Smith (1999, pp. 439-440) demonstrated that the
175 approaches based on the log ratios versus male mass as response variable are statistically
176 equivalent and suggested that the correct method is using log SSD ratio as response and
177 controlling for log size. We thus followed this latter approach. However, because the measure
178 of SSD remains a controversial issue among evolutionary ecologists (see e.g. Table 1 in
179 Tidière et al. 2015 for a review of SSD metrics commonly used), we replicated the main
180 analysis using an alternative method (i.e. male size as response variable while controlling for
181 female size in the model) to check the robustness of our results.

182 To test whether the results are sensitive to conversion of length to mass in reptiles, we
183 replicated the main analyses (1) with SSD calculated from body length ($\log_{10}(\text{male length} /$
184 $\text{female length})$) of reptiles, and (2) with SSD calculated from body mass for a subset (31
185 species) of reptiles that has sex-specific mass data available from Myhrvold et al. (2015).
186 Whatever approach was used to assess the degree of SSD the results were qualitatively
187 unchanged (see Results). In the main text we thus report results based on body mass estimated
188 from body length for reptiles.

189

190 **Sex ratio**

191 We followed Wilson and Hardy (2002) and Ancona et al. (2017) in expressing ASR as the
192 proportion of males in the adult population. We defined the adult population here broadly as
193 adult individuals living in the study area during ASR sampling. Wilson and Hardy (2002)
194 showed that analysing sex ratios as a proportion variable is appropriate when sex ratios are
195 estimated from samples of ≥ 10 individuals and the dataset has ≥ 50 sex ratio estimates. These
196 conditions were more than fully met in our analyses because sample sizes for ASR estimates

197 were always larger than 10 individuals per species (and typically much larger), and our
198 overall dataset included nine times more than the requirement of 50 species.

199 ASR data from Pipoly *et al.* (2015) were augmented with new species and updated
200 with more recent and/or better quality information (e.g. based on a more reliable method or a
201 larger sample size) for some reptiles. ASR estimates were collected by different observers for
202 the different taxa: reptiles by V.B. and I.P. (Pipoly *et al.* 2015; Bókony *et al.* 2019), birds by
203 A.L. (Liker *et al.* 2014), and mammals by Berger and Gompper (1999), Donald (2007) and
204 Anile and Devillard (2018). Details of data selection criteria are given in the original
205 publications (see also Ancona *et al.* 2017). Mean values were calculated for species with
206 multiple ASR data. ASR estimates are repeatable between populations of the same species as
207 measured by the intraclass correlation coefficient (ICC), although the magnitude of
208 repeatability varies among taxa: reptiles with genetic and environmental sex determination:
209 ICC= 0.55 and 0.14, respectively (Bókony *et al.* 2019), birds: ICC= 0.64 (Ancona *et al.* 2017),
210 mammals: ICC= 0.60 (Valentine Federico, J-F.L., J-M.G., A.L., I.P., T.S. unpublished
211 results). ASR estimates are not influenced by the sample size of the ASR studies (Székely *et*
212 *al.* 2014a; Bókony *et al.* 2019).

213

214 **Sex-specific mortality**

215 Annual mortality rates were collected from studies in which mortality (or survival) was
216 estimated for each of both sexes. Juvenile and adult mortality refer to age classes before and
217 after the age of first reproduction, respectively. For reptiles, data were collected by V.B.
218 (Bókony *et al.* 2019). Most adult mortality data on birds are taken from Székely *et al.* (2014a)
219 with the addition of new data for juvenile mortality by A.L. Reptile and bird mortality
220 includes estimates by various methods (capture-recapture, return rates, ...), although we used
221 better quality estimates (e.g. those from capture-recapture analyses) whenever we had a

222 choice (Székely *et al.* 2014a; Bókonyi *et al.* 2019). For mammals, all sex-specific estimates
223 were collected by J-M.G. and J-F.L. (Lemaître *et al.* 2020). Sex differences in juvenile and
224 adult mortality rates were calculated as the magnitude of male-biased mortality (i.e.
225 $\log_{10}(\text{juvenile or adult male mortality} / \text{juvenile or adult female mortality})$), also referred to as
226 ‘mortality bias’. These measures of mortality bias are not related to the overall mortality rate
227 of the species, as estimated by the average mortality rates of the sexes (phylogenetic
228 generalised least squares models, juvenile mortality bias: slope $\pm SE = -0.068 \pm 0.101$, $t =$
229 0.7 , $P = 0.497$, $n = 100$; adult mortality bias: slope $\pm SE = -0.05 \pm 0.08$, $t = 0.7$, $P = 0.513$, n
230 $= 230$).

231

232 **Other predictors**

233 We controlled for the potential effects of ecological variables and life-history traits related to
234 either ASR or SSD (or both) that may confound the assessment of their relationship. First, we
235 collected data on the type of sex determination system because it is associated with both ASR
236 (Pipoly *et al.* 2015) and SSD (Adkins-Regan and Reeve 2014). We divided the species into
237 three categories according to the Tree of Sex database (Ashman *et al.* 2014): male-
238 heterogametic (XY) or female-heterogametic (ZW) genetic sex determination, or temperature-
239 dependent sex determination (TSD). For species that were not included in the Tree of Sex
240 database we assumed the same type of sex determination as reported for the genus (or family,
241 respectively; Bókonyi *et al.* 2019) when the genus (or family) to which it belongs had
242 invariable sex determination system. All birds were assigned to ZW, and all mammals to XY
243 sex determination (Ashman *et al.* 2014).

244 Second, we controlled for the potential effects of environmental variation among
245 species by using two measures. Breeding latitude correlates with life history traits in many
246 organisms (as shown in pioneer work, Dobzhansky 1950) and may also influence the

247 potential for polygamy, hence also sexual selection (Fischer 1960; Isaac 2005;
248 Balasubramaniam and Rotenberry 2016). We used absolute values of the geographic latitude
249 of the ASR studies included in our dataset (i.e. average values for species with multiple ASR
250 estimates) to represent the distance from the Equator. When the authors did not report
251 latitude, we used Google Earth to estimate it as the center of the study sites based on the site
252 descriptions. For 30 birds and 10 mammals, accurate population locations were not reported,
253 hence, we used the latitudinal midpoint of the breeding ranges of these species (birds: V.
254 Remeš, A. Liker, R. Freckleton and T. Székely unpublished data, mammals: PanTHERIA
255 database).

256 In addition to latitude, we investigated environmental harshness as a second
257 environmental variable, which also has been hypothesized to influence SSD (Isaac 2005). We
258 quantified the harshness of the breeding environment using a proxy proposed by Botero *et al.*
259 (2014). This is the PC1 score extracted from Principal Component Analysis (PCA) performed
260 on a set of climatic and ecological variables (e.g. temperature and precipitation, net primary
261 productivity, habitat heterogeneity; see Botero *et al.* 2014 for a detailed description of the
262 variables and the analysis). The PC1 scores have higher values for a higher level of exposure
263 to drier, less productive environments, with colder, less predictable and more variable annual
264 temperatures (see Table 1 in Botero *et al.* 2014). In birds and mammals, we used the data
265 published in Botero *et al.* (2014), whereas for reptiles we calculated PC1 scores by
266 performing a PCA with the same set of variables.

267 Third, we characterized courtship displays in birds because earlier studies showed that
268 birds with aerial displays have less male-biased SSD compared to species with ground
269 displays, probably because selection favors male agility in aerially displaying species
270 constraining male body size (Jehl and Murray 1986; Székely *et al.* 2007). We followed
271 Székely *et al.* (2007) and divided species into two display groups: (1) mating displays that

272 may favor male agility, including species that mainly have aerial displays (both non-acrobatic
273 and acrobatic, categories 4 and 5 in Székely *et al.* 2007), and (2) displays that may not favor
274 male agility, including all other display types, typically performed on ground (categories 1-3
275 in Székely *et al.* 2007). Although SSD can also be influenced by display type and display
276 habitat in reptiles and mammals (e.g. see Agha *et al.* 2018), we were not able to collect
277 reliable data for these taxa, therefore we analyzed the effect of display type only in birds.

278 Fourth, we tested for the potential effect of social mating system, because the scope
279 for mating competition may be more limited in monogamous than in polygamous species
280 (Andersson 1994). Thus, although there is ASR variation among monogamous species that
281 can generate some variation in mating competition and/or opportunity, the relationship
282 between ASR and SSD is expected to be weaker in monogamous than in polygamous species.
283 To test this idea, we characterized social mating system for birds and mammals, because we
284 found reliable information in these taxa for most species (Liker *et al.* 2014; Lukas and
285 Clutton-Brock 2013). Although polygamous mating system differs from promiscuous mating
286 system, we pooled these mating systems because sexual selection is consistently stronger in
287 polygamous than in monogamous species, whereas the relative intensity of sexual selection in
288 polygynous vs. promiscuous species is not easy to assess. We thus categorized species as
289 either socially monogamous or polygamous (most often polygynous) according to the
290 sources, as previously done (see e.g. Clutton-Brock and Isvaran 2007). Although In birds,
291 mating system was originally scored on a five point scale (Liker *et al.* 2014), and here we
292 considered a species monogamous if it had score 0 or 1 (polygamy frequency <1%) for both
293 sexes.

294 Finally, in reptiles, the evolution of viviparity and reduced reproductive frequency are
295 generally correlated with shifts toward female-biased SSD due to fecundity selection for large
296 female size (Pincheira-Donoso and Hunt 2017). To control for its potential effect on SSD, we

297 categorized the reproductive mode of reptiles as either viviparous or oviparous (Uetz et al.
298 2019).

299

300 **Statistical analyses**

301 Phylogenetic generalized least squares (PGLS) models were built to conduct bivariate and
302 multi-predictor analyses. To control for phylogenetic relationships among taxa, we used the
303 composite phylogeny applied in Pipoly *et al.* (2015) with the addition of new species
304 according to the family-level (Sarre et al. 2011) and other recent phylogenies (Squamata:
305 Nicholson *et al.* 2012, Pyron *et al.* 2013, Gamble *et al.* 2014; Testudines: Barley *et al.* 2010,
306 Guillon *et al.* 2012, Spinks *et al.* 2014; Crocodylia: Oaks 2011; mammals: Fritz *et al.* 2009,
307 Meredith *et al.* 2011). Since composite phylogenies do not have true branch lengths, we used
308 three methods to generate branch lengths (Nee's method, Pagel's method, and unit branch
309 lengths, using the PDAP:PDTREE module of Mesquite; Midford *et al.* 2011), and repeated
310 key analyses with these alternative trees. We present results with Nee's branch lengths in the
311 paper, except for the sensitivity analyses (see Results). Freckleton et al. (2002) showed that
312 PGLS is relatively insensitive to branch length assumptions. In each model we used the
313 maximum-likelihood estimate of phylogenetic dependence (Pagel's λ). PGLS models were
314 run using the 'caper' R package (Orme et al. 2013).

315 First, using all species, we applied bivariate PGLS models to test interspecific
316 associations between ASR, SSD and sex differences in juvenile and adult mortality rates.
317 **When SSD was the response variable in the model, we also included mean body mass as a**
318 **second predictor, as recommended by Smith (1999) (hence we termed these models as**
319 **'separate predictor models' instead of bivariate models in the rest of the paper).** Then we built
320 two multi-predictor models. In Multi-predictor model 1, we tested the relationship between
321 ASR and SSD while controlling for potential confounding effects of mean mass, sex

322 determination system, and breeding latitude. In Multi-predictor model 2, we tested the ASR -
323 SSD relationships while controlling for the effects of sex differences in juvenile and adult
324 mortality rates, and mean mass. We built these two separate multi-predictor models because
325 we have much lower sample sizes for sex-specific mortalities than for the other predictors,
326 thus the statistical power would be reduced for variables of Multi-predictor model 1 if all
327 predictors were combined in a single model. We ran the models in two alternative versions in
328 which either SSD or ASR was the dependent variable, respectively, since we had no *a priori*
329 knowledge about the cause-effect direction of these relationships and results may differ
330 between these analyses if the two models have different values for Pagel's λ (see Appendix
331 S3).

332 We investigated whether the ASR – SSD relationship, which is the main focus of our
333 study, differed among taxa by testing the interaction between ASR and the taxonomic class.
334 To explore differences among taxa in the multivariate relationships, we repeated all analyses
335 separately for reptiles, birds and mammals. In taxon-specific Multi-predictor models 1, we
336 included reproductive mode for reptiles and display type for birds as further predictors. In
337 reptiles, we also tested whether the relationship between ASR and SSD is sensitive (1) to the
338 inclusion of species that have environmental sex determination, because ASR shows low
339 repeatability in such reptiles (Bókony et al. 2019), and (2) to the inclusion of species in which
340 the type of sex determination was inferred from data on related species in the genus or family.
341 Finally, we ran two additional separate analyses to test whether social mating system and
342 environmental harshness confounded the ASR - SSD relationship. All numeric variables were
343 standardized before analyses to make parameter estimates comparable, and model
344 assumptions were also checked and met. We report two-tailed statistics. Sample sizes differed
345 between models because not all variables were available for all species (see Appendix S1).

346 In addition to PGLS models, we used phylogenetic path analyses (Santos 2012;
347 Gonzalez-Voyer and von Hardenberg 2014) to compare two sets of path models
348 corresponding to different hypotheses for the relationships linking ASR, SSD and sex
349 differences in mortality. Although path analyses – unlike experiments – cannot infer causality,
350 it is a suitable method to compare alternative scenarios representing different causal
351 relationships between variables (Shipley 2016). Model 1 assumes that sex-biased mortality
352 influences ASR, which in turn influences SSD through its effects on mating competition (as
353 proposed by the mating opportunity hypothesis; Fig. 1). Three variants of this model were
354 tested: Model 1a assumes that sex differences in both juvenile and adult mortality rates
355 influence ASR, while Models 1b-c include only one of these mortality effects. Model 2
356 assumes that SSD has sex-specific effects on juvenile and/or adult mortality, which then
357 drives ASR variation (representing the mortality cost hypothesis; Fig. 1). We tested all the
358 three variants of this latter scenario, assuming SSD effects on both juvenile and adult
359 mortality (Model 2a) or only on one mortality component (Models 2b-c).

360 We followed the approach proposed by Santos (2012) for phylogenetic path analyses.
361 In the first step, we conducted phylogenetic transformation on the data to control for effects of
362 phylogenetic relatedness among species. For this purpose, we (1) determined λ separately for
363 each variable by maximum likelihood, (2) used this variable-specific λ value to re-scale the
364 phylogenetic tree to a unit tree, and (3) used the transformed tree to calculate phylogenetically
365 independent contrasts for the variable (using ‘pic’ function of the R package ‘ape’; Paradis
366 2012). We repeated this process for each variable, and the resulting phylogenetically
367 transformed values were used for fitting path models. In the second step of the analyses, we
368 evaluated model fit using d-separation method (Shipley 2016) as implemented in the R
369 package ‘piecewiseSEM’ (Lefcheck 2016). In this method, Fisher’s *C* statistic is used to test
370 the goodness of fit of the whole path model, and the model is rejected (i.e. it does not provide

371 a good fit to the data) if the result of this C statistic is statistically significant (and conversely
372 a statistically non-significant result means acceptable fit; Lefcheck 2016). We compared
373 model fit between the six path models by their AICc values. Note that this approach ensures
374 that the same variables (i.e. the contrasts with the same phylogenetic signal) are used in each
375 path model, and that correlations are non-directional (i.e. for a pair of variables X and Y , $r_{XY} =$
376 r_{YX} as assumed in path analysis).

377 To test the robustness of the results, we repeated the path analyses using two other
378 methods. First, we repeated the above procedure (i.e. followed Santos 2012) except that we
379 used the covariance matrix comparison method for model fit instead of d-separation, as
380 implemented in the R package ‘lavaan’ (Rosseel 2012). Second, we repeated the analyses
381 using the method developed by von Hardenberg and Gonzalez-Voyer (2013). Unlike Santos’
382 (2012) method, in this latter approach a single value of Pagel’s λ is estimated for the residuals
383 of a regression of each pair of traits in a directional statistical model, rather than a value of λ
384 for each variable (see the Discussion and Appendix S3). We used the R package ‘phylopath’
385 (van der Bijl 2018) for this latter analysis, which relies on the d-separation method for model
386 fitting (similarly to ‘piecewiseSEM’, see above). We provide additional analyses to test the
387 robustness of the path analysis’ results in Appendix S3.

388

389 **RESULTS**

390 **Mating competition versus mating opportunity hypotheses**

391 Consistent with the mating opportunity hypothesis, and in contrast to the mating competition
392 hypothesis, we found a negative relationship between our measures of ASR and SSD: the size
393 of males relative to females increases when ASR becomes more female-skewed (Fig. 2, Table
394 1). This correlation was statistically significant when all species were analyzed together and
395 did not differ among the three amniote classes (ASR \times class interaction on SSD: $F_{2,456} =$

396 0.935, $P= 0.393$). The increase of SSD with increasingly female-skewed ASR was
397 statistically significant within birds and mammals but was not in reptiles when the three taxa
398 were analyzed separately (Fig. S1, Tables S1-4). These results remained consistent when we
399 used SSD estimates based on length instead of estimated mass in reptiles (Tables S1, S2 and
400 S5), when SSD for reptiles were estimated from published body mass data (Table S5), and
401 also when male mass was used as response variable (Table S5).

402 These results are robust because the direction of the ASR - SSD relationship and its
403 statistical significance were not sensitive to branch length assumptions (Table S6), and to the
404 inclusion of other predictors (Table 1). In multi-predictor models (Table 1), mean body mass
405 was positively related to SSD, supporting the Rensch rule (Abouheif and Fairbairn 1997), and
406 the type of sex determination influenced ASR variation as previously reported by Pipoly *et al.*
407 (2015). Nevertheless, ASR remained negatively associated with SSD when the effects of
408 mass and sex determination systems were accounted for (Table 1). This result also did not
409 change when environmental variation was included in the models using either breeding
410 latitude (Table 1) or environmental harshness (Table S5). Finally, excluding reptiles with
411 TSD (that have the lowest consistency in ASR; Bókony *et al.* 2019) or with assumed sex
412 determination also did not influence the relationship (Table S5).

413 The multi-predictor model for birds showed that species with aerial courtship displays
414 have lowered SSD as found in earlier studies (Jehl and Murray 1986; Székely *et al.* 2007);
415 however, the relationship between ASR and SSD remained statistically significant and
416 negative when this effect was included in the model (Table S3). Furthermore, data in birds
417 and mammals showed that, as expected, the relationship was weaker in monogamous than in
418 polygamous species, although the same trend occurred in both mating systems (Table S7).
419 Finally, reproductive mode was not associated with SSD or ASR in reptiles in our dataset
420 (Tables S1-2).

421

422 **Mating opportunity versus mortality costs hypotheses**

423 Both the mating opportunity hypothesis and the mortality cost hypothesis predict female-
424 skewed ASRs in species with male-biased SSD. However, our results are more consistent
425 with the mating opportunity hypothesis for two reasons. First, ASR but not SSD was
426 associated with the extent of sex differences in juvenile or adult mortality, and ASR remained
427 strongly and negatively correlated with SSD when sex differences in juvenile and adult
428 mortality were statistically controlled for (Table 1). Second, phylogenetic path analyses
429 showed that models of the mating opportunity hypothesis provided better fit to the data
430 (Models 1a-c, Fisher' C statistic: $P = 0.07 - 0.97$) than models corresponding to the mortality
431 cost hypothesis (Models 2a-c, $P < 0.001$; Table 2). The strongest support was for Model 1a
432 because it had the lowest AICc ($\Delta\text{AICc} = 4.1 - 43.2$; Table 2). This model proposes that sex-
433 biased mortality in both juveniles and adults generates skewed ASR, which in turn leads to
434 SSD biased towards the rarer sex (Fig. 3). These results are robust because we obtained the
435 same results when the analyses were repeated using two other implementations of the path
436 analysis (see Table S8 for the results obtained using 'phylopath', and Appendix S3 for the
437 results obtained using 'lavaan'). Finally, path analyses that excluded reptiles (for which the
438 ASR - SSD relationship was not statistically significant, see above) also yielded results
439 qualitatively consistent with the full dataset (Table S9).

440

441 **DISCUSSION**

442 Our analyses provided three major findings: (1) adult sex ratio is related to SSD among
443 amniote species, although the association is the opposite of the one proposed by Darwin; (2)
444 sex-biased mortality is unrelated to the extent of SSD in amniotes; and (3) confirmatory path
445 analyses indicate that sex-biased mortality influences ASR, which in turn induces changes in

446 SSD. Collectively, these findings support the mating opportunity hypothesis, indicating that
447 selection is likely to favor an increased resource allocation toward mating competition (by
448 growing and maintaining a large body mass) in the rarer sex, which has a higher chance of
449 getting mates than the other sex.

450 Theoretical models show that skewed ASRs can promote evolutionary changes that
451 may generate this association between ASR and SSD. First, models of sex role evolution
452 showed that skewed ASR can result in divergences in reproductive roles between the sexes
453 leading to less parental care and more frequent desertion and remating in the rarer sex and
454 opposite changes (i.e. more parental care and less frequent remating) in the more abundant
455 sex (Queller 1997; McNamara et al. 2000). Similarly, a demographic analysis based on the
456 relationships between mating systems and sex ratio, sex-specific patterns of survivorship, age
457 of first reproduction, and annual fecundity predicts that skewed ASRs promote the evolution
458 of polygamy (i.e. polygyny and polyandry in female-biased and male-biased populations,
459 respectively; Murray 1984). Since both frequent remating and polygamy can intensify sexual
460 selection, the above effects of skewed ASR can promote the evolution of SSD by favoring
461 increased body size in the rare sex. In line with the predictions of these models, an increasing
462 number of recent studies in birds and humans show that polygyny is more frequent and
463 parental care by males is reduced in female-skewed populations (Liker et al. 2013, 2014,
464 2015; Remeš et al. 2015; Schacht and Borgerhoff Mulder 2015; Eberhart-Phillips et al. 2018;
465 Grant and Grant 2019). Our results are also concordant with experimental studies in voles and
466 lizards, which reported that female-skewed ASRs exert directional selection for large body
467 size in males (Klemme et al. 2007; Fitze and Le Galliard 2008), and increase variance in male
468 reproductive success (Dreiss et al. 2010).

469 Theoretical models predict that the effects of ASR may depend on other life history
470 and behavioral traits of the populations. For example, Fromhage and Jennions (2016)

471 highlighted the importance of the specific processes generating ASR skews for the outcomes
472 of sex role evolution, and that a coevolutionary feedback between parental care and sexually
473 selected traits can greatly amplify sex role divergence. In addition, sexual competition for
474 mates may favor different traits in species with distinct ecology and behavior, leading to
475 inconsistent relationships between sex differences in mating competition and sexual
476 dimorphisms in behavioral or morphological trait across species (Clutton-Brock 2017).
477 Collectively, these factors may account for the relatively low amount of variation in SSD
478 explained by ASR in some of our analyses.

479 The association between intense sexual selection in males and female-skewed ASRs
480 was proposed decades ago by avian evolutionary ecologists (e.g. Mayr 1939), although it was
481 usually explained by the mortality cost hypothesis (Wittenberger 1976). Our analyses do not
482 support this hypothesis because sex-biased SSD is not associated with sex-biased juvenile or
483 adult mortality in the studied amniote species, and the results of the confirmatory path
484 analyses are also inconsistent with the mortality cost hypothesis. We propose that the lack of
485 relationship between SSD and sex differences in mortality may be explained by variation in
486 the environmental context (Lemaître et al. 2020). Studies in birds and mammals showed that
487 having a large body size may only be costly in terms of mortality in populations subjected to
488 harsh environmental conditions (Toigo and Gaillard 2003; Kalmbach and Benito 2007; Jones
489 et al. 2009; Clutton-Brock 2017). The effect of SSD may thus be reduced or absent when the
490 sex-specific mortality estimates correspond to average conditions, that may often be the case
491 in wild populations.

492 The ASR - SSD relationship may also be influenced by sex differences in the time of
493 maturation because longer maturation time in the larger sex can result in a shortage of that sex
494 in the adult population (Lovich et al. 2014) because immature life stages are generally
495 characterized by higher mortality (e.g. Gaillard et al. 2000). Furthermore, Fromhage &

496 Jennions (2016) showed that female-skewed sex ratios at maturation (MSR) can result in the
497 evolution of increased female care and male allocation to traits facilitating mating success.
498 Thus, if variation in ASR is determined at least in part by MSR, then the effects of sex-biased
499 MSR on sex roles can contribute to the observed association of ASR with the intensity of
500 mating competition, and, hence, SSD. This latter mechanism would deserve further
501 investigations.

502 Although the relationship between ASR and SSD is not statistically significant in
503 reptiles, it is qualitatively consistent with our findings in birds and mammals. Other selective
504 processes (e.g. fertility selection for large female size in indeterminate growers, Cox *et al.*
505 2007) might have masked the influence of sexual selection on SSD in reptiles. Consistent
506 with this explanation, selection often favors delayed maturation in female reptiles, which
507 enables them to produce larger clutches, which in turn also influences their body size and the
508 extent of SSD (Shine 2005; Agha *et al.* 2018). Follow-up studies using different proxies of
509 sexual selection are needed to investigate further how sexual selection is related to ASR in
510 reptiles.

511 Biased estimates of ASR may generate spurious relationship with SSD, which may
512 potentially affect our results. For example, the larger sex may have lower detectability in
513 polygamous species if some members of that sex are excluded from breeding sites (Ancona *et*
514 *al.* 2017). However, highly polygamous species in which populations have been thoroughly
515 surveyed showed skewed ASR even when all individuals in the population were accurately
516 counted (Granjon *et al.* 2017), and fairly consistent ASR estimates were obtained when both
517 breeding and non-breeding individuals were included (Emlen and Wrege 2004). In general,
518 ASR estimates show a moderate but statistically significant repeatability across populations in
519 most of the studied taxa, except reptiles with temperature-dependent sex determination
520 (Ancona *et al.* 2017; Bókony *et al.* 2019; Valentine Federico, J-F.L., J-M.G., A.L., I.P., T.S.

521 unpublished result), and in 80% of bird species the direction of ASR skew is the same for all
522 repeated estimates (Székely et al. 2014a).

523 The paths of causality in comparative data are difficult to untangle. Path analysis is a
524 valuable tool for contrasting different causal models, although it cannot reveal causality
525 (Shipley 2016). Path analysis assumes that each variable includes independent variations or
526 ‘errors’ and that these errors are independent among variables. This is not true for
527 comparative data, because the errors will be correlated across species. Our approach follows
528 Santos (2012), an innovative but overlooked method that satisfies the assumptions of path
529 analysis better than an alternative method based on phylogenetic regressions proposed by von
530 Hardenberg and Gonzalez-Voyer (2013). This latter approach is problematic because it is not
531 robust to changes in the specification of the model: if variable Y is regressed on X and λ
532 estimated, then the estimates of the partial correlations and λ may be different from those
533 obtained if Y is regressed on X with λ estimated (Appendix 3). The approach we have taken
534 avoids this problem. However, there is still room for methodological improvement. For
535 instance, our approach has the drawback of being a ‘subtractive’ comparative method (sensu
536 Harvey and Pagel 1991). The question of how to robustly fit complex path models for data on
537 multiple traits with different levels of phylogenetic signal is not straightforward.

538

539 *Concluding remarks*

540 Our findings indicate that sex-specific selection for large body size is associated with skewed
541 ASRs across amniotes, and this process appears to produce SSD biased towards the rare sex
542 in birds and mammals. Although this conclusion contrasts with Darwin’s initial suggestion
543 that intense sexual selection among males occurs when there is a surplus of males in the
544 population (Darwin 1871), theoretical and empirical work have suggested mechanisms that
545 can favor large size in the rare sex (Murray 1984; Klemme et al. 2007; Fitze and Le Galliard

546 2008; Dreiss et al. 2010). Further analyses of these processes and their application to species
547 with differing mating systems offer exciting opportunities for future investigations of the
548 interplay among sexual selection, SSD and ASR across the tree of life.
549

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800 **Table 1.** Phylogenetically-corrected analyses of sexual size dimorphism (SSD) and adult sex
801 ratio (ASR) in amniotes (reptiles, birds and mammals).

| Predictors | $b \pm SE$ | t | P | R^2 | A | n |
|---------------------------------------------|-----------------|-------|--------------|-------|----------|-----|
| (A) Response: sexual size dimorphism | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1 | | | | | | |
| ASR | - 0.168 ± 0.035 | 4.835 | < 0.001 | 0.119 | 0.868 ** | 462 |
| Mean body mass | 0.515 ± 0.086 | 5.980 | < 0.001 | | | |
| Model 2 | | | | | | |
| Juvenile mortality bias | 0.041 ± 0.065 | 0.629 | 0.531 | 0.129 | 0.703 ** | 100 |
| Mean body mass | 0.529 ± 0.131 | 4.051 | < 0.001 | | | |
| Model 3 | | | | | | |
| Adult mortality bias | - 0.021 ± 0.047 | 0.454 | 0.650 | 0.095 | 0.932 * | 230 |
| Mean body mass | 0.596 ± 0.117 | 5.090 | < 0.001 | | | |
| Multi-predictor model 1: | | | | | | |
| ASR | - 0.160 ± 0.035 | 4.555 | < 0.001 | 0.126 | 0.869 ** | 457 |
| Mean body mass | 0.515 ± 0.087 | 5.950 | < 0.001 | | | |
| Latitude | 0.004 ± 0.038 | 0.103 | 0.918 | | | |
| Sex determination, TSD ¹ | - 0.297 ± 0.251 | 1.184 | 0.237 | | | |
| Sex determination, ZW ¹ | - 0.685 ± 0.264 | 2.592 | 0.010 | | | |
| Multi-predictor model 2: | | | | | | |
| ASR | - 0.271 ± 0.061 | 4.452 | < 0.001 | 0.273 | 0.841 * | 97 |
| Mean body mass | 0.377 ± 0.134 | 2.824 | 0.006 | | | |
| Juvenile mortality bias | 0.001 ± 0.060 | 0.011 | 0.992 | | | |
| Adult mortality bias | - 0.019 ± 0.067 | 0.277 | 0.783 | | | |
| (B) Response: adult sex ratio | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1: SSD | | | | | | |
| SSD | - 0.234 ± 0.051 | 4.593 | < 0.001 | 0.042 | 0.359 ** | 462 |
| Model 2: Juvenile mortality bias | | | | | | |
| Juvenile mortality bias | - 0.214 ± 0.099 | 2.151 | 0.034 | 0.035 | 0.281 ** | 100 |
| Model 3: Adult mortality bias | | | | | | |
| Adult mortality bias | - 0.257 ± 0.060 | 4.313 | < 0.001 | 0.071 | 0.288 ** | 230 |
| Multi-predictor model 1: | | | | | | |
| SSD | - 0.188 ± 0.050 | 3.727 | < 0.001 | 0.071 | 0.247 ** | 457 |
| Mean body mass | - 0.106 ± 0.080 | 1.330 | 0.184 | | | |
| Latitude | - 0.095 ± 0.045 | 2.135 | 0.033 | | | |
| Sex determination, TSD ¹ | 0.481 ± 0.221 | 2.178 | 0.030 | | | |
| Sex determination, ZW ¹ | 0.712 ± 0.205 | 3.471 | < 0.001 | | | |
| Multi-predictor model 2: | | | | | | |
| SSD | - 0.457 ± 0.120 | 3.794 | < 0.001 | 0.402 | 0.030 # | 97 |
| Mean body mass | - 0.249 ± 0.108 | 2.316 | 0.023 | | | |
| Juvenile mortality bias | - 0.146 ± 0.086 | 1.702 | 0.092 | | | |
| Adult mortality bias | - 0.259 ± 0.100 | 2.591 | 0.011 | | | |

802 Results of separate predictor and multi-predictor phylogenetic generalized least-squares
803 (PGLS) models with either (A) SSD ($\log_{10}(\text{male mass}/\text{female mass})$) or (B) ASR (proportion
804 of males in the adult population) as dependent variable. **Separate predictor models with SSD**
805

806 as dependent variable also include $\log_{10}(\text{mean mass})$ as predictor (see Methods). Mortality
807 biases were calculated as $\log_{10}(\text{male mortality}/\text{female mortality})$ for juveniles and adults,
808 respectively. $b \pm SE$ is the model's parameter estimate with its standard error (intercepts are
809 not shown), t and P are the associated test statistic and its significance, λ is Pagel's lambda, n
810 is number of species.
811 * λ statistically different from 0, # λ statistically different from 1.
812 ¹ Differences from species with XY sex determination; overall effect of sex determination on
813 SSD: $F_{2,451} = 3.411$, $P = 0.034$; on ASR: $F_{2,451} = 6.135$, $P = 0.002$.

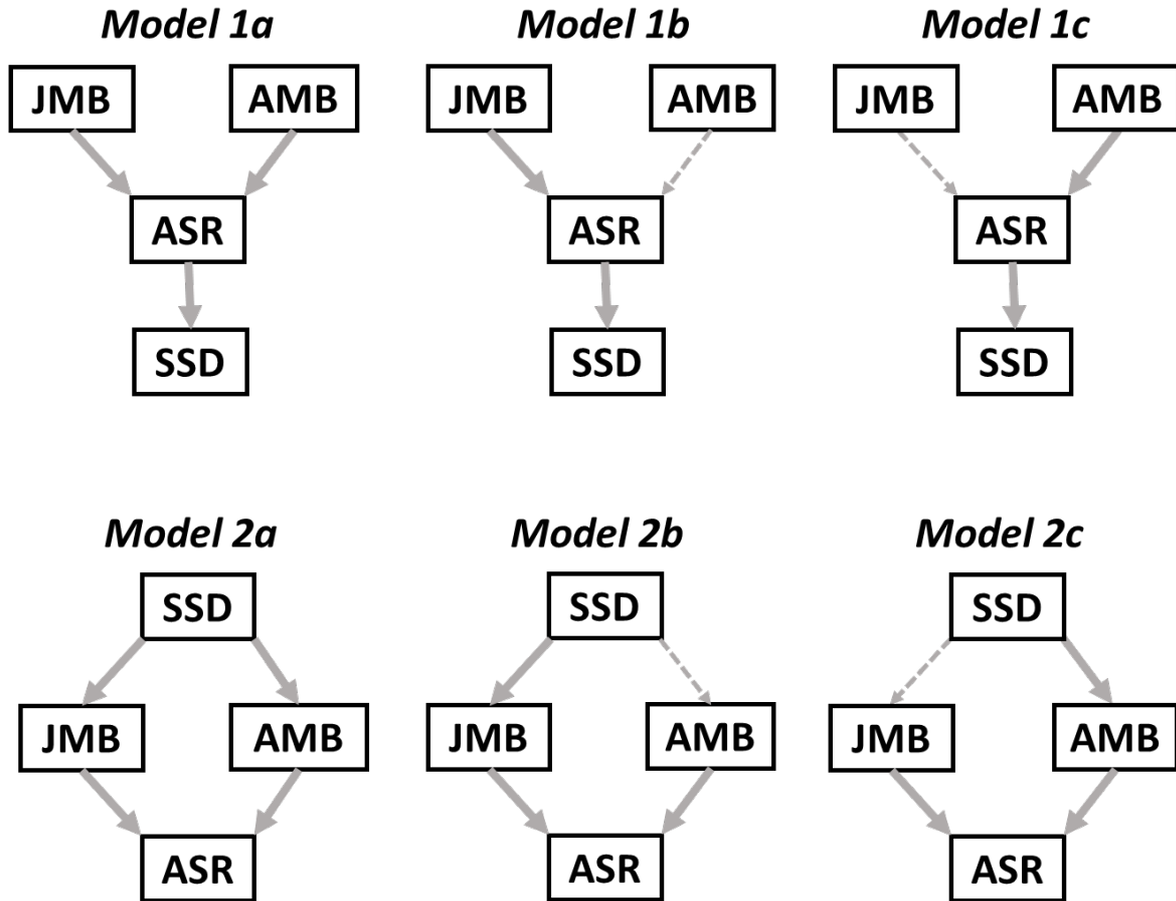
814 **Table 2.** Phylogenetic path models of the mating opportunity hypothesis (Models 1a-c) and
815 the mortality cost hypothesis (Models 2a-c) in amniotes (reptiles, birds and mammals).
816

| Model/Path | Path coefficient \pm SE | Z | P |
|--------------------------------------|----------------------------------------------------------------------------------------------------------------|---------|-------------------|
| Model 1a | $P_C = 0.972$, $df = 4$, $AICc = 15.8$, $\Delta AICc = 0.0$ | | |
| AMB \rightarrow ASR | - 0.340 \pm 0.113 | - 3.000 | 0.004 |
| JMB \rightarrow ASR | - 0.205 \pm 0.104 | - 1.970 | 0.052 |
| ASR \rightarrow SSD | - 0.425 \pm 0.074 | - 5.723 | < 0.001 |
| Model 1b | $P_C = 0.065$, $df = 6$, $AICc = 25.7$, $\Delta AICc = 9.9$ | | |
| (AMB \rightarrow ASR) ¹ | 0 | - | - |
| JMB \rightarrow ASR | - 0.258 \pm 0.107 | - 2.417 | 0.018 |
| ASR \rightarrow SSD | - 0.425 \pm 0.074 | - 5.723 | < 0.001 |
| Model 1c | $P_C = 0.376$, $df = 6$, $AICc = 19.9$, $\Delta AICc = 4.1$ | | |
| AMB \rightarrow ASR | - 0.378 \pm 0.113 | - 3.334 | 0.001 |
| (JMB \rightarrow ASR) ¹ | 0 | - | - |
| ASR \rightarrow SSD | - 0.425 \pm 0.074 | - 5.723 | < 0.001 |
| Model 2a | $P_C = 0.0$, $df = 4$, $AICc = 59.0$, $\Delta AICc = 43.2$ | | |
| SSD \rightarrow AMB | 0.171 \pm 0.105 | 1.631 | 0.106 |
| SSD \rightarrow JMB | 0.111 \pm 0.115 | 0.958 | 0.341 |
| AMB \rightarrow ASR | - 0.340 \pm 0.113 | - 3.000 | 0.004 |
| JMB \rightarrow ASR | - 0.205 \pm 0.104 | - 1.970 | 0.052 |
| Model 2b | $P_C = 0.0$, $df = 4$, $AICc = 50.4$, $\Delta AICc = 34.6$ | | |
| SSD \rightarrow JMB | 0.111 \pm 0.115 | 0.958 | 0.341 |
| AMB \rightarrow ASR | - 0.340 \pm 0.113 | - 3.000 | 0.004 |
| JMB \rightarrow ASR | - 0.205 \pm 0.104 | - 1.970 | 0.052 |
| Model 2c | $P_C = 0.0$, $AICc = 50.4$, $\Delta AICc = 34.6$ | | |
| SSD \rightarrow AMB | 0.171 \pm 0.105 | 1.631 | 0.106 |
| AMB \rightarrow ASR | - 0.340 \pm 0.113 | - 3.000 | 0.004 |
| JMB \rightarrow ASR | - 0.205 \pm 0.104 | - 1.970 | 0.052 |

849 Model structures are shown in Figure S1. SSD: sexual size dimorphism, ASR: adult sex ratio,
850 JMB and AMB: juvenile and adult mortality biases, respectively (variables are explained in
851 footnotes of Table 1). P_C is P -value for Fisher's C statistic for model fit, with non-significant
852 values (> 0.05) indicating an acceptable fit. $\Delta AICc$ indicates difference in $AICc$ values
853 between the most supported model (lowest $AICc$, Model 1a) and the focal models. $\Delta AICc > 2$
854 indicates substantially higher support for the best model than for the other models. The
855 analyses include 97 species of reptiles, birds and mammals with data for all for variables.

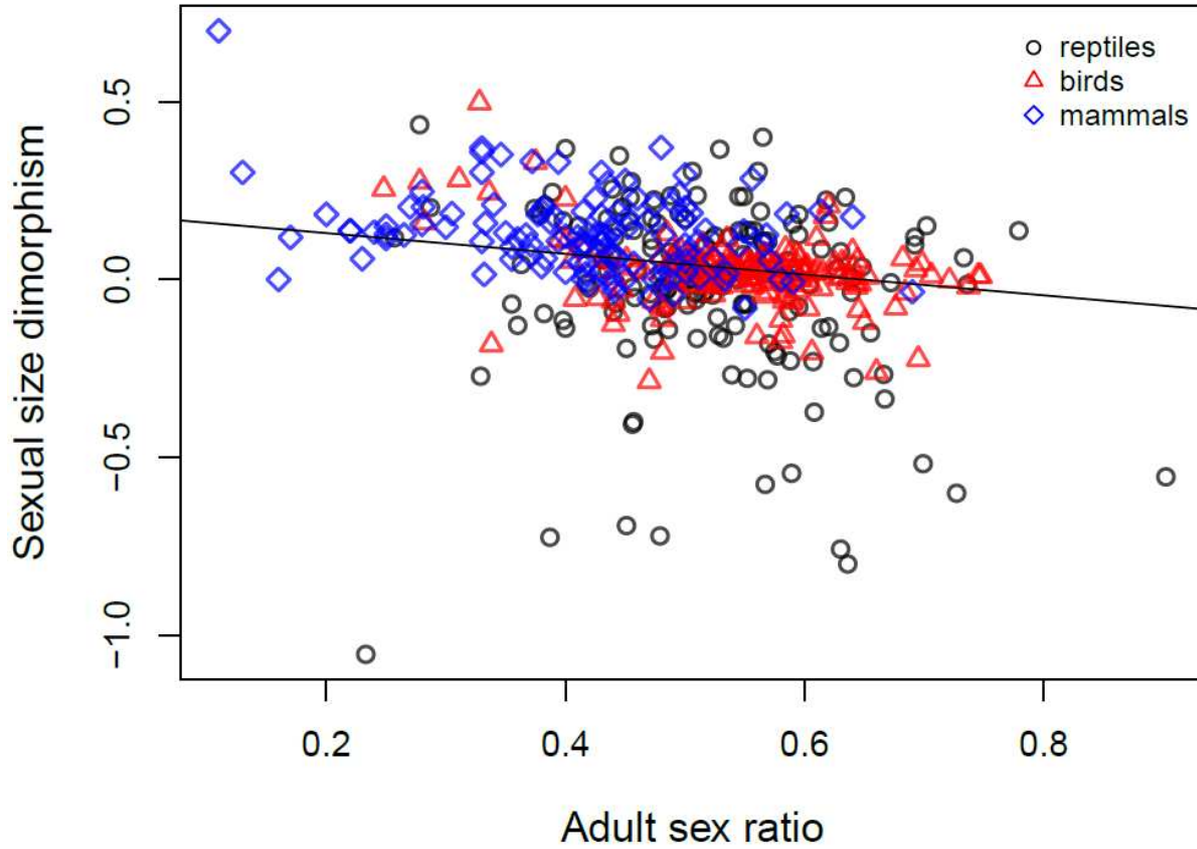
856 ¹ Path coefficient set to zero to keep the variable in the model.

857 **Figure 1.** Path models tested in the phylogenetic path analyses. SSD: sexual size dimorphism,
 858 ASR: adult sex ratio, JMB: juvenile mortality bias, AMB: adult mortality bias. Dashed arrows
 859 indicate paths with coefficients set to zero to keep the variable in the model. Models 1a-c and
 860 2a-c represent relationships as predicted by the mating opportunity hypothesis and the
 861 mortality cost hypothesis, respectively.
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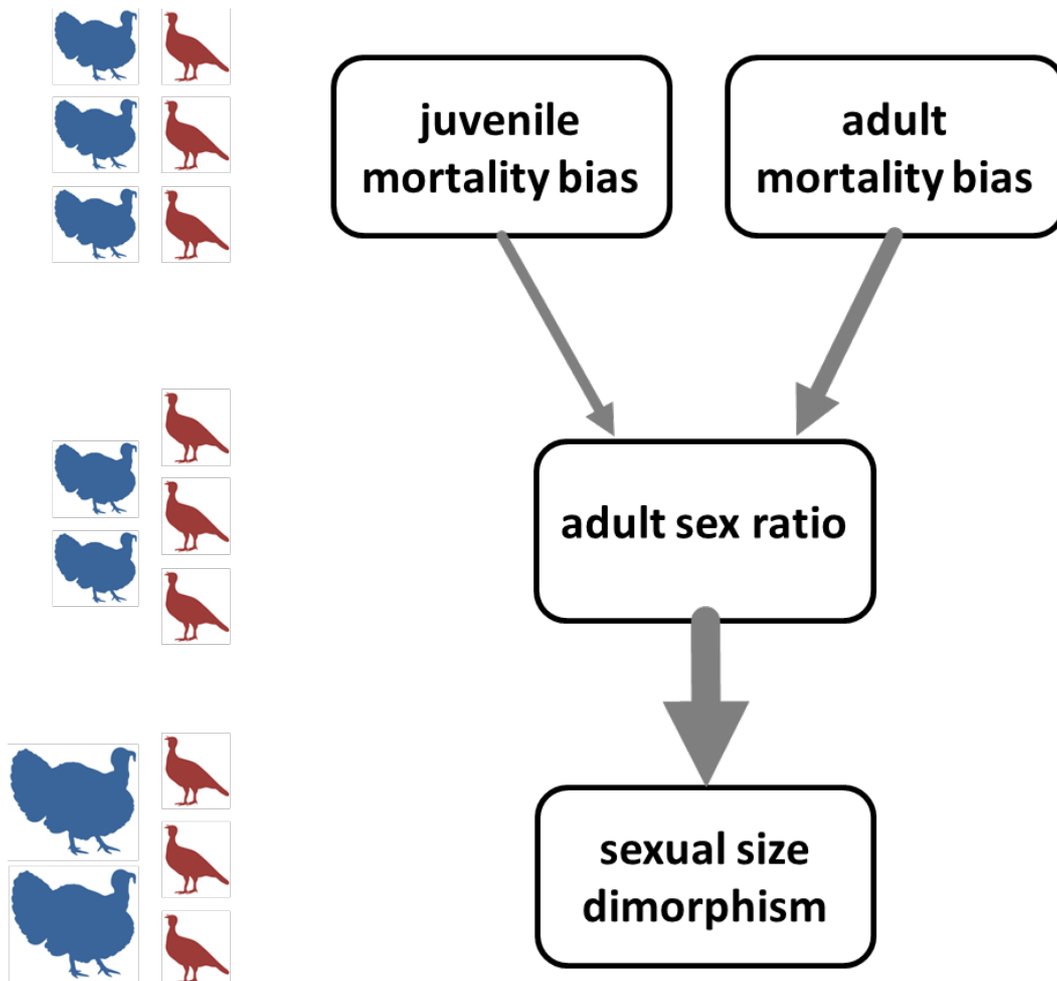
867 **Figure 2.** Sexual size dimorphism (SSD) in relation to adult sex ratio (ASR) in amniotes.
868 SSD was calculated as $\log_{10}(\text{male mass}/\text{female mass})$; ASR is the proportion of males in the
869 adult population. Each data point represents a species; the regression line is fitted by
870 phylogenetic generalized least-squares (PGLS) model (see Table 1 for statistics).
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875 **Figure 3.** Path diagram of the best-fitting phylogenetic path model (Model 1a in Table 2, $n =$
 876 97 species of reptiles, birds and mammals). The model supports the scenario that sex-biased
 877 juvenile and adult mortalities lead to skewed adult sex ratio, which in turn results in increased
 878 size dimorphism by sexual selection. Width of the arrows is proportional to path coefficients
 879 (see Table 2 for statistical details of the model). Bird pictures on the left illustrate the case
 880 when differential mortality generates female-skewed ASR, which then leads to a more male-
 881 biased SSD (i.e. larger body size in males relative to females). The path analyses were based
 882 on the approach proposed by Santos (2012), see Appendix S3 for details.
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Electronic Supporting Information: tables and figures

888 **Table S1.** Relationship between SSD, ASR and sex-biased mortalities in reptiles, using
889 estimated body mass data for SSD calculation.

| Predictors | $b \pm SE$ | t | P | R^2 | λ | n |
|---------------------------------------------|-----------------|-------|---------|---------|-----------|-----|
| (A) Response: sexual size dimorphism | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1 | | | | | | |
| ASR | - 0.123 ± 0.075 | 1.641 | 0.103 | 0.082 | 0.948 * | 155 |
| Mean body mass | 0.668 ± 0.177 | 3.774 | < 0.001 | | | |
| Model 2 | | | | | | |
| Juvenile mortality bias | - 0.414 ± 0.337 | 1.228 | 0.240 | 0.005 | 0.0 | 17 |
| Mean body mass | - 0.500 ± 0.440 | 1.136 | 0.275 | | | |
| Model 3 | | | | | | |
| Adult mortality bias | - 0.151 ± 0.117 | 1.287 | 0.203 | 0.092 | 1.0 * | 62 |
| Mean body mass | 0.737 ± 0.317 | 2.324 | 0.024 | | | |
| Multi-predictor model 1: | | | | 0.116 | 0.956 * | 153 |
| ASR | - 0.090 ± 0.075 | 1.203 | 0.231 | | | |
| Mean body mass | 0.715 ± 0.178 | 4.019 | < 0.001 | | | |
| Latitude | - 0.175 ± 0.126 | 1.389 | 0.167 | | | |
| Reproductive mode ¹ | 0.348 ± 0.313 | 1.112 | 0.268 | | | |
| Sex determination, TSD ² | - 0.463 ± 0.384 | 1.206 | 0.230 | | | |
| Sex determination, ZW ² | - 1.003 ± 0.313 | 2.344 | 0.020 | | | |
| Multi-predictor model 2: | | | | < 0.001 | 0.0 | 17 |
| ASR | - 0.022 ± 0.252 | 0.086 | 0.933 | | | |
| Mean body mass | - 0.452 ± 0.523 | 0.865 | 0.404 | | | |
| Juvenile mortality bias | - 0.500 ± 0.374 | 1.339 | 0.205 | | | |
| Adult mortality bias | 0.284 ± 0.429 | 0.662 | 0.520 | | | |
| (B) Response: adult sex ratio | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1: SSD | | | | | | |
| SSD | - 0.074 ± 0.061 | 1.209 | 0.228 | 0.003 | 0.171 # | 155 |
| Model 2: Juvenile mortality bias | | | | | | |
| Juvenile mortality bias | - 0.480 ± 0.415 | 1.156 | 0.266 | 0.021 | 0.0 | 17 |
| Model 3: Adult mortality bias | | | | | | |
| Adult mortality bias | - 0.159 ± 0.092 | 1.732 | 0.088 | 0.032 | 0.155 # | 62 |
| Multi-predictor model 1: | | | | 0.078 | 0.0 # | 153 |
| SSD | - 0.049 ± 0.055 | 0.891 | 0.374 | | | |
| Mean body mass | 0.173 ± 0.108 | 1.599 | 0.112 | | | |
| Latitude | - 0.001 ± 0.109 | 0.013 | 0.990 | | | |
| Reproductive mode ¹ | - 0.140 ± 0.216 | 0.650 | 0.517 | | | |
| Sex determination, TSD ² | 0.209 ± 0.224 | 0.934 | 0.352 | | | |
| Sex determination, ZW ² | 0.667 ± 0.216 | 3.091 | 0.002 | | | |
| Multi-predictor model 2: | | | | 0.165 | 0.0 # | 17 |
| SSD | - 0.028 ± 0.331 | 0.086 | 0.933 | | | |
| Mean body mass | 0.929 ± 0.556 | 1.671 | 0.121 | | | |
| Juvenile mortality bias | - 0.044 ± 0.459 | 0.095 | 0.926 | | | |
| Adult mortality bias | - 0.641 ± 0.465 | 1.377 | 0.194 | | | |

890

891 * Pagel's lambda statistically different from 0, # lambda statistically different from 1.

892 ¹ Differences from oviparous species.

893 ² Differences from XY species; overall effect of sex determination on SSD: $F_{2,146} = 2.8$, $P =$

894 0.066; on ASR: $F_{2,146} = 5.2$, $P = 0.006$.

895 For further explanation, see the footnotes of Table 1 in the main text.

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Table S2. Relationship between SSD, ASR and sex-biased mortalities in reptiles, using body length data for SSD calculation.

| Predictors | $b \pm SE$ | t | P | R^2 | λ | n |
|---------------------------------------------|-----------------|-------|---------|---------|-----------|-----|
| (A) Response: sexual size dimorphism | | | | | | |
| Separate predictor models: | | | | | | |
| Model 1 | | | | | | |
| ASR | - 0.008 ± 0.005 | 1.587 | 0.114 | 0.073 | 0.935 * | 155 |
| Mean body mass | 0.040 ± 0.011 | 3.562 | < 0.001 | | | |
| Model 2 | | | | | | |
| Juvenile mortality bias | - 0.035 ± 0.024 | 1.472 | 0.163 | 0.073 | 0.0 | 17 |
| Mean body mass | - 0.046 ± 0.031 | 1.485 | 0.160 | | | |
| Model 3 | | | | | | |
| Adult mortality bias | - 0.010 ± 0.007 | 1.402 | 0.166 | 0.086 | 1.0 * | 62 |
| Mean body mass | 0.044 ± 0.020 | 2.156 | 0.035 | | | |
| Multi-predictor model 1: | | | | | | |
| ASR | - 0.048 ± 0.042 | 1.126 | 0.262 | 0.122 | 0.952 * | 153 |
| Mean body mass | 0.391 ± 0.100 | 3.891 | < 0.001 | | | |
| Latitude | - 0.103 ± 0.071 | 1.459 | 0.147 | | | |
| Reproductive mode ¹ | 0.179 ± 0.177 | 1.015 | 0.312 | | | |
| Sex determination, TSD ² | - 0.223 ± 0.216 | 1.032 | 0.304 | | | |
| Sex determination, ZW ² | - 0.633 ± 0.241 | 2.628 | 0.010 | | | |
| Multi-predictor model 2: | | | | | | |
| ASR | - 0.059 ± 0.161 | 0.368 | 0.720 | < 0.001 | 0.0 | 17 |
| Mean body mass | - 0.347 ± 0.334 | 1.038 | 0.320 | | | |
| Juvenile mortality bias | - 0.354 ± 0.239 | 1.484 | 0.164 | | | |
| Adult mortality bias | 0.092 ± 0.274 | 0.337 | 0.742 | | | |
| (B) Response: adult sex ratio | | | | | | |
| Separate predictor models: | | | | | | |
| Model 1: SSD | | | | | | |
| SSD | - 0.131 ± 0.109 | 1.209 | 0.229 | 0.003 | 0.169 # | 155 |
| Model 2: Juvenile mortality bias | | | | | | |
| Juvenile mortality bias | - 0.480 ± 0.415 | 1.156 | 0.266 | 0.021 | 0.0 | 17 |
| Model 2: Adult mortality bias | | | | | | |
| Adult mortality bias | - 0.159 ± 0.092 | 1.732 | 0.088 | 0.032 | 0.155 # | 62 |
| Multi-predictor model 1: | | | | | | |
| SSD | - 0.085 ± 0.098 | 0.870 | 0.386 | 0.078 | 0.0 # | 153 |
| Mean body mass | 0.172 ± 0.109 | 1.588 | 0.114 | | | |
| Latitude | - 0.001 ± 0.109 | 0.007 | 0.994 | | | |
| Reproductive mode ¹ | - 0.141 ± 0.216 | 0.654 | 0.514 | | | |
| Sex determination, TSD ² | 0.214 ± 0.223 | 0.958 | 0.340 | | | |
| Sex determination, ZW ² | 0.667 ± 0.216 | 3.089 | 0.002 | | | |
| Multi-predictor model 2: | | | | | | |
| SSD | - 0.188 ± 0.512 | 0.368 | 0.720 | 0.174 | 0.0 # | 17 |
| Mean body mass | 0.867 ± 0.570 | 1.522 | 0.154 | | | |
| Juvenile mortality bias | - 0.096 ± 0.463 | 0.208 | 0.839 | | | |
| Adult mortality bias | - 0.624 ± 0.457 | 1.366 | 0.197 | | | |

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* Pagel's lambda statistically different from 0, # lambda statistically different from 1.

¹ Differences from oviparous species.

901 ² Differences from XY species; overall effect of sex determination on SSD: $F_{2,146} = 3.7$, $P =$
902 0.028; on ASR: $F_{2,146} = 5.2$, $P = 0.006$.
903 For further explanation, see the footnotes of Table 1 in the main text.
904

905 **Table S3.** Relationship between SSD, ASR and sex-biased mortalities in birds.

| Predictors | $b \pm SE$ | t | P | R^2 | λ | n |
|---------------------------------------------|-----------------|-------|---------|-------|-----------|-----|
| (A) Response: sexual size dimorphism | | | | | | |
| Separate predictor models: | | | | | | |
| Model 1 | | | | | | |
| ASR | - 0.242 ± 0.037 | 6.625 | < 0.001 | 0.250 | 0.812 *# | 185 |
| Mean body mass | 0.293 ± 0.105 | 2.798 | 0.006 | | | |
| Model 2 | | | | | | |
| Juvenile mortality bias | 0.066 ± 0.073 | 0.898 | 0.374 | 0.130 | 0.095 * | 47 |
| Mean body mass | 0.735 ± 0.270 | 2.722 | 0.009 | | | |
| Model 3 | | | | | | |
| Adult mortality bias | 0.068 ± 0.051 | 1.335 | 0.184 | 0.072 | 0.708 *# | 123 |
| Mean body mass | 0.372 ± 0.130 | 2.870 | 0.005 | | | |
| Multi-predictor model 1: | | | | | | |
| ASR | - 0.242 ± 0.038 | 6.390 | < 0.001 | 0.272 | 0.720 ** | 178 |
| Mean body mass | 0.261 ± 0.100 | 2.599 | 0.010 | | | |
| Latitude | - 0.020 ± 0.034 | 0.584 | 0.560 | | | |
| Display type, agile ¹ | - 0.338 ± 0.090 | 3.748 | < 0.001 | | | |
| Multi-predictor model 2: | | | | | | |
| ASR | - 0.346 ± 0.080 | 4.318 | < 0.001 | 0.386 | 1.0 * | 47 |
| Mean body mass | 0.424 ± 0.246 | 1.719 | 0.093 | | | |
| Juvenile mortality bias | - 0.032 ± 0.065 | 0.489 | 0.627 | | | |
| Adult mortality bias | - 0.068 ± 0.089 | 0.763 | 0.450 | | | |
| (B) Response: adult sex ratio | | | | | | |
| Separate predictor models: | | | | | | |
| Model 1: SSD | | | | | | |
| SSD | - 0.746 ± 0.114 | 6.520 | < 0.001 | 0.184 | 0.480 ** | 185 |
| Model 2: Juvenile mortality bias | | | | | | |
| Juvenile mortality bias | - 0.354 ± 0.115 | 3.084 | 0.003 | 0.156 | 0.0 # | 47 |
| Model 3: Adult mortality bias | | | | | | |
| Adult mortality bias | - 0.384 ± 0.079 | 4.866 | < 0.001 | 0.157 | 0.0 # | 123 |
| Multi-predictor model 1: | | | | | | |
| SSD | - 0.717 ± 0.116 | 6.183 | < 0.001 | 0.239 | 0.244 # | 178 |
| Mean body mass | - 0.191 ± 0.136 | 1.406 | 0.161 | | | |
| Latitude | - 0.127 ± 0.058 | 2.201 | 0.029 | | | |
| Display type, agile ¹ | - 0.589 ± 0.161 | 3.667 | < 0.001 | | | |
| Multi-predictor model 2: | | | | | | |
| SSD | - 0.382 ± 0.153 | 2.499 | 0.016 | 0.397 | 0.0 # | 47 |
| Mean body mass | - 0.128 ± 0.198 | 0.646 | 0.522 | | | |
| Juvenile mortality bias | - 0.199 ± 0.109 | 1.831 | 0.074 | | | |
| Adult mortality bias | - 0.468 ± 0.139 | 3.368 | 0.002 | | | |

906 * Pagel's lambda statistically different from 0, # lambda statistically different from 1.

907 ¹ Difference from non-agile species.

908 For further explanation, see the footnotes of Table 1 in the main text.

909

910

911 **Table S4.** Relationship between SSD, ASR and sex-biased mortalities in mammals.

| Predictors | $b \pm SE$ | t | P | R^2 | λ | n |
|---------------------------------------------|-----------------|-------|-------------------|---------|-----------|-----|
| (A) Response: sexual size dimorphism | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1 | | | | | | |
| ASR | - 0.170 ± 0.058 | 2.946 | 0.004 | 0.143 | 0.313 ** | 122 |
| Mean body mass | 0.385 ± 0.129 | 2.979 | 0.004 | | | |
| Model 2 | | | | | | |
| Juvenile mortality bias | 0.089 ± 0.123 | 0.719 | 0.477 | 0.078 | 0.233 # | 36 |
| Mean body mass | 0.515 ± 0.233 | 2.214 | 0.034 | | | |
| Model 3 | | | | | | |
| Adult mortality bias | 0.025 ± 0.103 | 0.244 | 0.809 | 0.056 | 0.217 # | 45 |
| Mean body mass | 0.424 ± 0.202 | 2.093 | 0.042 | | | |
| Multi-predictor model 1: | | | | | | |
| ASR | - 0.153 ± 0.058 | 2.646 | 0.009 | 0.166 | 0.342 ** | 120 |
| Mean body mass | 0.418 ± 0.131 | 3.191 | 0.002 | | | |
| Latitude | 0.106 ± 0.053 | 2.016 | 0.046 | | | |
| Multi-predictor model 2: | | | | | | |
| ASR | - 0.374 ± 0.129 | 2.900 | 0.007 | 0.250 | 0.0 # | 33 |
| Mean body mass | 0.209 ± 0.237 | 0.880 | 0.386 | | | |
| Juvenile mortality bias | 0.088 ± 0.116 | 0.762 | 0.452 | | | |
| Adult mortality bias | - 0.016 ± 0.119 | 0.134 | 0.894 | | | |
| (B) Response: adult sex ratio | | | | | | |
| <i>Separate predictor models:</i> | | | | | | |
| Model 1: SSD | | | | | | |
| SSD | - 0.460 ± 0.130 | 3.539 | < 0.001 | 0.087 | 0.252 ** | 122 |
| Model 2: Juvenile mortality bias | | | | | | |
| Juvenile mortality bias | - 0.032 ± 0.166 | 0.195 | 0.847 | < 0.001 | 0.0 # | 36 |
| Model 3: Adult mortality bias | | | | | | |
| Adult mortality bias | - 0.076 ± 0.155 | 0.493 | 0.624 | < 0.001 | 0.0 # | 45 |
| Multi-predictor model 1: | | | | | | |
| SSD | - 0.375 ± 0.140 | 2.670 | 0.009 | 0.093 | 0.320 ** | 120 |
| Mean body mass | - 0.314 ± 0.209 | 1.500 | 0.136 | | | |
| Latitude | - 0.075 ± 0.083 | 0.907 | 0.366 | | | |
| Multi-predictor model 2: | | | | | | |
| SSD | - 0.617 ± 0.213 | 2.900 | 0.007 | 0.293 | 0.0 # | 33 |
| Mean body mass | - 0.494 ± 0.294 | 1.678 | 0.104 | | | |
| Juvenile mortality bias | - 0.043 ± 0.150 | 0.285 | 0.778 | | | |
| Adult mortality bias | 0.022 ± 0.153 | 0.142 | 0.888 | | | |

912
 913 * Pagel's lambda statistically different from 0, # lambda statistically different from 1.
 914 For further explanation, see the footnotes of Table 1 in the main text.
 915
 916

917 **Table S5.** Sensitivity analyses of the relationship between sexual size dimorphism (SSD,
918 dependent variable in all models) and adult sex ratio (ASR). Table shows results when (A)
919 male mass (instead of $\log_{10}(\text{male mass} / \text{female mass})$ is used as response variable, (B)
920 reptiles are included with SSD based on body length, (C) reptiles are included with SSD
921 calculated from sex-specific body mass, (D) reptiles with temperature-dependent sex
922 determination (TSD) are excluded, (E) reptiles with assumed sex determination, based on
923 related species, are excluded, and (F) environmental harshness is included in the model.
924

| Predictors | $b \pm SE$ | t | P | R^2 | λ | n |
|----------------------------------------------------------------------------------------|----------------|---------|---------|-------|-----------|-----|
| (A) Male body mass as response variable (all species): | | | | 0.957 | 0.846*# | 462 |
| ASR | -0.020 ± 0.004 | 4.953 | < 0.001 | | | |
| Female body mass | 1.008 ± 0.010 | 100.658 | < 0.001 | | | |
| (B) Reptiles' SSD calculated from body length (all species): | | | | 0.139 | 0.703*# | 462 |
| ASR | -0.234 ± 0.038 | 6.231 | < 0.001 | | | |
| Mean body mass | 0.473 ± 0.085 | 5.575 | < 0.001 | | | |
| (C) Reptiles' SSD calculated from body mass¹ (all species): | | | | 0.144 | 0.761*# | 338 |
| ASR | -0.271 ± 0.050 | 5.437 | < 0.001 | | | |
| Mean body mass | 0.495 ± 0.112 | 4.426 | < 0.001 | | | |
| (D) TSD reptiles excluded (all species): | | | | 0.132 | 0.791*# | 402 |
| ASR | -0.250 ± 0.043 | 5.767 | < 0.001 | | | |
| Mean body mass | 0.507 ± 0.105 | 4.814 | < 0.001 | | | |
| (E) Reptiles with assumed sex determination excluded² (all species): | | | | 0.125 | 0.860*# | 409 |
| ASR | -0.167 ± 0.036 | 4.669 | < 0.001 | | | |
| Mean body mass | 0.502 ± 0.088 | 5.710 | < 0.001 | | | |
| (F) Effect of environmental harshness³: | | | | | | |
| birds and mammals: | | | | 0.141 | 0.763*# | 219 |
| ASR | -0.164 ± 0.037 | 4.440 | < 0.001 | | | |
| Environmental harshness | 0.039 ± 0.045 | 0.855 | 0.394 | | | |
| Mean body mass | 0.263 ± 0.088 | 2.966 | 0.003 | | | |
| reptiles: | | | | 0.026 | 0.957* | 58 |
| ASR | -0.081 ± 0.144 | 0.562 | 0.576 | | | |
| Environmental harshness | 0.105 ± 0.064 | 1.624 | 0.110 | | | |
| Mean body mass | 0.294 ± 0.302 | 0.975 | 0.334 | | | |
| all species: | | | | 0.111 | 0.867*# | 277 |
| ASR | -0.153 ± 0.038 | 4.012 | < 0.001 | | | |
| Environmental harshness | 0.076 ± 0.033 | 2.295 | 0.023 | | | |
| Mean body mass | 0.297 ± 0.091 | 3.256 | 0.001 | | | |

925
926 * Pagel's lambda statistically different from 0, # lambda statistically different from 1

927 ¹ Sex-specific body mass data from Myhrvold et al. (2015).

928 ² Sex determination mechanism assumed to be the same type as reported for the genus or
929 family (see Methods).

930 ³ The influence of environmental harshness was tested in birds and mammals using data from
931 Botero et al. (2014), in reptiles using data calculated in this study (following the method of
932 Botero et al 2014), and in all species by pooling the harshness scores from the two studies.

933 **Table S6.** Analyses of the relationship between SSD (dependent variable) and ASR with
 934 branch lengths calculated by three different methods for the phylogeny used in the PGLS
 935 models. The analyses included reptiles, birds, and mammals.
 936

| Predictors | <i>b</i> ± <i>SE</i> | <i>t</i> | <i>P</i> | <i>R</i>² | <i>λ</i> | <i>n</i> |
|-------------------------------|-----------------------------|-----------------|-----------------|-----------------------------|-----------------|-----------------|
| (A) Nee's method | | | | | | |
| | | | | <i>0.119</i> | <i>0.868 *#</i> | <i>462</i> |
| ASR | - 0.168 ± 0.035 | 4.835 | < 0.001 | | | |
| Mean body mass | 0.515 ± 0.086 | 5.980 | < 0.001 | | | |
| (B) Pagel's method | | | | | | |
| | | | | <i>0.124</i> | <i>0.869 *#</i> | <i>462</i> |
| ASR | - 0.166 ± 0.034 | 4.826 | < 0.001 | | | |
| Mean body mass | 0.564 ± 0.090 | 6.282 | < 0.001 | | | |
| (C) Unit branch length | | | | | | |
| | | | | <i>0.148</i> | <i>1.0 *</i> | <i>462</i> |
| ASR | - 0.179 ± 0.032 | 5.577 | < 0.001 | | | |
| Mean body mass | 0.565 ± 0.085 | 6.682 | < 0.001 | | | |

937
 938 * Pagel's lambda statistically different from 0, # lambda statistically different from 1
 939 ¹ See Methods for details of branch length calculations
 940
 941
 942
 943

944 **Table S7.** Analyses of the relationship between SSD (dependent variable) and ASR in
 945 socially monogamous and socially polygamous species, respectively. The analyses included
 946 birds and mammals.
 947

| Predictors | <i>b</i> ± <i>SE</i> | <i>t</i> | <i>P</i> | <i>R</i>² | <i>λ</i> | <i>n</i> |
|-----------------------|-----------------------------|-----------------|-----------------|-----------------------------|-----------------|-----------------|
| (A) Monogamy | | | | | | |
| | | | | <i>0.022</i> | <i>1.0 *</i> | <i>109</i> |
| ASR | - 0.066 ± 0.038 | 1.717 | 0.089 | | | |
| Mean body mass | 0.087 ± 0.083 | 1.044 | 0.299 | | | |
| (B) Polygamy | | | | | | |
| | | | | <i>0.222</i> | <i>0.418 *#</i> | <i>162</i> |
| ASR | - 0.223 ± 0.048 | 4.607 | < 0.001 | | | |
| Mean body mass | 0.399 ± 0.105 | 3.790 | < 0.001 | | | |

948
 949 * Pagel's lambda statistically different from 0, # lambda statistically different from 1
 950 ¹ When monogamous and polygynous species are analyzed together, there is a statistically
 951 significant interaction between the effects of mating system and ASR (*b* ± *SE* = 0.218 ± 0.087,
 952 *t* = 2.510, *P* = 0.013).

953 **Table S8.** Results of the phylogenetic path analyses using the R package ‘phylopath’. Models
 954 represent the mating opportunity hypothesis (Models 1a-c) and the mortality cost hypothesis
 955 (Models 2a-c). Analyses based on data of all species (birds, mammals, and reptiles; $n=97$
 956 species).
 957

| Model | <i>k</i> | <i>q</i> | <i>C</i> | <i>P</i> | <i>CICc</i> | Δ<i>CICc</i> |
|------------------------|-----------------|-----------------|-----------------|-----------------|--------------------|---------------------------------------|
| <i>Model 1a</i> | 3 | 7 | 6.4 | 0.383 | 21.6 | 0.0 |
| <i>Model 1b</i> | 4 | 6 | 18.7 | 0.017 | 31.6 | 10.0 |
| <i>Model 1c</i> | 4 | 6 | 11.2 | 0.188 | 24.2 | 2.6 |
| <i>Model 2a</i> | 2 | 8 | 32.4 | <0.001 | 50.0 | 28.4 |
| <i>Model 2b</i> | 3 | 7 | 34.8 | <0.001 | 50.0 | 28.4 |
| <i>Model 2c</i> | 3 | 7 | 36.6 | <0.001 | 51.9 | 30.3 |

959 Model structures are shown in Figure 1. SSD: sexual size dimorphism, ASR: adult sex ratio,
 960 JMB and AMB: juvenile and adult mortality biases, respectively. The table shows the number
 961 of independence claims (*k*), the number of parameters (*q*), Fisher’s *C* statistic (*C*) and its
 962 accompanying probability (*P*), C-statistic information criterion corrected for small sample
 963 sizes (*CICc*), and the difference in *CICc* from the top model (Δ *CICc*). A *P*-value less than
 964 0.05 indicates a poor model fit (i.e. rejection of the model), whereas a Δ *CICc* > 2 indicates
 965 substantial support for the top path model over the alternative models.
 966
 967

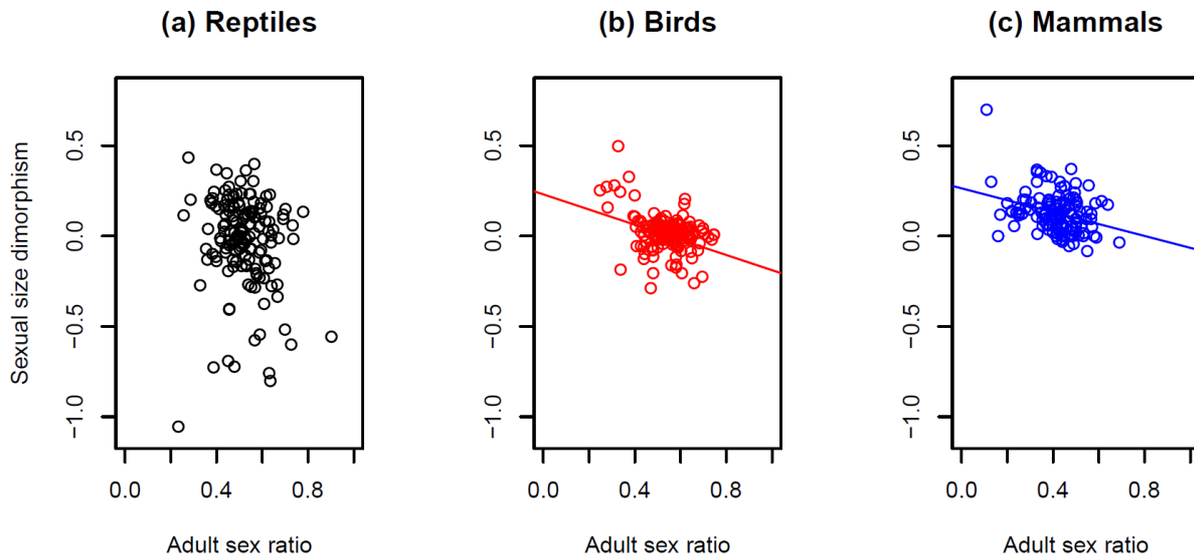
968 **Table S9.** Phylogenetic path models representing the mating opportunity hypothesis (Models
 969 1a-c) and the mortality cost hypothesis (Models 2a-c). Analyses with data of birds and
 970 mammals (i.e. excluding reptiles; $n= 81$ species).
 971

| Model/Path | Path coefficient $\pm SE$ | Z | P |
|--------------------------------------|-----------------------------------------------------------------------|---------|-------------------|
| Model 1a | $P_C= 0.991, df= 4, AIC_c= 15.9, \Delta AIC_c= 0.0$ | | |
| AMB \rightarrow ASR | - 0.321 \pm 0.121 | - 2.662 | 0.009 |
| JMB \rightarrow ASR | - 0.210 \pm 0.109 | - 1.920 | 0.059 |
| ASR \rightarrow SSD | - 0.719 \pm 0.122 | - 5.887 | < 0.001 |
| Model 1b | $P_C= 0.142, df= 6, AIC_c= 23.7, \Delta AIC_c= 7.8$ | | |
| (AMB \rightarrow ASR) ¹ | 0 | - | - |
| JMB \rightarrow ASR | - 0.257 \pm 0.112 | - 2.289 | 0.025 |
| ASR \rightarrow SSD | - 0.719 \pm 0.122 | - 5.887 | < 0.001 |
| Model 1c | $P_C= 0.428, df= 6, AIC_c= 19.7, \Delta AIC_c= 3.8$ | | |
| AMB \rightarrow ASR | - 0.358 \pm 0.121 | - 2.958 | 0.004 |
| (JMB \rightarrow ASR) ¹ | 0 | - | - |
| ASR \rightarrow SSD | - 0.719 \pm 0.122 | - 5.887 | < 0.001 |
| Model 2a | $P_C= 0.0, df= 4, AIC_c= 59.7, \Delta AIC_c= 43.8$ | | |
| SSD \rightarrow AMB | 0.117 \pm 0.077 | 1.503 | 0.137 |
| SSD \rightarrow JMB | 0.102 \pm 0.086 | 1.187 | 0.239 |
| AMB \rightarrow ASR | - 0.321 \pm 0.121 | - 2.662 | 0.009 |
| JMB \rightarrow ASR | - 0.210 \pm 0.109 | - 1.920 | 0.059 |
| Model 2b | $P_C= 0.0, df= 4, AIC_c= 50.5, \Delta AIC_c= 34.6$ | | |
| SSD \rightarrow JMB | 0.102 \pm 0.086 | 1.187 | 0.239 |
| AMB \rightarrow ASR | - 0.321 \pm 0.121 | - 2.662 | 0.009 |
| JMB \rightarrow ASR | - 0.210 \pm 0.109 | - 1.920 | 0.059 |
| Model 2c | $P_C= 0.0, AIC_c= 50.5, \Delta AIC_c= 34.6$ | | |
| SSD \rightarrow AMB | 0.117 \pm 0.077 | 1.503 | 0.137 |
| AMB \rightarrow ASR | - 0.321 \pm 0.121 | - 2.662 | 0.009 |
| JMB \rightarrow ASR | - 0.210 \pm 0.109 | - 1.920 | 0.059 |

1004 Model structures are shown in Figure 1. SSD: sexual size dimorphism, ASR: adult sex ratio,
 1005 JMB and AMB: juvenile and adult mortality biases, respectively (variables are explained in
 1006 footnotes of Table 1). P_C is P -value for Fisher's C statistic for model fit, with non-significant
 1007 values (> 0.05) indicating an acceptable fit. ΔAIC_c indicates difference in AIC_c values
 1008 between the most supported model (lowest AIC_c , Model 1a) and the focal models. $\Delta AIC_c > 2$
 1009 indicates substantially higher support for the best model than for the other model.

1010 ¹ Path coefficient set to zero to keep the variable in the model.
 1011

1012 **Figure S1.** Sexual size dimorphism in relation to adult sex ratio in (a) reptiles (PGLS, $b \pm SE$
1013 $= -0.123 \pm 0.075$, $P=0.103$, $n=155$ species), (b) birds ($b \pm SE = -0.242 \pm 0.037$, $P < 0.001$,
1014 $n=185$), and (c) mammals ($b \pm SE = -0.170 \pm 0.058$, $P=0.004$, $n=122$). Each data point
1015 represents a species, and lines show statistically significant regressions fitted by PGLS (see
1016 Tables S1-4 for further statistical details).
1017
1018



1019

Electronic Supporting Information: Appendix S1

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1021

1022 References for data sources are provided in a separate excel file. The full dataset
1023 will be published together with references after the manuscript is accepted for
1024 publication.
1025

Electronic Supporting Information: Appendix S2

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Parameters of the allometric equations between body length and body mass ($\log_{10}(\text{mass in g}) = a + b \cdot \log_{10}(\text{length in mm})$) used for the calculation of mass estimates in reptiles. n is the number of species included in the analyses.

| Taxon | Intercept (a) | Slope (b) | n | Reference |
|----------------------------|-----------------------------------|-------------------------------|-----------------------|--------------------------|
| Snakes | -5.773 | 2.786 | 336 | Feldman and Meiri (2013) |
| Squamates and crocodilians | -4.52 | 2.923 | 600 | Meiri (2010) |
| Turtles | -3.535 | 2.887 | 199 | Regis and Meik (2017) |

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References:

Feldman, A., and S. Meiri. 2013. Length-mass allometry in snakes. *Biol. J. Linn. Soc.* 108:161–172.

Meiri, S. 2010. Length-weight allometries in lizards. *J. Zool.* 281:218–226.

Regis, K. W., and J. M. Meik. 2017. Allometry of sexual size dimorphism in turtles: a comparison of mass and length data. *PeerJ* 5:e2914.

Electronic Supporting Information: Appendix S3

Methodological notes on path analyses applied to comparative data

Several approaches have been proposed for applying path analysis in phylogenetic comparative studies (von Hardenberg and Gonzalez-Voyer 2013). However, some methodological aspects of this method still pose challenges for its phylogenetic applications. Here we discuss two of such aspects: (1) the problem of using bivariate phylogenetic generalized least squares (PGLS) models to estimate correlations between variables in the path models, and (2) the reliability of AIC statistics to compare non-nested path models. We suggest ways to avoid these problems which may help further applications of path analysis to phylogenetic data.

1. The problem of using PGLS in path analyses

The analysis of multivariate dependent data is a notoriously thorny problem. In this dataset we, as in common with many similar analyses, had to deal with the problem of phylogenetic non-independence (Harvey and Pagel 1991). Techniques for analyzing cause-effect relationships using linear models are well developed (Felsenstein 1985; Harvey and Pagel 1991; Hansen and Martins 1996; Pagel 1997; Freckleton et al. 2002). These techniques are designed for analysis of data in which there is a dependent variable of interest, which is modelled as a function of independent predictors. In these models the effect of phylogeny is accounted for by modelling phylogenetic dependence in the residual term, and we have used these for several analyses (e.g. Table 1 in the main text). However, analyses of data in which variables are treated as multivariate responses are much less common. Phylogenetic principal components analysis is one exception, although recent research has stressed that this may be more complex than previously realized (Uyeda et al. 2015).

Path analysis is a method of multivariate trait analysis that allows complex dependencies among variables to be modelled. von Hardenberg & Gonzalez-Voyer (2013) presented a method for performing path analysis on phylogenetically dependent data. This approach accounts for phylogenetic non-independence through constructing a series of bivariate PGLS models in which one variable is treated as the dependent variable, and the other is the independent variable. Importantly, this approach permits variable levels of

1073 dependence to be modelled through estimating Pagel's λ , which accounts for varying
1074 contributions of phylogeny to trait variation.

1075 However, the analysis of such data is complex. Consider a simple example, in which
1076 we have three variables x , y , and z . If we model $x \rightarrow y$ (i.e. where x is the predictor and y is
1077 the response variable) using the PGLS- λ approach, we might well estimate a different
1078 correlation than if we model $y \rightarrow x$. This is because in PGLS the estimate of λ depends on the
1079 direction of the relationship, and this affects the estimate of the correlation. In path analysis
1080 correlations between pairs of variables are the input and the directionality of the statistical
1081 model should not be an issue. Specifically, the net correlation r_{xz} should then be the product
1082 of the component pairwise correlations, i.e. $r_{xz} = r_{xy} \times r_{yz}$. However, if we use PGLS and $\lambda \neq 0$
1083 and $\lambda \neq 1$ (as in most cases in our analyses, see Table 1 in the main text), then in general $r_{xz} =$
1084 $r_{xy} \times r_{yz} \neq r_{yx} \times r_{yz} = r_{xz}$. We would therefore prefer to avoid an arbitrary decision about the
1085 directionality of the model to affect the estimate of the coefficient.

1086 In our analyses we overcame the above difficulties by avoiding the use of bivariate
1087 PGLS models to estimate correlations between the variables in the path models. We followed
1088 the approach proposed by Santos (2012), in which first we calculated λ for each individual
1089 variable and conducted a phylogenetic transformation on the variable using that estimate of λ .
1090 Then we fitted path models to these already transformed data using ordinary fitting methods
1091 developed for fitting path models to non-phylogenetic data. Details of the analyses are
1092 described in the Methods section of the main text.

1093

1094 **2. The problem of comparing non-nested path models by AIC statistics**

1095 In phylogenetic comparative studies the direction of causality between variables is often
1096 unknown, and different evolutionary hypotheses may propose opposing cause - effect
1097 relationships (like the mating competition and the mortality cost hypotheses in our study, see
1098 Fig. S1). These hypotheses may be represented by different path models, and then their fit to
1099 the data can be compared by some comparative fit indices, most commonly by AIC (West et
1100 al. 2012). However, simulations suggest that conclusions of path model comparisons based on
1101 information theory approach (like AIC) can be unreliable (Preacher and Merkle 2012). In
1102 addition the competing models can be non-nested (non-hierarchical) (e.g. Models 1a versus 2a
1103 in Fig. S1), for which AIC-based comparison should be applied with caution (Kline 2015).

1104 To explore the problem of model comparison in the context of our study, first we
1105 fitted our path models to the real dataset by two alternative methods: (1) by covariance matrix

1106 comparison, as implemented in the R package *lavaan* (Rosseell 2012), and (2) by piecewise
 1107 structural equation modelling (or d-separation) method, as implemented in the *piecewiseSEM*
 1108 (Lefcheck 2016) package. We compared path coefficient estimates and various model fit
 1109 indices between these two methods to evaluate whether they produce consistent conclusions.
 1110 Second, we used the same two methods and R implementations to fit the models to simulated
 1111 datasets, and tested which of the methods produces more reliable (less biased) model
 1112 comparisons.

1113

1114 2.1. Fitting path models to real data

1115 The general steps of model fitting procedure we followed in this study are described in the
 1116 Methods section of the main text. We performed model fitting with the two R packages
 1117 *piecewiseSEM* and *lavaan*. In *piecewiseSEM* and *lavaan* the global model fit for each
 1118 individual path model is evaluated by Fisher's *C* and χ^2 statistics, respectively, where a
 1119 statistically non-significant result means acceptable fit. In *lavaan*, several other measures for
 1120 model fit of individual models are also available, and here we report four of the most widely
 1121 used indices (TLI, CFI, RMSEA, SRMR). It has been proposed that that the values of TLI
 1122 and CFI > 0.95, RMSEA < 0.06, and SRMR < 0.08 indicate acceptable/good fit of models to
 1123 the data (West et al. 2012).

1124 We found that the two methods produced highly consistent estimates for the
 1125 standardized path coefficients in all path models (*piecewiseSEM*: Table 1 in the main text,
 1126 *lavaan*: Table S9 below). The effect of juvenile mortality on ASR was marginally not
 1127 significant in most *piecewiseSEM* models whereas it was significant with all *lavaan* models.
 1128 For all other relationships the two methods produced consistent results.

1129

1130 **Table S9.** Estimates of standardized path coefficients for the six path models representing
 1131 various relationships between SSD, ASR, and sex biases in adult (AMB) and juvenile (JMB)
 1132 mortality, obtained by the R package *lavaan* (see Fig. S1 for model details). Significant
 1133 relationships are highlighted in bold.

1134

| Model/Path | Path coefficient ± SE | Z | P |
|-----------------|--------------------------|----------------|--------------|
| <i>Model 1a</i> | | | |
| AMB → ASR | - 0.340 ± 0.112 | - 3.048 | 0.002 |
| JMB → ASR | - 0.205 ± 0.102 | - 2.002 | 0.045 |
| ASR → SSD | - 0.657 ± 0.107 | - 6.144 | 0.000 |
| | | | |
| <i>Model 1b</i> | | | |

| | | | |
|--------------------------|------------------------|----------------|--------------|
| (AMB → ASR) ¹ | 0 | - | - |
| JMB → ASR | - 0.258 ± 0.105 | - 2.443 | 0.015 |
| ASR → SSD | - 0.657 ± 0.107 | - 6.144 | 0.000 |
| | | | |
| <i>Model 1c</i> | | | |
| AMB → ASR | - 0.378 ± 0.112 | - 3.370 | 0.001 |
| (JMB → ASR) ¹ | 0 | - | - |
| ASR → SSD | - 0.657 ± 0.107 | - 6.144 | 0.000 |
| | | | |
| <i>Model 2a</i> | | | |
| SSD → AMB | 0.117 ± 0.070 | 1.680 | 0.093 |
| SSD → JMB | 0.089 ± 0.077 | 1.157 | 0.247 |
| AMB → ASR | - 0.340 ± 0.110 | - 3.092 | 0.002 |
| JMB → ASR | - 0.205 ± 0.101 | - 2.031 | 0.042 |
| | | | |
| <i>Model 2b</i> | | | |
| SSD → JMB | 0.089 ± 0.077 | 1.157 | 0.247 |
| AMB → ASR | - 0.340 ± 0.110 | - 3.092 | 0.002 |
| JMB → ASR | - 0.205 ± 0.101 | - 2.031 | 0.042 |
| | | | |
| <i>Model 2c</i> | | | |
| SSD → AMB | 0.117 ± 0.070 | 1.680 | 0.093 |
| AMB → ASR | - 0.340 ± 0.110 | - 3.092 | 0.002 |
| JMB → ASR | - 0.205 ± 0.101 | - 2.031 | 0.042 |

1135

1136 ¹ Path coefficient set to zero

1137

1138

1139

1140

1141 The two methods also produced highly consistent results for model fit as evaluated by global

1142 fit indices (i.e. *C* and χ^2 statistics, respectively, see Table S10). The only difference was that

1143 for Model 1b *piecewiseSEM* indicated 'marginally acceptable' model fit whereas *lavaan*

1144 indicated poor model fit for this path model. The other fit indices (TLI, CFI, RMSEA, and

1145 SRMR) suggest conclusions that are fully consistent with *C* statistics and χ^2 tests, i.e.

1146 acceptable fit for Models 1a and 1c by all of these indices and unacceptable fit for all other

1147 models (Table S10).

1148

1149

1150 **Table S10.** Fit indices for the six path models, obtained by *piecewiseSEM* and *lavaan*. Values

1151 indicating acceptable fit are highlighted in bold.

1152

| Model | piecewiseSEM | | | lavaan | | | | | | |
|-------|--------------|-----------|----------------------|-------------|-----------|----------------------------------------|--------------|--------------|--------------|--------------|
| | <i>C</i> | <i>df</i> | <i>P_c</i> | χ^2 | <i>df</i> | <i>P_{χ^2}</i> | TLI | CFI | RMSEA | SRMR |
| 1a | 0.29 | 4 | 0.972 | 0.02 | 2 | 0.991 | 1.119 | 1.000 | 0.000 | 0.004 |
| 1b | 11.6 | 6 | 0.065 | 8.9 | 3 | 0.031 | 0.764 | 0.858 | 0.143 | 0.101 |
| 1c | 6.2 | 6 | 0.376 | 3.9 | 3 | 0.267 | 0.962 | 0.977 | 0.057 | 0.065 |
| 2a | 34.7 | 4 | 0.000 | 30.6 | 2 | 0.000 | -0.978 | 0.341 | 0.386 | 0.154 |
| 2b | 34.7 | 4 | 0.000 | 30.6 | 2 | 0.000 | -0.719 | 0.313 | 0.386 | 0.154 |
| 2c | 34.7 | 4 | 0.000 | 30.6 | 2 | 0.000 | -0.661 | 0.336 | 0.386 | 0.154 |

1153

1154

1155 2.2. AIC-based model comparisons using real and simulated data

1156 To assess which of these models provides the best account of the data, first we calculated the
1157 AIC value for each model (in *piecewiseSEM* this is corrected for small sample size, i.e. AIC_c)
1158 using the real dataset. Second, we used simulated data to test which of the two methods
1159 produces less biased conclusions. For this latter purpose, we generated simulated datasets
1160 using the R function ‘rnorm’. The simulated datasets have the same number of variables and
1161 sample size as the phylogenetically transformed real dataset. We fitted path models with both
1162 *piecewiseSEM* and *lavaan* to obtain the AIC (or AIC_c) values. Then we compared Model 1a
1163 (the model that got the highest support for model fit by the global fit indices, see Table S10)
1164 to the other five models (Models 1b, 1c, 2a, 2b, and 2c), thus conducted five pairwise
1165 comparisons, repeated with the two methods. These paired comparisons between models
1166 mimic the comparison we conducted with the real dataset in our study (Table 2 in the main
1167 text). We calculated Δ AIC for each comparison as the difference between AIC values of the
1168 two models (i.e. AIC of compared model - AIC of Model 1a, thus a positive Δ AIC value
1169 indicates better fit for Model 1a). We repeated this procedure with 1000 simulated datasets
1170 that resulted in 1000 Δ AIC values for each pairwise comparison. To assess whether the
1171 comparison of two particular models produces biased results with simulated data we
1172 calculated (1) the mean Δ AIC value of the 1000 runs (Δ AIC_{simulation}), and (2) the probability
1173 that the simulated Δ AIC was larger than the Δ AIC value we got with the real dataset
1174 ($P_{\geq \Delta \text{AIC}_{\text{sim}}}$).

1175 Using real data, *piecewiseSEM* gave the lowest AIC_c for Model 1a (Table S11), a
1176 result consistent with global model fit evaluation (see Table S10). Δ AIC_c values suggested

1177 strong support for this model in all comparisons ($\Delta AIC_c \geq 4.1$, Table S11). In contrast,
 1178 *lavaan* results were inconsistent with global model fit evaluation because it gave very strong
 1179 support for Model 2c (Table S11), a model that had an unacceptable fit by all fit indices (see
 1180 Table S10).

1181

1182 **Table S11.** AIC-based model comparison using real and simulated data by the two methods.
 1183 AIC_c (*piecewiseSEM*) and AIC (*lavaan*) values provided for all models are based on analyses
 1184 of our real data. ΔAIC_{data} and $\Delta AIC_{simulation}$ show differences from Model 1a in pairwise
 1185 comparisons, based on analyses of real or simulated data, respectively. $P_{\geq \Delta AIC_{sim}}$ indicates the
 1186 probability that analyses of random data result in as large or larger AIC differences in support
 1187 for Model 1a than the ΔAIC values obtained with real data.

1188

| Model | piecewiseSEM | | | | lavaan | | | |
|-------|------------------|---------------------|---------------------------|-----------------------------|--------------|---------------------|---------------------------|-----------------------------|
| | AIC _c | ΔAIC_{data} | $\Delta AIC_{simulation}$ | $P_{\geq \Delta AIC_{sim}}$ | AIC | ΔAIC_{data} | $\Delta AIC_{simulation}$ | $P_{\geq \Delta AIC_{sim}}$ |
| 1a | 15.8 | 0.0 | - | - | 382.4 | 0.0 | - | - |
| 1b | 25.7 | 9.9 | -0.2 | 0.003 | 389.3 | 6.9 | -1.0 | 0.004 |
| 1c | 19.9 | 4.1 | -0.9 | 0.042 | 384.3 | 1.9 | -1.1 | 0.041 |
| 2a | 59.0 | 43.2 | 7.4 | 0.0 | 521.3 | 138.9 | 274.3 | 1.0 |
| 2b | 50.4 | 34.6 | -0.1 | 0.0 | 360.0 | -22.4 | 273.4 | 1.0 |
| 2c | 50.4 | 34.6 | 0.4 | 0.0 | 341.7 | -40.7 | 273.4 | 1.0 |

1189

1190

1191 Using simulated data, we found that *piecewiseSEM* produced less biased results than *lavaan*.
 1192 First, in most cases mean simulated ΔAIC values were small and there was no strong bias in
 1193 favor of one specific model (see $\Delta AIC_{simulation}$ in Table S11), as one would expect with
 1194 random data. The only exception was the comparison between Model 1a and Model 2a in
 1195 which simulated ΔAIC produced by *piecewiseSEM* was 7.4, favoring Model 1a. Importantly,
 1196 however, these simulations indicated only a low probability for random data resulting in as
 1197 large or larger AIC differences (43.2) in support for Model 1a than the ΔAIC values we
 1198 obtained with real data (see low $P_{\geq \Delta AIC_{sim}}$ values in Table S11), suggesting that support for
 1199 Model 1a was unlikely the result of biased AIC estimates.

1200

1201

1202

In contrast, simulations showed that *lavaan* produced highly biased ΔAIC values in all
 non-nested comparisons (see the high $\Delta AIC_{simulation}$ and $P_{\geq \Delta AIC_{sim}}$ values for Models 2a, 2b
 and 2c in Table S9). On the other hand, for nested model comparisons (i.e. with Models 1b

1203 and 1c) *lavaan* produced unbiased results similarly to those we got with *piecewiseSEM*
1204 (Table S11).

1205 These analyses suggest that the two methods gave consistent results for (1) path
1206 coefficients estimates and for (2) evaluating model fit of individual path models by global fit
1207 indices (using *C* statistics in *piecewiseSEM*, and χ^2 , TLI, CFI, RMSEA, and SRMR in
1208 *lavaan*). On the other hand, simulation results indicate that AIC-based model comparisons are
1209 less biased when performed by the piecewise structural equation modelling method, at least
1210 for comparisons between non-nested models.

1211

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