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**TITLE: The complex effects of mass extinctions on morphological  
disparity**

Short title: Mass extinctions and morphological disparity

**Mark N Puttick, Thomas Guillaume, Matthew A Wills**

1 **ABSTRACT**

2 Studies of biodiversity through deep time have been a staple for biologists and  
3 palaeontologists for over 60 years. Investigations of species richness (diversity) revealed that  
4 at least five mass extinctions punctuated the last half billion years, each seeing the rapid  
5 demise of a large proportion of contemporary taxa. In contrast to diversity, the response of  
6 morphological diversity (disparity) to mass extinctions is unclear. Generally, diversity and  
7 disparity are decoupled, such that diversity may decline as morphological disparity increases,  
8 and vice versa. Here, we develop simulations to model disparity changes across mass  
9 extinctions using continuous traits and birth-death trees. We find no simple null for disparity  
10 change following a mass extinction but do observe general patterns. The range of trait values  
11 decreases following either random or trait-selective mass extinctions, whereas variance and  
12 the density of morphospace occupation only decline following trait-selective events. General  
13 trends may differentiate random and trait-selective mass extinctions, but methods struggle to  
14 identify trait selectivity. Long-term effects of mass extinction trait selectivity change support  
15 for phylogenetic comparative methods away from the simulated Brownian motion towards  
16 Ornstein-Uhlenbeck and Early Burst models. We find that morphological change over mass  
17 extinction is best studied by quantifying multiple aspects of morphospace occupation.

18 **Keywords:** disparity, mass extinctions, phylogenetic comparative methods, traits,  
19 macroevolution

## 20 INTRODUCTION

21 Evolutionary biologists and paleobiologists have long quantified diversity in terms of  
22 species numbers or species richness, making comparisons both horizontally between clades  
23 and higher taxa (Wiens 2017) and vertically throughout evolutionary time (Smith 2007).  
24 However, diversity takes no account of the morphological differences between species, a  
25 property known as morphological disparity (Wills et al. 1994). Researchers have attempted to  
26 formally define disparity in different ways, but most of the indices that derive from these  
27 definitions quantify variation in morphology or phenotype (Wills 2001; Hopkins and Gerber  
28 2017; Guillerme et al. 2020a).

29 There have been numerous empirical analyses of the diversity of the global biota  
30 through evolutionary time (Raup and Sepkoski 1982; Alroy 2010) with an overall null  
31 expectation of symmetry as clades rise and fall in diversity (Gilinsky and Good 1989; Liow et  
32 al. 2010). For disparity, by contrast, there is no such simple null (Pie and Weitz 2005;  
33 Hughes et al. 2013; Oyston et al. 2015, 2016) but see (Foote 1996). Quantifying disparity  
34 alongside diversity is essential to fully understand the evolution of biodiversity (Roy and  
35 Foote 1997) as a series of empirical studies have demonstrated that diversity and disparity are  
36 largely decoupled (Wills et al. 1994; Fortey et al. 1996; Bapst et al. 2012). Analyses of  
37 disparity have accordingly proven invaluable for studying the tempo and mode of evolution  
38 (Simpson 1944), how clades diversify through time and throughout morphological “form  
39 space” (Gould 1990), and the patterns of drift and selection that have produced the  
40 distribution of living diversity (Raup 1981).

41 Extinction has had an immediate and potentially catastrophic role in sculpting patterns  
42 of biodiversity through time (Sepkoski 1981), particularly mass extinctions (Bond and  
43 Grasby 2017). Mass extinctions are defined, by convention, as geologically brief events that  
44 remove at least 75% of contemporary diversity (Barnosky et al. 2011). Researchers accept

45 that the effect of a mass extinction upon disparity will depend upon whether species are  
46 selectivity or randomly removed by extinction (Foote 1997; Korn et al. 2013). Discrete  
47 events in which extinction acts selectively are expected to decrease disparity by eliminating  
48 the majority of an enclosed area of morphospace. Non-selective events, by contrast, may not  
49 lead to disparity decreases (Foote 1991, 1993; Roy and Foote 1997; Villier and Korn 2004;  
50 Korn et al. 2013), particularly when disparity is measured using a variance-based index (Korn  
51 et al. 2013).

52 Empirical studies have demonstrated cases where mass extinctions acted both  
53 selectively and non-selectively with regards to particular traits (Foote 1993; Roy 1996;  
54 Lockwood 2004; Erwin 2007; Halliday and Goswami 2016). Phylogenetic comparative  
55 studies indicate that there is trait selectivity on some groups, such as vascular plants (Green et  
56 al. 2011), but there is little evidence to link traits and extinction susceptibility during mass  
57 crises in the fossil record (Friedman 2009; Puttick et al. 2017; Allen et al. 2019). Many  
58 studies reveal a phylogenetic signal of extinction (Hardy et al. 2012; Harnik et al. 2014; Krug  
59 and Patzkowsky 2015; Puttick et al. 2017; Soul and Friedman 2017), without demonstrating  
60 links to trait selectivity. We note that in studies of fossil record disparity, it is only possible to  
61 analyze traits with fossilization potential, which excludes soft body parts and behavior.

62 In order to investigate the consequences of extinctions on traits, one approach would  
63 be to analyze empirical data, as simulations may lack biological realism. Importantly,  
64 however, simulations provide an underpinning framework for such empirical analyses (Foote  
65 1991, 1997; Ciampaglio et al. 2001; Bapst et al. 2012; Korn et al. 2013), as it is possible to  
66 test scenarios in which we know the definitive underlying patterns; an impossibility with  
67 empirical data. Here we use a novel simulation approach to determine the expected, null  
68 patterns of disparity change through mass extinction events and to investigate how quickly  
69 diversity and disparity might be expected to recover to pre-extinction levels under many

70 scenarios. We simulate birth-death trees (Ma 2010; Stadler 2010; Mooers et al. 2012) and  
71 traits under Brownian motion with a variety of parameter values. During our simulations, a  
72 mass extinction removes a proportion of contemporary diversity. We show that disparity  
73 generally reduces following a mass extinction event, but that all patterns are variable.  
74 Alternative disparity indices imply different patterns of disparity change, such that there is a  
75 fundamental link between how we conceptualize and quantify disparity (Wills 1998b; Korn  
76 et al. 2013). Finally, we assess how mass extinctions bias comparative phylogenetic models  
77 of trait evolution in the extant lineages that survive such events. Phylogenetic regressions do  
78 not distinguish selective and non-selective mass extinction events under the simplest  
79 simulation models and the selective removal of taxa according to trait value causes  
80 comparative models to support non-BM models in preference to the (true) simulated BM  
81 process.

82

## 83 **METHODS**

### 84 *Overview*

85 We summarize the full simulation procedure in Fig 1. We simulated birth-death trees  
86 and Brownian motion traits simultaneously. We ran simulations until 50, 100, or 200  
87 contemporary lineages were present, after which an extinction event removed 50, 75, or 90%  
88 of standing diversity. Simulations finished when diversity had recovered to pre-extinction  
89 levels (i.e., 50, 100, 200 tips). We chose the severity of mass extinctions (50, 75, 90%) to  
90 reflect known extinction values in the fossil record (Stanley 2016; Bond and Grasby 2017).  
91 The upper figure (90%) reflects the severity often quoted for the end-Permian event, although  
92 this has been questioned (Stanley 2016). The middle figure (75% of species lost) reflects the  
93 severest estimates for end-Ordovician and end-Cretaceous events and is accepted as the  
94 necessary threshold for a mass extinction (Barnosky et al. 2011). We used 50% for a

95 ‘smaller’ extinction events, such as the Frasnian extinction in the Late Devonian (Stanley  
96 2016; Bond and Grasby 2017).

97 We simulated extinction in two different ways. Taxa were either eliminated at random  
98 in a non-selective event or removed according to their trait values to simulate a selective  
99 extinction. In separate analyses, we ran models to simulate one, two, and five traits. We then  
100 assessed disparity through time and analyzed the ability of models to recover the signal of  
101 trait change across extinctions. We wrote new code in *R* to simulate data; users can freely  
102 download this as part of the package *Phylogenetic Evolution of Traits and Extinction in R*  
103 (PETER) from *GitHub* (<https://github.com/PuttickMacroevolution/PETER>).

104 The first objective of this study is to provide a simple null model for change in  
105 disparity across mass extinctions, given a number of varying parameters. In our simulations,  
106 we have an extremely high resolution of extinction through time as we record the trait values  
107 for all simulated internal nodes and tips. In the fossil record, however, it is often necessary to  
108 aggregate data into time bins. Therefore, we also undertook steps to study binned values of  
109 disparity through time. For the majority of our analyses, we focused on studies with one  
110 continuous trait, as there is often just a single measurement available for a large group of  
111 species (e.g., body size in fossil vertebrates). For these simulations, we attempted to control  
112 for properties that are likely to vary across the tree of life, such as rates of background  
113 extinction and the total number of species. Our multivariate approach was designed to extend  
114 the single trait approach, albeit that selectivity was simulated as acting on a single trait.  
115 Evolution in this directly selected focal trait was correlated with evolution in the other  
116 trait(s), so that these latter traits experienced indirect selectivity (Lande and Arnold 1983).  
117 We also ran simulations in which traits evolved independently.

118 The second objective is to examine the effectiveness of methods commonly used to  
119 investigate mass extinctions. For example, we determine whether it is possible to differentiate

120 trait-selective and trait-independent mass extinctions, and how mass extinctions influence  
121 models of continuous trait evolution through time.

### 122 ***Simulation of trees and traits***

123 We simulated trees evolving from the root to the tips with time-homogeneous  
124 speciation ( $\lambda$ ) and extinction rates ( $\mu$ ). We used the same speciation rate ( $\lambda = 1$ ) for all  
125 simulations, but different levels of background extinction ( $\mu = 0, 0.4, 0.8$ ).

126 We simulated traits under a Brownian motion process (Felsenstein 1973, 1985). At  
127 the start of each simulation, we sampled the root state from a normal distribution with a mean  
128 of zero and a standard deviation of one. In simulations with two and five traits, we set the co-  
129 variance either to zero such that the traits were independent or to a co-variance of 0.75 so that  
130 the evolution of traits was linked but not identical. The use of a moderate co-variance (0.75)  
131 implies that although selectivity (when present) would act on one trait only, the mass  
132 extinction would also affect the remaining, non-directly selected traits. We designed this to  
133 mimic a scenario in which selectivity acts on one analyzed trait that is known to co-vary with  
134 many other traits, such as body size.

### 135 ***Disparity indices***

136 We summarized diversity using lineage-through-time plots, and we used three indices  
137 of disparity (Fig 1C): *median pairwise distance*, which is the median pairwise Euclidean  
138 distance between all points (Wills et al. 1994); *Sum of Variances* (SOV) from each trait  
139 (Foote 1992a); and *Sum of Ranges* (SOR) (Wills 1998a, 2001; Brusatte et al. 2011; Ruta et al.  
140 2013). We use these disparity indices to elucidate different aspects of morphospace  
141 occupation: we interpret *median pairwise distance* as an index of the density of morphospace  
142 occupation; and we consider *Sum Of Variances* as an index of dispersion, and *Sum Of Ranges*  
143 as an index of the overall magnitude or volume of morphospace occupation (Guillerme et al.



144 2020b). Unless stated, all analyses were conducted using the R package *dispRity* (Guillerme  
145 2018).

146 Our simulations produced trait estimates for every node in the tree. We estimated  
147 disparity indices using these traits, but for most analyses, we binned trait values into equal-  
148 size time bins and subsequently calculated disparity for each bin using a time-slicing  
149 approach. We explored the impacts of using different bin sizes (4, 8, 10, 16-time bins), and  
150 used the highest resolution, sixteen-time bins to assess if there were significant differences in  
151 using each bin size. We used sixteen time-bins throughout the remaining analyses, but we  
152 observed the same general trends with all bin sizes (Table 1). After each simulation, we  
153 scaled disparity and diversity to unity by dividing each by their maximum values.

#### 154 **Simulation of mass extinctions**

155 For a given quantile of extinction, we set different strengths of selectivity:

156 (i) *Strict selectivity*. All lineages with traits larger than the cut-off went extinct and all  
157 other lineages survived.

158 (ii) *Strong selectivity*. To investigate events in which the vast majority of extinction  
159 related to the size of traits, lineages larger than the cut off had a probability of  
160 0.99 to go extinct whilst lineages smaller than the cut off had a probability of 0.01  
161 to go extinct.

162 (iii) *Random selectivity*. Trait values had no influence on extinction susceptibility.

163 (iv) *Weak selectivity*. An intermediate scenario in which lineages larger than the cut off  
164 had a probability of 0.75 of extinction whilst lineages smaller than the cut off had  
165 an extinction probability of 0.25 (i.e, there was a 25% chance of random  
166 extinction).

167 In each case of selectivity strength, we set the cut-off by using the contemporary  
168 distribution of traits at the extinction boundary; lineages with trait values above the 0.5, 0.25,

169 or 0.1 quantiles of the contemporary distribution were more prone to extinction in those  
170 simulations with *directional selectivity*. The quantiles were chosen to directly reflect the  
171 selectivity of extinction. So, for simulations with *strict selectivity* there was a zero probability  
172 of extinction for taxa with trait values below a certain cut-off value. As an example, for  
173 simulations with 75% loss of taxa at an extinction boundary and *strict selectivity*, there was a  
174 zero probability of extinction for taxa with trait values below the 0.25 quantile of all  
175 contemporary trait values.

176 We binned and summarized simulated data by pooling all parameters:  $n$  lineages lost,  
177 background extinction, and the number of tips for *strict*, *strong*, and *random* extinctions. We  
178 used three levels of background extinction in all other simulations (i.e., 0, 0.4, 0.8). In  
179 analyses with one trait only, we also ran simulations with a background extinction rate of 1.

180 Finally, we ran *disruptive selectivity* mass extinction simulations for a single trait in  
181 which lineages with trait values closer to the mean were prone to extinction, such that  
182 lineages with smaller or larger trait values were more likely to survive. For these simulations,  
183 we set the selective extinctions to remove lineages from the 0.5, 0.75, and 0.9 quantiles  
184 symmetrical around the mean.

### 185 **Phylogenetic analysis of mass extinctions**

186 We tested whether survivors and extinct lineages differed significantly in their trait  
187 values using Ordinary Least Squares (OLS) and Phylogenetic Generalized Least Squares  
188 (pGLS) regressions (Grafen 1989). In these models, we set the response as the continuous  
189 trait value and coded the predictor as a binary variable indicating whether or not the lineage  
190 survived the mass extinction. We pruned the phylogeny and trait values to include only those  
191 lineages that existed at the boundary (Puttick et al. 2017), such that we did not consider  
192 values from lineages that were lost before the extinction or arose after it. We estimated the

193 phylogenetic signal in trait values using Pagel's lambda (Pagel 1997, 1999) alongside the  
194 regression parameters in the R package *motmot* (Puttick et al. 2020).

195 We tested the phylogenetic signal in extinction itself, without consideration of trait  
196 values, using the *phyloD* statistic (Fritz and Purvis 2010). This method used phylogenetic  
197 contrasts to estimate internal node values of binary traits (zero = survivors, one = extinct  
198 lineages), and used these node values to estimate the number of binary transitions. The  
199 number of changes indicated the phylogenetic signal, with the number of changes and signal  
200 strength being inversely correlated (more changes means a weaker phylogenetic signal). A  
201 Brownian motion model will produce a *d* value close to 0 under the *phyloD* statistic. Values  
202 larger than 0 denote increasingly random signal, and values under 0 are indicative of over-  
203 conserved signals.

## 204 **Phylogenetic comparative models of trait distributions that have recovered from a mass** 205 **extinction**

206 The signal of past mass extinctions may be present in the trait distribution of extant  
207 lineages. To explore these patterns, we pruned simulated datasets to yield ultrametric trees  
208 comprising only extant taxa (i.e., lineages in existence at the end of the simulation). We  
209 applied commonly-used likelihood phylogenetic comparative models of trait evolution to  
210 these pruned data: Brownian motion (Felsenstein 1973, 1985) to represent the null model;  
211 evolution under a constraint using an Ornstein-Uhlenbeck (OU) model (Hansen 1997; Butler  
212 and King 2004; Blomberg et al. 2020) in which variance was constrained to the ancestral  
213 value according to the strength of  $\alpha$ ; and the Early Burst model, in which the rate variance  
214 exponentially decreased through time (Blomberg et al. 2003; Harmon et al. 2010).

215 The whole procedure for running these simulations is available and documented on  
216 *GitHub* (<https://github.com/PuttickMacroevolution/PETER>).

## 217 **RESULTS**

218 ***The variance around simulated disparity is higher than the variance around diversity***

219 We ran initial analyses without a mass extinction event. When plotted through time,  
220 relative disparity showed an overall pattern of increase (Fig 2). Trends are similar for each  
221 index, but there is a more substantial difference between disparity and diversity when the  
222 former is measured using the Sum Of Ranges rather than the Sum of Variances or median  
223 pairwise distance. We observed similar trends in simulations with two and five traits  
224 (Supplementary Fig S1).

225 ***Mass extinctions usually cause reductions in disparity, and the precise patterns are related***  
226 ***to the index of disparity***

227 As a general rule, selective mass extinctions caused a reduction in disparity (Fig 3,  
228 Supplementary Figure S2). However, this pattern was not universal, and changed according  
229 to the simulation model (e.g., different strengths of selectivity) and the disparity index used  
230 (Fig 4).

231 All disparity indices showed iterations with both increases and decreases across a  
232 mass extinction, such that there is no universal pattern. Pooling the results of all simulations  
233 that entailed strict directional selectivity (irrespective of background extinction, number of  
234 tips, or extinction severity) we found a median decrease in disparity indexed by median  
235 pairwise distance (90%), Sum of Variances (90%), and Sum of Ranges (75%). When  
236 selectivity was absent from the simulations, all indices showed a median disparity change of  
237 zero across the mass extinction, except for the Sum Of Ranges for which most simulations  
238 showed a decrease in disparity across the boundary (75%). For those simulations with one  
239 trait, we ran additional simulations that used the special case of  $\lambda = \mu = 1$ , and these showed  
240 patterns similar to those in simulations with other levels of extinction (Supplementary Table  
241 S1, Supplementary Figure S2).

242 For a single trait, we also investigated the effects of varying the simulation parameters  
243 individually (viz., strength of selectivity, the proportion of lineages lost, rate of background  
244 extinction, number of tips, and disparity index used) (Supplementary Fig S3). Simulations  
245 with more tips showed more extreme reductions in disparity across the mass extinction  
246 boundary, irrespective of the disparity index used. Only in certain circumstances did all  
247 iterations indicate a decrease in disparity across an extinction boundary. For example, there  
248 was a consistent decrease in disparity across a mass extinction in simulations with more tips,  
249 more severe lineage loss, and higher rates of background extinction (including a background  
250 extinction of  $\mu = 1$ ). The Sum Of Ranges index is very sensitive to sample size differences,  
251 and a fall in the Sum of Ranges would be expected alongside a reduction in diversity in the  
252 wake of a mass extinction. For the *directional selectivity* simulations with one trait, we ran  
253 additional simulations that used the special case of  $\lambda = \mu = 1$  and these showed similar  
254 patterns to simulations with other levels of extinction (Supplementary Table S1).

255 For simulations with an intermediate level of extinction selectivity (*weak selectivity*  
256 with 25% probability of random extinction), the results are generally closer to *random*  
257 *selectivity* models rather than to other models with directional selectivity (Supplementary  
258 Table S1).

259 In simulations with multiple traits, only the extinctions with strict selectivity showed a  
260 consistent decrease in disparity across a mass extinction boundary (Fig 4B-C). The median  
261 decreases in disparity for multiple traits were smaller than in simulations with one trait, and  
262 some multiple trait simulations indicated an increase in disparity across the extinction event.  
263 The number of iterations that showed a decrease in disparity across the mass extinction  
264 boundary was lower in simulations with no trait co-variance compared to analyses with 0.75  
265 trait co-variance (Table 1A). Again, an exception was the Sum Of Ranges, in which the  
266 majority of iterations showed a decrease in disparity with any selectivity strength, and nearly

267 all traits had a decrease in disparity with five traits (Table 1B). We note that increases in  
268 range-based indices are only possible because of time binning effects: the *immediate* wake of  
269 an extinction event (if not time averaged and therefore excluding new lineages) could only  
270 show a decrease or no change in range. However, we seek to simulate the temporal sampling  
271 that pertains in typical paleontological data sets. The tendency for a disruptive extinction was  
272 for all disparity indices to show an increase or no change across the mass extinction boundary  
273 (Fig S4).

#### 274 ***Comparative phylogenetic methods do not distinguish selective and non-selective*** 275 ***extinctions***

276 pGLS models were more conservative at detecting the true relationship between trait  
277 values and extinction selectivity compared to OLS models, but pGLS models have lower  
278 type-two error rates in comparison to OLS models (Fig 5). In all simulations with *strict* and  
279 *strong* selectivity 99% of OLS models support the correct relationship, compared to only a  
280 median of 86% (*strict*) and 68% (*strong*) of pGLS models. With *weak selectivity*, few pGLS  
281 (median 6%) models show a significant relationship between extinction and traits, which is  
282 much lower than the OLS relationship (80%). When there is no relationship between trait  
283 values and extinction selectivity, pGLS models have a lower error rate (median 1%)  
284 compared to pGLS models (median 6%).

#### 285 ***Models consistently detect the phylogenetic signal of extinction***

286 There was a strong phylogenetic signal, as measured by the *phyloD* statistic, when  
287 selectivity was *strict* (median 0.07) and *strong* (median 0.12). Conversely, there was a  
288 random signal when selectivity was *weak* (median 0.82) or *random* (median 1.0). Only the  
289 strength of selectivity and magnitude of lineage loss were significant factors in explaining the  
290 differences in phylogenetic signal. There was no significant difference in the phylogenetic  
291 signal for different numbers of tips or different levels of background extinction.

292 ***Mass extinctions have long-term impacts on trait distributions***

293 High levels of extinction selectivity increased support for non-Brownian models when  
294 analyzed using exclusively extant data that had ancestrally suffered a mass extinction (Fig 6,  
295 Supplementary Fig S5). For a single trait with no mass extinction, most simulations  
296 supported a Brownian motion model (median 82%), with relatively few supporting Early  
297 Burst (3%) and OU (15%) models.

298 For simulations of *directional selectivity*, strong or strict selectivity yielded much  
299 lower support for Brownian motion (median strict 51%, strong 70%), and higher support for  
300 the OU model (strict 49%, strong 28%). When there was no selectivity on the trait value at  
301 the mass extinction boundary, the relative support for models was comparable to support for  
302 models in simulations with no mass extinctions (BM support, no selectivity 80%). Across all  
303 analyses that supported the OU model, the strength of selectivity  $\alpha$  had a median half-life of  
304 1.69 (within a range of values from 0.52 to 17.25). Similar patterns were seen in simulations  
305 with two traits compared to simulations with one trait.

306 With higher levels of background selectivity in the simulations with *disruptive*  
307 *selectivity* at mass extinctions, there was lower support for the OU model compared to BM.  
308 This was because there was a high turnover of lineages such that the ‘crown’ of these  
309 simulated phylogenies emerged after the mass extinction. When only those trees that had at  
310 least one ‘crown’ node *predating* the mass extinction were analyzed, the support for OU  
311 model was higher (Fig S6). *Disruptive selectivity*, in contrast to directional trait selectivity,  
312 resulted in higher support for an early burst model compared with the simulated Brownian  
313 motion process.

314 **DISCUSSION**

315 Few putative macroevolutionary rules withstand scrutiny (McShea 1998; Hone and  
316 Benton 2005; McShea and Brandon 2010; Benson et al. 2018). However, there do appear to

317 be statistical generalities concerning the manner in which major clades evolve through time,  
318 with most groups achieving maximum or near maximal morphological disparity relatively  
319 early in their existence (Foote 1992a, 1994, 1996; Hughes et al. 2013; Oyston et al. 2015,  
320 2016). The largest environmental crises often appear to disrupt this pattern, with those clades  
321 that go extinct coincident with a mass event typically being truncated and having maximum  
322 disparity much later in their evolutionary trajectories (Hughes et al. 2013). Despite these  
323 empirical observations, there are no null models for the manner in which we expect the  
324 disparity of clades to change through time, still less for clades truncated by or (as here)  
325 surviving such events with reduced diversity (Korn et al. 2013). Hence, simulations provide a  
326 powerful way of analyzing the role of mass extinctions in shaping disparity.

327         We acknowledge that there is no single, universally agreed index for disparity.  
328 Moreover, all empirical assessments of disparity are necessarily relative and constrained  
329 within the context of a particular set of descriptors or data set. We also note that disparity is  
330 indexed with reference to the constituent entities (species or other operational taxonomic  
331 units) within a group (e.g., subclade or time bin), with no reference to species or other entities  
332 outside of that group. As such, identical indices of disparity can be reported for clades  
333 occupying very different regions of morphospace, or for clades that migrate through that  
334 morphospace through time. Strikingly different distributions of points (local densities and  
335 clustering structures) can also yield identical disparity indices (Wills et al. 2012).  
336 Comparisons of diversity and disparity can nevertheless yield insights into the dynamics of  
337 evolutionary change (Foote 1991, 1993, 1994, 1997), while comparisons of a variety of  
338 disparity indices can be used to classify patterns of disparity change across mass extinction  
339 boundaries (Korn et al. 2013). There are many aspects of evolutionary dynamics (e.g., the  
340 directionality or otherwise of selectivity, modes of morphological evolution or speciation)  
341 that are more effectively tested and modelled directly rather than via the distributions of taxa



342 in morphospace.

343 By definition, a mass extinction destroys standing diversity, but our results suggest  
344 that it need not always precipitate a fall in disparity (Foote 1991). Across a selective  
345 extinction the variance and range of morphological occupation typically decreases (Fig. 3);  
346 but the variance is mostly unchanged across a non-selective extinction, even though the range  
347 of values still decreases. The patterns of disparity change that we observe depend upon the  
348 indices that we use to quantify it, so these indices must therefore be codifying different  
349 things. Only the simulations with little deviation from trait-based extinctions show general,  
350 predictable patterns of extinction through time. However, when there is more variance around  
351 mass extinction selectivity (*weak selectivity*), the observed patterns are much closer to those  
352 seen when extinction is random in regard to trait values.

353 Patterns of disparity are more complicated than patterns of diversity. Here, we  
354 principally discuss patterns of directionally selective mass extinctions, but we note that  
355 disruptive selectivity can yield disparity increases across extinction boundaries (Fig S4).  
356 Many parameters complicate the pattern of disparity change across mass extinction events,  
357 including the number of analyzed traits, co-variance between traits, and the index of disparity  
358 (Fig 3; Fig S3). For example, when the co-variation between multiple traits is high (0.75), the  
359 observed patterns of disparity through time (Fig 4B) are similar to those observed with just  
360 one trait (Fig 4A). Thus, there is no single expectation of disparity change across a mass  
361 extinction for all clades and all events. Rather, we suggest that in order to understand patterns  
362 of disparity change across a mass extinction for a particular group, it is necessary to  
363 determine whether this pattern differs from those expected given a similar set of parameters.  
364 These expected patterns could be determined using simulations (Foote 1991; Harmon et al.  
365 2003; Slater et al. 2010; Green et al. 2011; Korn et al. 2013). We do note, however, that even

366 with simulation approaches it could be impossible to detect small-scale changes in selectivity  
367 at a large scale (Raup et al. 1973), especially in analyses with hundreds or thousands of traits.

368         Where we find decreases in disparity across mass extinction events, these are often  
369 non-linear (Fig 3). Clades evolve through an empirically realized (Stone 1997) or  
370 theoretically possible (Novack-Gottshall 2007) multidimensional trait morphospace via a  
371 process of branching cladogenesis that is inherently diffusive and with increasing degrees of  
372 freedom (McShea 1998). By contrast, random processes of lineage extinction cause a linear  
373 decline in diversity, but disparity is a function of the *distribution* of a group of entities, be this  
374 morphological range, variance or otherwise (Foote and Gould 1992; Wills et al. 1994; Wills  
375 2001; Ruta et al. 2013; Bazzi et al. 2018). Hence after an extinction event, disparity is  
376 expected to decline non-linearly and more slowly than diversity (Foote 1991; Ruta et al.  
377 2013). This expectation is generally echoed in our simulations (Fig 3), but the precise pattern  
378 is contingent on the index of disparity used.

379         Korn et al. (2013) demonstrated that it is possible to differentiate types of mass  
380 extinction selectivity by examining changes in morphospace occupation. For example, a  
381 random extinction that is non-selective with regards to lineage trait values leads to a minor  
382 decrease in the Sum Of Ranges, with no appreciable change in disparity indexed using the  
383 Sum Of Variances. For median pairwise distance and Sum Of Variances indices, a substantial  
384 decrease in disparity may be indicative of a directionally selective mass extinction. In some  
385 instances, these indices may not reflect a decrease in disparity or may reflect a change in the  
386 pattern of morphospace occupation (Ciampaglio et al. 2001). The Sum Of Ranges decreases  
387 with all types of extinction selectivity and so does not differentiate between selective and  
388 random extinctions as well as mean pairwise distance and Sum Of Variances. Because the  
389 Sum Of Ranges is sensitive to sample size effects (Foote 1991, 1992b; Butler et al. 2012) it  
390 may be an unsuitable index to capture patterns of disparity across a mass extinction event.

391 However, some have argued this is unimportant when sample sizes are large (Simon et al.  
392 2010), and that it can be adjusted using rarefaction approaches (Foote 1992b; Wills 1998a).  
393 Median pairwise distances and Sum Of Variances may be able to differentiate selective and  
394 non-selective extinction, but no one index of disparity will be sufficient to distinguish all  
395 aspects of evolutionary change (Ciampaglio et al. 2001). Moreover, we note that in studies  
396 with a higher number of traits than the five simulated here, it will be even more challenging  
397 to abstract generalities.

398 We have primarily investigated the patterns of a directionally selective extinction  
399 acting on one trait. Further studies could examine reductions in multi-variate shape space at  
400 extinction boundaries (Korn et al. 2013), by employing simulations of non-homogeneous  
401 patterns of tree and trait evolution, and patterns of discrete character trait change. Trait  
402 evolution in our models is simulated under Brownian motion, such that disparity levels tend  
403 to increase linearly through time, and, in the absence of extinction, disparity is higher than  
404 relative diversity in the early history of a clade (Figs 2). In the absence of selective extinction  
405 (Korn et al. 2013), clades are expected to have top-heavy disparity profiles in which disparity  
406 is higher later in time (Gould 1990; Hughes et al. 2013; Deline et al. 2018; Hill et al. 2018).  
407 This observations contrasts strikingly with the empirical observation that there is a significant  
408 but weak tendency for clades to have low diversity but high disparity relatively *early* in their  
409 evolution (Foote 1994; Hughes et al. 2013). In our simulations we employ homogeneous  
410 models of speciation and extinction, but alternative patterns are expected to arise from  
411 heterogeneous diversification rates through time. Additionally, simulations and evolutionary  
412 models can employ different relations between traits and extinction, such as modeling  
413 extinction as a logistic function of trait values (Slater et al. 2017).

414 Rapid speciation early in a clade's history is expected to result in disparity being  
415 higher between (rather than within) clades: this leads to a pattern in which average subclade

416 disparity decreases towards the present (Harmon et al. 2003). However, the patterns we  
417 present are not directly comparable to Disparity-Through-Time (DTT) approaches as DTT  
418 methods calculate disparity at each point in time using all tip descendants of nodes in a  
419 phylogenetic tree (Harmon et al. 2003). Here we estimate disparity based on all of the  
420 branches present at a point in time, or we otherwise pool these samples into time bins.

421         A key question in paleobiology concerns whether mass extinctions selectively remove  
422 taxa based on their trait values or whether extinction is random with regards to traits. To this  
423 end, previous phylogenetic approaches to detecting extinction selectivity at mass extinctions  
424 have utilized pGLS models to test whether surviving and extinct lineages differ significantly  
425 in trait values (Friedman 2009; Puttick et al. 2017; Allen et al. 2019). pGLS models are  
426 appropriate in this context to correct for non-independence of residuals in the regression as a  
427 result of shared phylogenetic history (Felsenstein 1985), but we show that pGLS models are  
428 not able to correctly support the hypothesis that trait values differ in extinct lineages  
429 compared to surviving lineage trait values (Fig 5). In contrast, non-phylogenetic Ordinary  
430 Least Squares analyses correctly support a significant difference between trait values in  
431 survivors and losers at a mass extinction boundary. The cost of Ordinary Least Squares  
432 approaches is that they have type-one error rates above 5%, so not using phylogenetic  
433 correction cannot be recommended (Felsenstein 1985). Thus, the lack of pGLS support for a  
434 selective extinction may not mean that the extinction was random but may reflect a type-two  
435 error. This may result from the strictly homogeneous pattern of evolution employed here,  
436 which may not reflect biological reality. However, the solution for this problem is unclear, as  
437 it is not straightforward to compare non-phylogenetic and phylogenetic modelling results  
438 (Freckleton 2009), and impossible to differentiate true signals from model error. One  
439 solution, albeit unsatisfactory, is to infer trait-selectivity by analyzing the phylogenetic signal  
440 in extinction, without considering traits directly. Previous research has indicated that it is

441 possible to differentiate selectivity types using the phylogenetic signal of extinction (Hardy et  
442 al. 2012; Harnik et al. 2014; Krug and Patzkowsky 2015; Puttick et al. 2017; Soul and  
443 Friedman 2017). We support these observations because selective and non-selective  
444 extinctions have significantly different phylogenetic signals in our simulations (measured  
445 using the *phylD* statistic), and because the amount of phylogenetic diversity lost during an  
446 extinction event is dramatically different for random and selective events. We note, however,  
447 that even low levels of random noise (25% extinction non-selective, *weak selectivity*)  
448 produces results much closer to a fully random extinction simulation rather than one with  
449 selectivity.

450         After recovery from mass extinctions, extant species trait distributions still carry the  
451 signature of trait-selective extinction. This selectivity of extinction towards specific  
452 morphologies leads to changes in the relative support of phylogenetic comparative methods  
453 away from the simulated Brownian motion model to support an Ornstein-Uhlenbeck process  
454 (Fig 6A,C). Selective mass extinctions remove the left hand of the trait distributions, so the  
455 variance of tip trait values is lower than expected under a Brownian motion model; this trait  
456 distribution then resembles the expected distribution under the Ornstein-Uhlenbeck process,  
457 so this model is supported (Fig 6E). At face value, this may appear to be a bias that does not  
458 support the simulation model (BM) over an erroneous alternative model (OU). If OU is  
459 interpreted as a measure of stabilizing selection (Hansen 1997), then support for this model is  
460 correct in the context of a selective mass extinction. Taxa in a selective extinction are  
461 removed as they possessed trait values that made them prone to extinction. The OU  $\alpha$  value  
462 from the models here can be interpreted as measuring a discrete event of stabilizing selection  
463 (Butler and King 2004) or as an evolutionary optimum (Hansen 1997) that confers survival to  
464 a mass extinction event. In an analogous situation, a disruptive extinction event leads to a bi-

465 modal distribution in which a distribution resembles an Early Burst pattern over alternative  
466 models (Fig 6b).

### 467 ***Conclusions***

468 Our results indicate that patterns of disparity change across a mass extinction boundary are  
469 more complex than patterns of diversity change. As would be expected, disparity generally  
470 decreases following a selective extinction, but most indices show both increases and  
471 decreases in disparity across boundaries.

472         Understanding whether mass extinctions are selective or random with regards to traits  
473 is a major open question in paleobiology. Our results suggest that differentiating selective  
474 and non-selective extinction is difficult with phylogenetic comparative methods, but it is  
475 easier to identify phylogenetic signal in extinction itself. Mass extinctions also have long-  
476 term impacts on trait distributions and comparative models of trait evolution, even when  
477 traits evolve by a simple, time-homogeneous process.

478         All conclusions here assume that disparity may represent a real biological signal or a  
479 phenomenological description of patterns of trait evolution. Whatever trait disparity is  
480 measuring, we suggest that multiple indices are necessary in order to capture all aspects of  
481 disparity change across an extinction boundary, and a number of parameters need to be  
482 considered when inferring extinction disparity patterns.

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681

682 **FIGURE LEGENDS**

683 **Fig 1.** Workflow of simulation and analyses. In the simulations, trees and traits are simulated  
684 with time-homogeneous birth-death and Brownian motion processes (A). After 50, 100, 200  
685 contemporary tips are present, a mass extinction removes 50, 75, or 90% of lineages (B). We  
686 analysed scenarios in which there is a *directional selectivity* so lineages with larger values  
687 above a cut-off are prone to extinction, and *disruptive selectivity* in which trait values closer  
688 to the mean are liable to extinction. Strict selectivity means only lineages within the cut-off  
689 go extinct, strong selectivity applies the same bias but with a non-zero probability of  
690 extinction for all lineages, and in a random extinction all lineages are equally susceptible to  
691 extinction. (B). For all simulations disparity is measured for traits using a number of indices  
692 (C), with summaries of the levels of diversity and disparity (D).

693 **Fig 2.** Summarises of relative and disparity through time in simulations with no mass  
694 extinctions for iterations run until 50, 100, and 200 contemporary tips are present. For each  
695 simulation time, disparity, and diversity measures are scaled to unity, and each line represents  
696 a single simulation. The relative disparity patterns through time from Sum Of Ranges are  
697 consistent across iterations, but the patterns from other metrics are more variable.

698 **Fig 3.** The relative difference between diversity (red lines) and disparity (grey lines)  
699 measured in simulations of one trait with 200 extant tips (background extinction 0.8). Results  
700 are scaled to show the extinction at the midpoint of each iteration that destroyed 0.5 or 0.9 of  
701 contemporary lineages; the median (dark line) and full range (shaded areas) of all iterations  
702 are summarised. Full all iterations the variance of disparity indices across all iterations is  
703 larger than the trends in diversity patterns. When extinctions are strict or strong diversity and  
704 disparity exhibit similar trends with large decreases following extinction, except when  
705 disparity is measured using median root distance. Non-selective extinctions do not generally  
706 lead to disparity decreases.

707 **Fig 4.** Relative change in disparity across a mass extinction boundary for simulations with  
708 one (A), two (B), and five (C) traits. The dark horizontal line at zero indicates a null model of  
709 no disparity change across an extinction, values below this line indicate a decrease in  
710 disparity, and values above indicate a disparity increase. The figures summarise data for all  
711 considered disparity indices, including the amount of lineage loss. For two (B) and five (C)  
712 traits the first, darker box shows the traits that evolved independently and the lighter, second  
713 box shows traits that were simulated with co-variance. Vertical shading denotes selectivity of  
714 trait values.

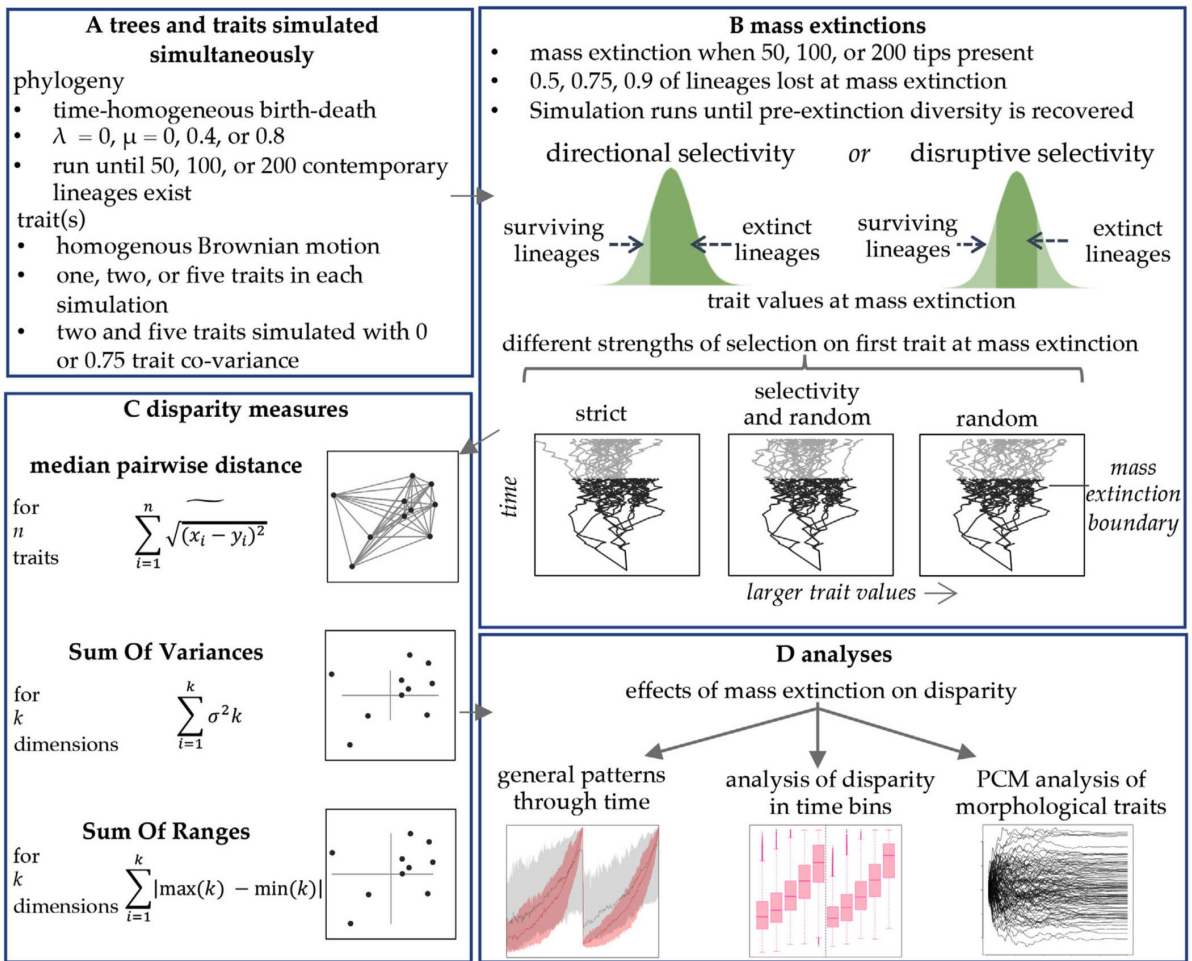
715 **Fig 5.** Summary of the number of pGLS (green) and OLS (pink) models that supported a  
716 significant relationship between the trait value and extinction for different selectivity levels  
717 and severity of extinction. For most simulations there was a true relationship between  
718 extinction and trait values (A-C) so it is expected the majority of models would support a  
719 significant relationship (gray shading). For the selectivity models (A-C) OLS models more  
720 consistently support a significant relationship compared to pGLS, and models perform more  
721 poorly as the number of lost lineages at a mass extinction increases. When extinction is  
722 random with regards to traits (D) most models should reject a relationship between trait  
723 values and extinction (i.e., below 5%, gray shading); OLS models generally have a high type-  
724 two error rate but pGLS models consistently and correctly reject a relationship.

725 **Fig 6.** The effects of trait distribution and phylogenetic comparative models applied to extant  
726 lineages on a tree that when through a deep-time mass extinction. The results are summarised  
727 for a directional selectivity (A) in which selectivity is directed towards lineages with larger  
728 trait values and disruptive selectivity (B) in which extinction selectivity is directed towards  
729 lineages with trait values at the tails of the distribution. All data were simulated under  
730 homogeneous Brownian motion, but models applied to extant data only show higher support  
731 for the OU model when selectivity is directional (C) and support for the EB model when



732 selectivity is disruptive (D). A directional selectivity leads to a trait distribution that  
733 resembles a distribution expected under an OU process (E), and a disruptive extinction  
734 resembles an EB-type distribution (F).

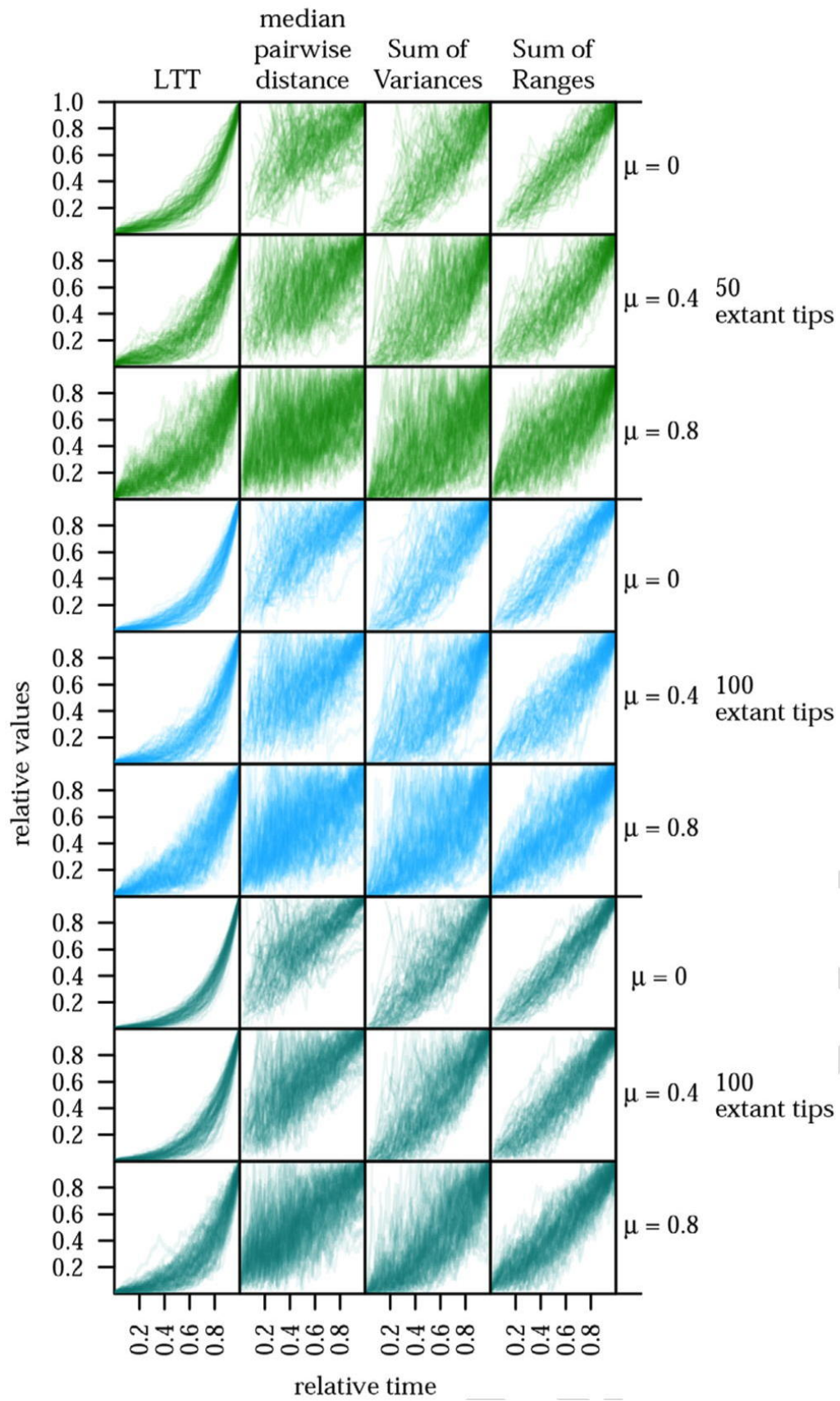
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736

737 Fig. 1

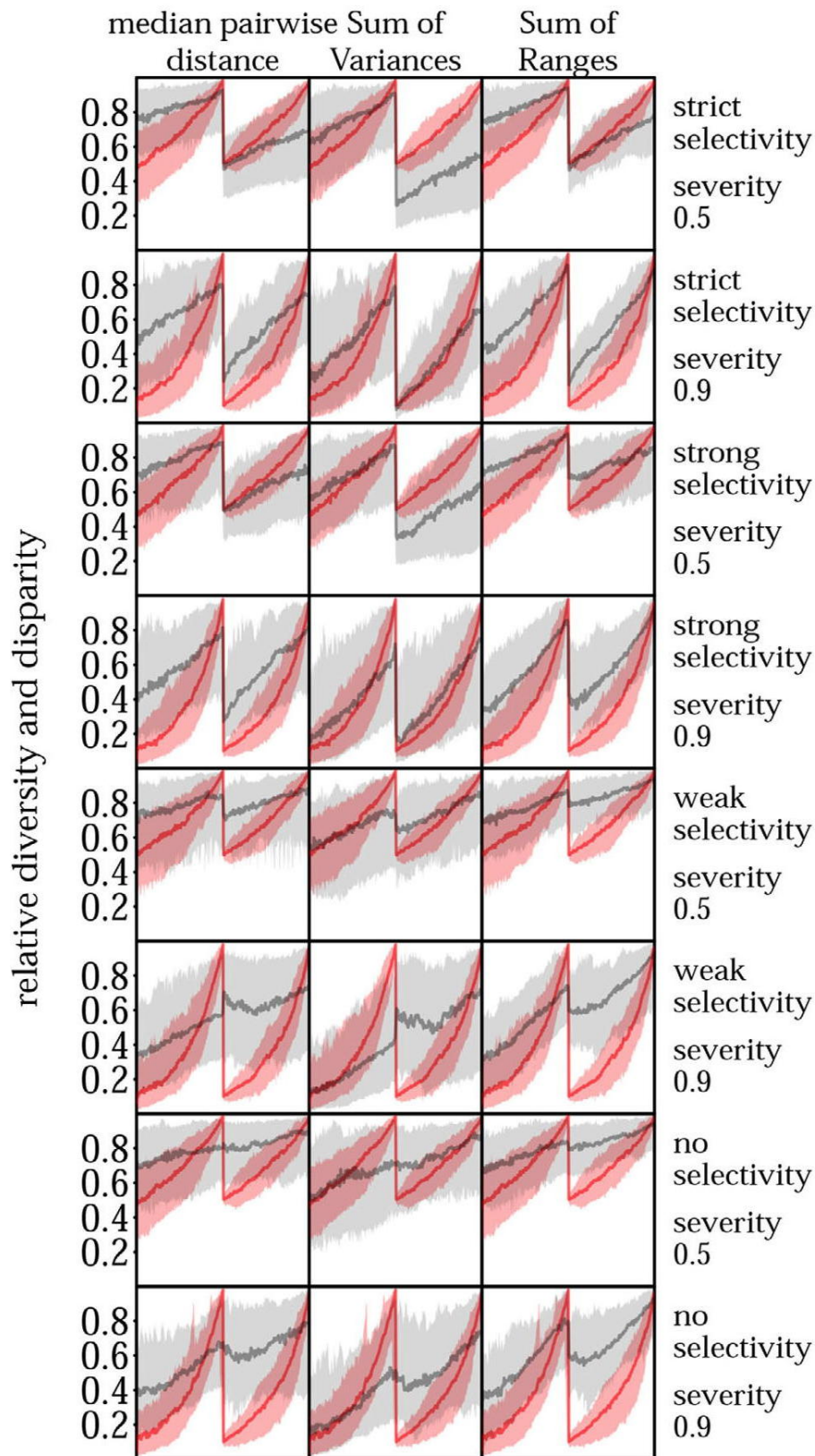
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740 Fig. 2.

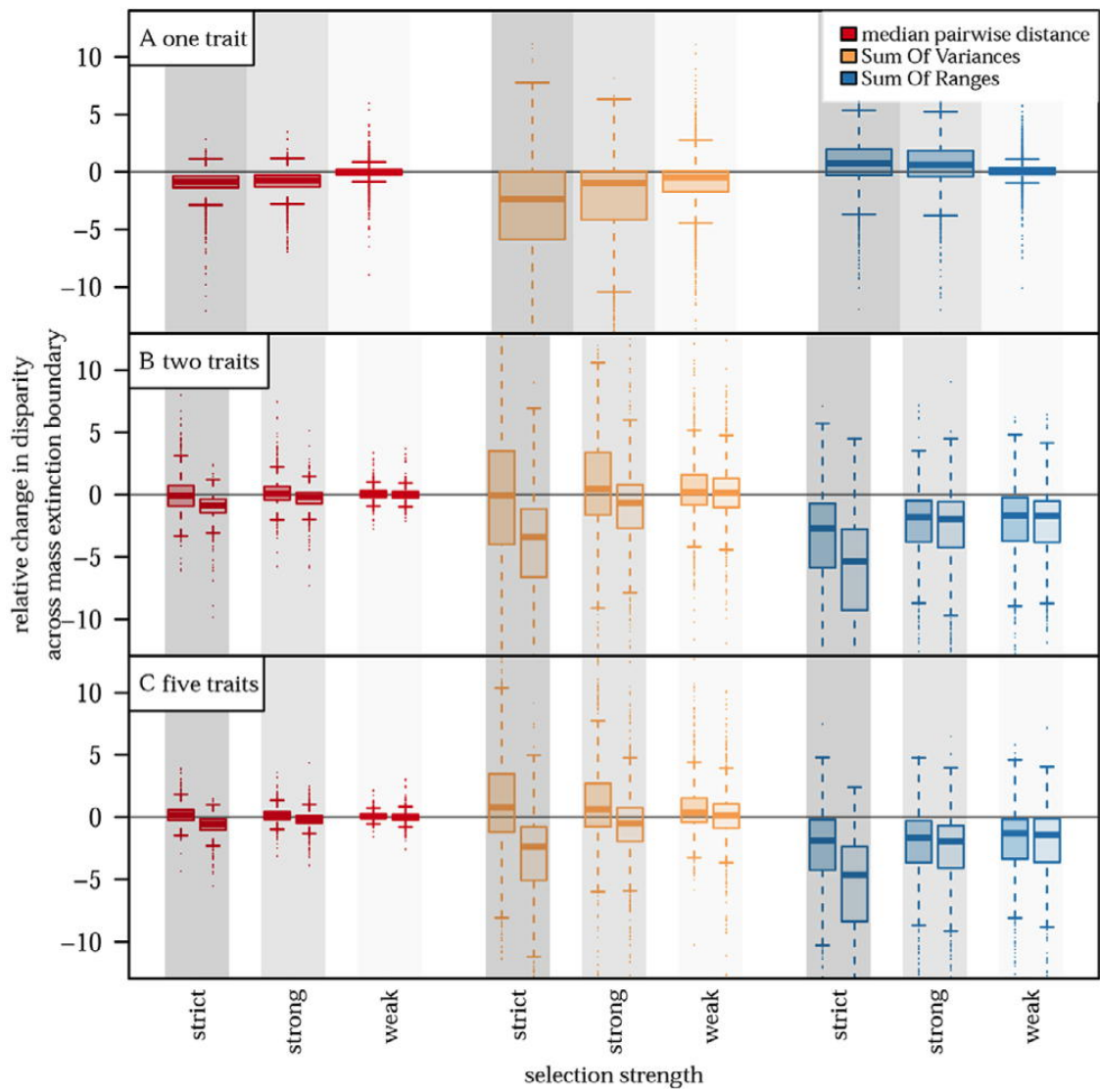
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743 Fig. 3.

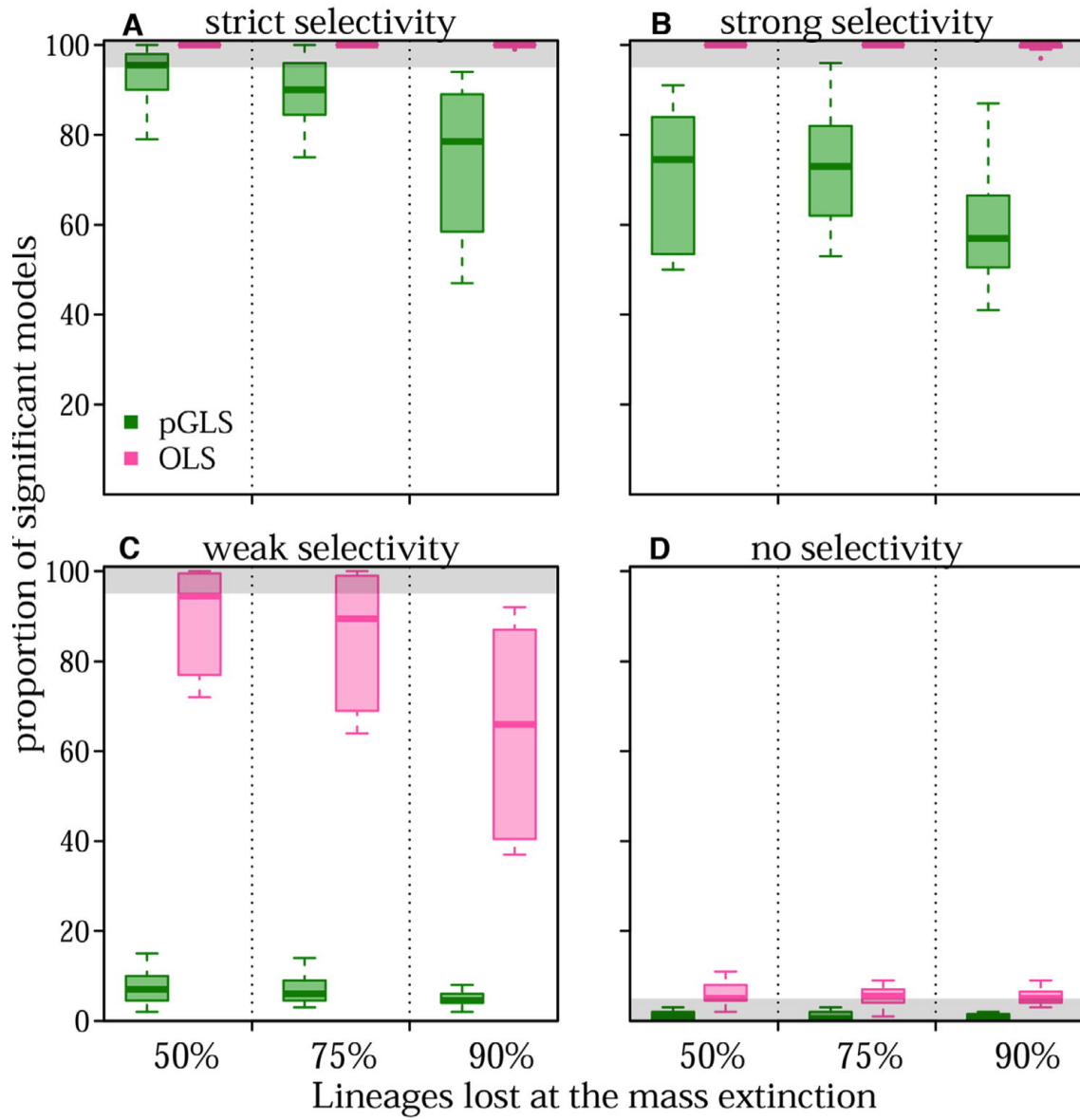
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746 Fig. 4.

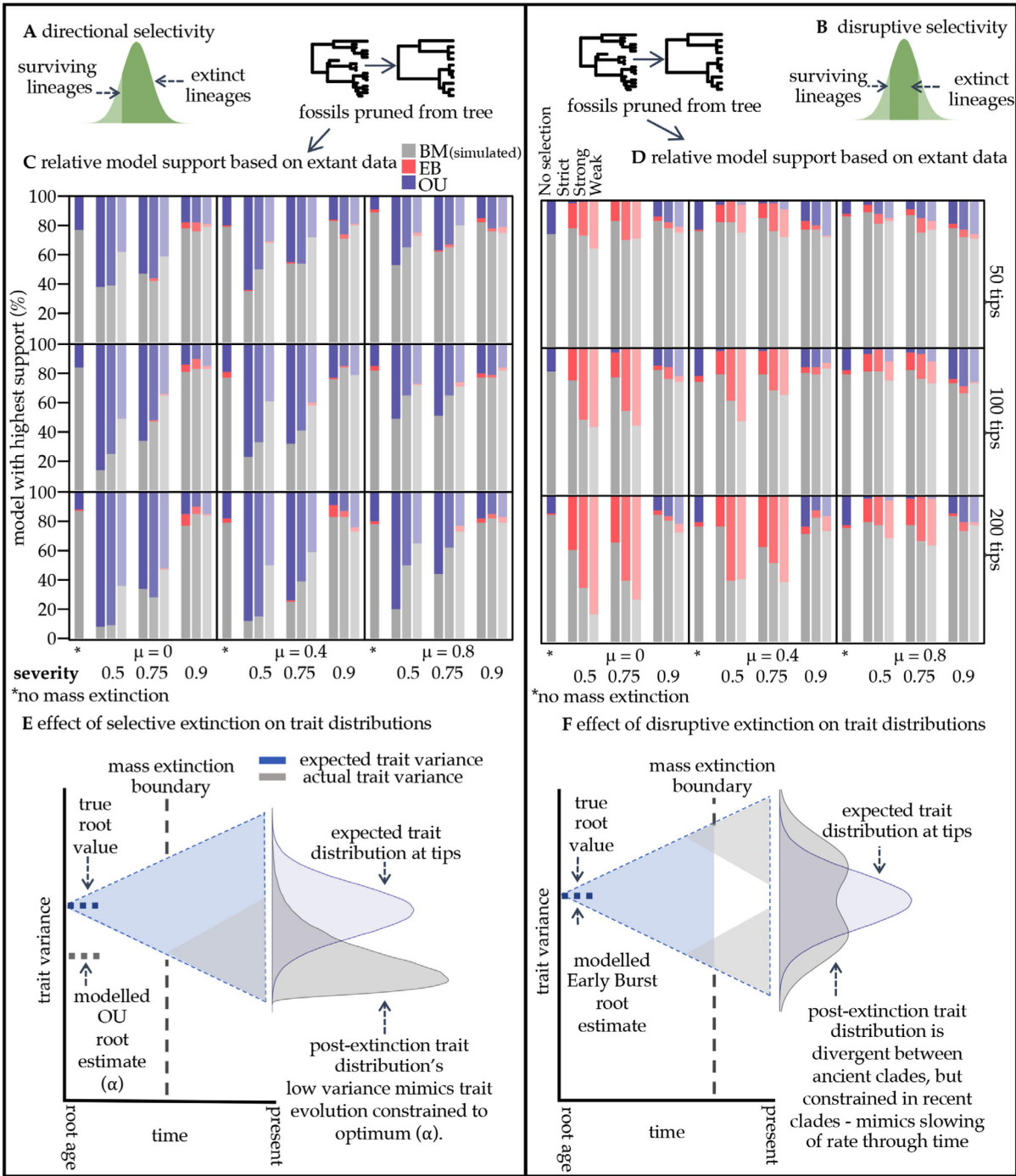
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749 Fig. 5.

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751

752 Fig. 6