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1 The evolution of multicellular complexity: the role of relatedness and

2 environmental constraints

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10 Keywords: major evolutionary transitions, multicellularity, scaling

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12 Abstract

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14 A major challenge in evolutionary biology has been to explain the variation in multicellularity 15 across the many independently evolved multicellular lineages, from slime moulds to 16 humans. Social evolution theory has highlighted the key role of relatedness in determining 17 multicellular complexity and obligateness, however there is a need to extend this to a 18 broader perspective incorporating the role of the environment. In this paper, we formally 19 test Bonner's 1998 hypothesis that the environment is crucial in determining the course of 20 multicellular evolution, with aggregative multicellularity evolving more frequently on land and 21 clonal multicellularity more frequently in water. Using a combination of scaling theory and 22 phylogenetic comparative analyses, we describe multicellular organisational complexity 23 across 139 species spanning 14 independent transitions to multicellularity and investigate 24 the role of the environment in determining multicellular group formation and in imposing 25 constraints on multicellular evolution. Our results, showing that the physical environment 26 has impacted the way in which multicellular groups form, could shed light on the role of the 27 environment for other major evolutionary transitions.

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31 Introduction

32 Macroscopic life on earth has been shaped by the evolution of multicellularity from 33 unicellular ancestors. Multicellularity is a complex and variable trait, ranging from simple cell aggregations found in yeast to differentiated metazoan organisms, with much diversity in 34 between [1]. For example, our bodies contain 10¹⁴ cells with more than 200 specialized 35 types [2] but Volvox is 10 orders of magnitude smaller and has just 2 cell types [3]. Some 36 37 lineages have become obligately multicellular, where cells only exist as part of a multicellular organism (e.g. animals), whereas others remain facultative, switching between 38 39 a unicellular and multicellular lifestyle (e.g. cellular slime moulds) [4].

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41 A major challenge in evolutionary biology has been to explain this variation in complexity among multicellular lineages. Social evolution theory has greatly advanced our 42 43 understanding of the evolution of multicellularity, primarily through clarifying the factors that favour the cooperation needed to become multicellular. We understand how relatedness 44 45 between cells is crucial in determining when altruism can evolve (for example, in 46 Dictyostelium slime moulds) [5], division of labour between cell types [6] and the 47 proliferation of cheaters [7-9]. It has also become clear that clonal relatedness (r = 1) is a necessary, albeit not sufficient, condition for the evolution of obligate multicellularity like we 48 49 see in animals and plants, and that these lineages have more cell types than those with 50 facultative multicellularity [4].

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52 However, there are limits to the variation in multicellular complexity that is explained by 53 relatedness. For example, both land plants and fungi have cells that are clonal and 54 obligately multicellular, but plants have approximately 10 times more cell types than fungi 55 [10] and it is unclear what can explain these differences. There are good reasons to 56 speculate that the environment could be an important factor shaping the first trajectories of 57 multicellular evolution with lasting consequences for later elaborations. Firstly, the 58 environment itself could determine the way in which multicellular groups form and hence 59 relatedness between cells. Bonner (1998) observed that clonal group formation, where daughter cells remain attached to mother cells after division, seems to be more common in 60 61 lineages that originated in the sea compared to species that originated on land [11]. If this is 62 the case, it would mean that the environment where multicellularity originates could have

63 profound consequences for subsequent evolutionary possibilities. Secondly, the physical 64 constraints associated with living in water or on land are likely to affect many aspects of 65 phenotypic evolution, for example the need for support and structural reinforcement tissues, 66 the diversity of dispersal mechanisms, and the sustaining the biomechanics of active 67 motility.

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69 Scaling theories provide powerful tools to test for such constraints, since an organism's 70 body size can accurately predict complex traits such as metabolic rate, lifespan, and growth 71 rate [12], and since the shapes of these relationships reflect fundamental physiological 72 constraints on how diverse organisms can evolve [13]. Scaling relationships can also reveal outlier taxa that highlight cases where evolutionary innovation fueled the breaking of 73 74 ecological and physiological constraints [14]. In practice, scaling parameters (*i.e.* the slope (b) and intercept (a) in the equation $y = aM^{b}$) represent mechanistic hypotheses that, for the 75 76 purpose of this study, relate the number of cell types (y) to the total number of cells (M). 77 Isometric scaling (b = 1) provides a null model, predicting that cell type and cell number 78 increase at the same rate (every added cell is a new type), and allometric scaling (b < 1) 79 would indicate that cell type increases at a slower rate than cell number, such that small 80 organisms have more cell types relative to their body size.

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82 There is a need to build on our understanding of the fundamental factors influencing 83 multicellular evolution - primarily the role of relatedness - and extend this to a broader 84 perspective incorporating the role of the environment. The objectives of this paper are to: 85 (1) describe the variation in multicellular organisational complexity across 139 species by 86 investigating the scaling relationships between body size (total number of cells) and number 87 of cell types; (2) use phylogenetically-controlled comparative analyses across 14 88 independent multicellular transitions to assess the extent to which the environment 89 determines how multicellular groups form and the consequences for whether obligate 90 multicellularity evolves; and (3) test whether constraints imposed by the environment can 91 explain why some lineages have reached higher levels of organisational complexity than 92 others, and can account for part of the variation in cell-type diversity and differences in 93 scaling relationships. We use the term organisational complexity to highlight that division of

94 labour is fundamental to multicellularity, that the number of cell types is a marker of division

95 of labour and that any form of division of labour requires organisational integration.

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98 **Results**

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100 Describing variation in body size and complexity

Across 139 species, representing 14 independent transitions to multicellularity, the scaling of cell type and cell number is strongly allometric (reduced major axis regression, RMA): slope = 0.14, CI = 0.13 - 0.16, R^2 = 0.64, Figure 1a & b). This means that despite a positive association, the number of cell types increases much more slowly with cell number than arithmetic proportionality (i.e. isometric scaling) would predict. In other words, small organisms are organisationally more complex for their size than large organisms.

107

We next found that there was a difference in the scaling relationship between number of cell 108 109 types and total number of cells for small versus large species (Table S1, S2). Specifically, we identified the estimated breakpoint in the regression as 6.3×10^4 total number of cells. 110 111 corresponding to 4.8 ± 0.9 cell types, where the scaling relationship changes. Small species (before the breakpoint) showed an allometric slope about twice as steep (RMA: slope = 112 0.27, CI = 0.23 - 0.33, R² = 0.61) as large species (above the breakpoint) (RMA: slope = 113 0.18, CI = 0.15 – 0.23, R^2 = 0.15). This implies that larger organisms face a unique set of 114 115 more stringent constraints on the accumulation of new cell types than small organisms, and 116 supports the observation that lineages consisting of small species gain new cell types more 117 quickly as they grow in size.

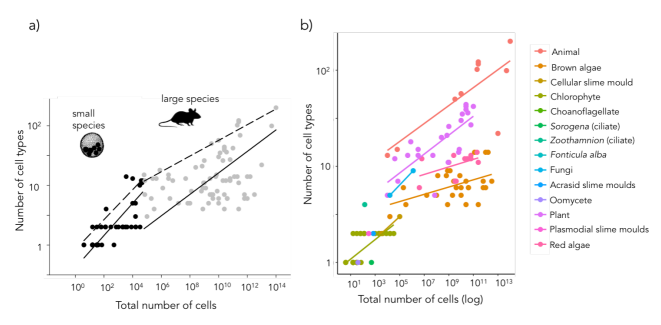
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We further sought to understand the substantial reduction in cell type variation explained by cell number for large organisms (i.e. $R^2 = 0.15$) using a technique called quantile regression. Regression through the upper 90% quantile of the dataset suggests that there is an upper threshold to the number of cell types a species can have for its size, whereas there is a lot of variation in the number of cell types below that threshold (Figure 1a, dashed lines). This suggests that: there could be other factors limiting the number of cell types below that threshold and these other limiting factors are especially important in larger

species since the slope describing this upper limit is far shallower (b = 0.13) than the upper limit for small species (b = 0.25) (Table S2).

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130 Figure 1(a) Scaling across multicellular organisms. The relationship between number of cell types and total number of cells for small (in black, between $4 - 10^4$ cells) and large multicellular 131 species (in grey, between $10^4 - 10^{14}$ cells) shown on logarithmic axes. Small species show a 132 steeper allometry (reduced major axis regression: slope = 0.27 (Cl 0.23 -0.33) compared to large 133 134 species (reduced major axis regression: slope = 0.18 (Cl 0.15 - 0.23). Solid lines show the reduced 135 major axis regression and dashed lines show regressions through the upper 90% quantile of the data. We estimated the breakpoint of 4.8 \pm 0.9 (corresponding to 6.3 x 10⁴ total cells) using the 136 137 'segmented' package in R. (b) Multicellular organisational complexity across different 138 multicellular lineages. Organisational complexity, measured as both the number of cell types and 139 the total number of cells, for each of the independently evolved multicellular lineages. These data 140 have been taken from the dataset of Bell & Mooers (1997) [10]. Original data are from the data set of 141 Bell & Mooers (1997) and images of Mus musculus and Volvox are from Phylopic 142 (http://phylopic.org/). The statistical results of the different regressions are given in Table S1.

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144 The origins of multicellularity in different environments

Our results show that the physical environment (whether or not a species lives in the water or on land) has had a major impact on both the origins and subsequent elaborations of multicellularity, both in determining how multicellular groups originally form and how organisational complexity subsequently evolves.

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150 We found that lineages in aquatic environments were significantly more likely to form 151 multicellular groups through daughter cells remaining attached to mother cells after division 152 (clonal group formation) (MCMCglmm, difference between aquatic & terrestrial: posterior mode = 5.74, credible intervals (CI) = 2.91 - 9.79, $p_{diff} = 0.0008$, $N_{species} = 139$, Figure 2a). 153 154 All of the multicellular lineages in our dataset that have their origins in water form 155 multicellular groups in this way, whereas two thirds of the lineages that originated on land 156 form groups through aggregation (non-clonal group formation) (Figure 2a). In fact, there are 157 only two lineages that originated on land that employ clonal group formation – the Fungi 158 and the plasmodial slime moulds and these tend to grow in terrestrial environments of 159 saturated humidity. This result confirms Bonner's original observation that clonal group 160 formation is more common in multicellular lineages originating in the sea [11].

161

162 Secondly, we found that the transition to obligate multicellularity was significantly more 163 likely to occur in aquatic environments compared to on land. Most (5 of 6) lineages that 164 evolved multicellularity on land remained facultatively multicellular (difference between 165 aguatic & terrestrial: posterior mode = 6.59, CI = 4.29 - 8.72, $p_{diff} = < 0.0001$, $N_{species} = 139$, 166 Figure 2b). The only multicellular lineage that has evolved obligate multicellularity on land is 167 the Fungi. This is consistent with this lineage also being a rare example of clonal group 168 formation that originated on land, as the resulting clonal relatedness between cells is 169 significantly associated with the transition to obligate multicellularity [4].

170

171 Table 1: At least 14 transitions to multicellularity occurred within the eukaryotes. Estimates 172 of the number of independent transitions in each lineage are given along with the environment 173 where the lineage originated, average number of cell types, and the corresponding references. We 174 have not included two other known transitions to multicellularity – the diatoms [15] and 175 *Sorodiplophrys* (Stramenopiles) [16] – due to a lack of data on cell types and environment of origin. 176 See Figure 3a.

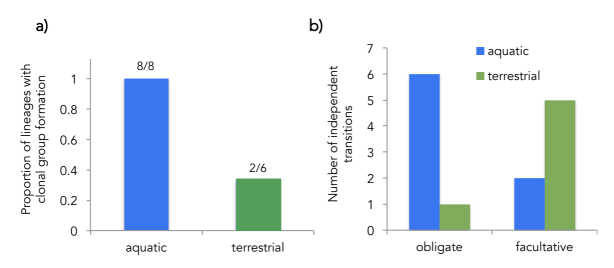
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Lineage	Number of	Average	Ancestral	Obligate or	Reference(s)
	transitions	number of	environment	facultative	
		cell types		multicellularit	
				у	

			1	fee liefter	14 71
Acrasid slime moulds	1	2	terrestrial	facultative	[17]
Brown algae	1	6.9	aquatic	obligate	[18,19]
Cellular slime	1	2	terrestrial	facultative	[20]
moulds					
Chlorophyte algae	1 - 4	1.5	aquatic	facultative &	[21]
				obligate	
Choanoflagellates	1	1	aquatic	facultative	[22]
Ciliates	2	2.5	terrestrial &	facultative &	[23-25]
			aquatic	obligate	
Fonticula alba	1	2	terrestrial	facultative	[26]
(Fonticulida)					
Fungi	2	7	terrestrial	facultative &	[27,28]
				obligate	
Metazoa	1	101.6	aquatic	obligate	[28]
Oomycetes	1	1	aquatic	facultative &	[29]
				obligate	
Plants	1	22.2	aquatic	obligate	[28]
Plasmodial slime	1	2	terrestrial	facultative	[30]
moulds					
Red algae	1+	10.8	aquatic	obligate	[28]

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179



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Figure 2: The origins of multicellularity in different environments. (a) The proportion of lineages that have clonal group formation that originated in aquatic and terrestrial environments. All multicellular lineages that originated in the sea have clonal group formation (8/8 lineages) whereas most of the multicellular lineages that originated on land have non-clonal group formation (4/6 lineages). (b) Multicellular lineages that originated in water more commonly evolve obligate
 multicellularity (6/8 lineages) compared to lineages that originated on land, which more often remain
 facultatively multicellular (5/6 lineages).

188

189 *Multicellular organisational complexity on land versus in water*

190 We found that the number of cell types of multicellular species currently found on land was significantly higher than those currently found in aquatic environments (Figure 3a), whilst 191 192 controlling for the total number of cells (posterior mode = -0.77, CI = -1.42 to -0.11, p_{diff} = 193 0.02, $N_{\text{species}} = 137$, Figure 3b). The average number of cell types for aquatic lineages is 8 194 whereas for terrestrial lineages it is 25. Species on land were however not significantly 195 larger in size than those found in the sea (posterior mode = -2.79, CI = -9.04 to 1.81, p_{diff} = 0.12, $N_{\text{species}} = 137$). Overall, there was a significant phylogenetic correlation between 196 197 number of cell types and total number of cells, meaning that species with more cell types also tend to be bigger due to their shared ancestry (posterior mode = 0.90, CI = 0.72 to 198 199 0.96, $p_{diff} = \langle 0.0001, N_{species} = 137 \rangle$. However, we also found a significant phenotypic correlation between these two variables, meaning that the association is also a result of a 200 201 shared environment (posterior mode = 0.56, CI = 0.19 to 0.76, p_{diff} = 0.004, $N_{species}$ = 137). 202

203

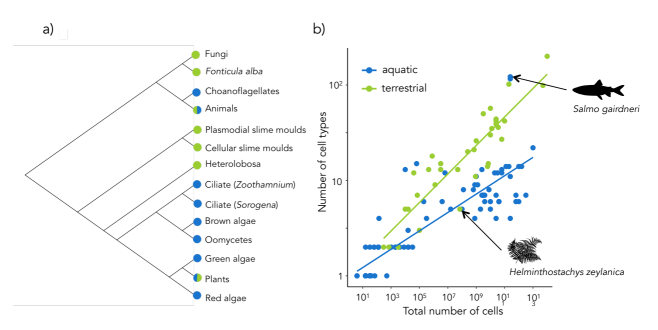


Figure 3: Organisational complexity and environmental constraints. a) A summarised phylogram of the lineages that have independently evolved multicellularity. The current environment is shown as terrestrial (in green), aquatic (in blue) or both (half green, half blue) for species that

have a substantial number of species in both environments. b) Multicellular complexity, measured
as both the number of cell types and the total number of cells, for species currently found in aquatic
and terrestrial environments. Two notable outliers (*Salmo gairdneri* and *Helminthostachys zeylanica*) are highlighted with black arrows and images and further interpreted in the Discussion.
These data have been taken from the dataset of [10].

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215 **Discussion**

216

217 We were interested in how multicellular complexity scales with body size and the role the 218 physical environment could play in shaping the course of multicellular evolution. Overall, we 219 found that the number of cell types scales allometrically with the total number of cells 220 (echoing Bonner's observations [1]), and that the specific scaling relationship is different for 221 small versus large species. Our comparative analyses also show that the environment 222 (aquatic or terrestrial) has a crucial impact on the trajectory of multicellular evolution. Firstly, 223 we found that clonal group formation giving rise to obligate multicellularity has been 224 significantly more common in lineages that evolved in aquatic environments. Secondly, we 225 showed that current environmental conditions have an impact on multicellular evolution, 226 with species living on land having a higher number of cell types compared to species found 227 in aquatic environments.

228

229 Bonner (1998) observed that clonal group formation was more common in multicellular 230 lineages that evolved in the sea whereas aggregation was more common in terrestrial 231 lineages. Our results provide formal support for this observation by including additional 232 lineages and using phylogenetically controlled comparative analyses. Bonner speculated 233 that this pattern could be because of water currents, meaning that cells in water need to 234 stick together after they divide if they want to reap the benefits of being in a group [11]. This 235 is not the case on land, where cells must use active motility (e.g. cilia, flagella, amoeboid 236 movement) in order to form multicellular groups. It is clear from the species in our dataset (Dictyostelium, Sorogena, Physarum, 237 that the multicellular lineages found on land 238 Fonticula) all have some form of motile cell stage, and even the Fungi have an ancestral 239 lineage with motile cells [31]. Therefore, it seems plausible that the biophysics of moving

through air and water has had a profound impact on the way in which multicellular groupscould form on land and in the sea.

242

243 It is likely that our dataset underestimates the number of lineages that have facultative 244 multicellularity. These species have transient multicellular phenotypes and so many species 245 are still identified as unicellular. For example, Saccharomyces cerevisiae has a variety of 246 multicellular phenotypes [32] and yet is still often not recognised as being facultatively 247 multicellular [33]. Other lineages, notably the green algae, have evolved facultative 248 multicellularity many times [21] and there are likely new examples to be found on land as 249 well. However, it is unlikely that we have underestimated the number of lineages with 250 obligate multicellularity. This is because these species tend to be bigger and therefore more 251 visible and complex [4] and potentially better studied [18,34]. There is no obvious reason to 252 assume that under- or overestimation would be biased towards terrestrial or aquatic 253 species. By inflating the number of facultative lineages, we would therefore not alter the 254 pattern and the result we find – that obligate multicellularity has evolved much more often in 255 water compared to on land.

256

257 Our study reveals a number of intriguing outliers. Firstly, not all species fit the overall 258 pattern of higher organizational complexity on land. There are outlier species in our dataset. 259 including several living in aquatic environments that display levels of complexity more 260 similar to terrestrial species. For example, Salmo gairdneri (Rainbow trout) has an estimated 116 cell types and a total size of 2.51 x 10^{11} cells, which is more similar to the 261 262 terrestrial Mus musculus (mouse) than to other aquatic species. Another example is 263 Helminthostachys zeylandica (a member of the fern family) that has much lower complexity 264 for its size than other terrestrial multicellular species (just 5 cell types) (Figure 3b). 265 Secondly, a major and strikingly unusual lineage are the Fungi, that develop multicellularity 266 through clonal group formation but display (mostly) simple multicellularity. However, our 267 dataset only included 3 species from Kingdom Fungi and there is also a lack of data on 268 fungal multicellularity in the wider literature. Perhaps a closer look at the Fungi as 269 'exceptions to the rule' could help to unravel the relationship between the environment and 270 multicellular complexity.

271

272 Not only does the environment affect how multicellular groups form, but we show that it also 273 has a major impact on the scaling relationships between size and complexity. Species that live on land tend to be more complex for their size compared to species that live in water 274 275 (i.e., with a higher slope, Figure 3b) and this could be for several reasons. Land dwelling 276 organisms need more support structures than their aquatic counterparts – this is because 277 water provides natural support through buoyancy whereas air does not. Organisms living on 278 land therefore needed to increasingly invest in stems and skeletons to 'hold themselves' up as their body size increases (i.e. skeleton mass ~ $M^{b>1}$, [12]), possible leading to greater 279 280 diversification of cell types and tissues than organisms in the sea. This scaling logic can 281 further be extended to resource allocation dynamics within organisms (e.g., vascular 282 networks,), although systematic effects on cell diversity and differences between land and 283 water remain to be elucidated. There are also other potentially confounding physiological 284 parameters, for example that autotrophic lineages compete for light, both on land and in the 285 water, whereas heterotrophic lineages do not, but that support tissues on land are more 286 costly to maintain (e.g. rain forest trees versus kelp forest).

287

288 The parallel evolutionary events towards obligate multicellularity are examples of major 289 evolutionary transitions in individuality [35]. A key aim of major transition research is to 290 identify common patterns across different transitions (e.g. the evolution of prokaryote and 291 eukaryote cells, obligately multicellular organisms, and colonial superorganisms). The fact 292 that these all arose through clonal group formation or as full sibling families (initiated by 293 strictly monogamous pairs) implied that reproductive allocation conflicts did not play a role. 294 as they usually do in promiscuous or chimeric associations that do not make such major 295 transitions [36,37] [4,38]. Strict vertical transmission of symbionts, including mitochondria 296 and plastids, was also a potent force to avoid conflict [39]. Our results, showing that the 297 physical environment has impacted the way in which multicellular groups form, could 298 therefore shed light on the role of the environment for other major evolutionary transitions. 299 For example, how have physical conditions across nesting habitats (e.g. subterranean 300 versus arboreal; [40]) influenced the necessary and sufficient conditions for insect colonies 301 to commit to obligate division of labour via specialized and physically differentiated castes? 302

- 303

304 Material and Methods

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306 Data collection

307 The data used in this study were originally published in Fisher *et al.* (2013) and are stored 308 data depository Dryad (original data be found in the can here: 309 https://datadryad.org/resource/doi:10.5061/dryad.27q59). In summary, we conducted an 310 extensive literature search on multicellular species, searching specifically for information on 311 multicellular complexity, the ways in which groups formed and whether or not they were 312 obligately or facultatively multicellular. We used data from Bell & Mooers (1997) on the 313 number of cell types and total number of cells to estimate multicellular complexity for each 314 multicellular species [10] as this is the most taxonomically-representative dataset on cell 315 types, to our knowledge [41]. Our full dataset can be found in Table S6.

316

317 In this study, we expanded this original dataset by adding information on the ancestral and 318 current environment of each species. We considered any species found on land as 319 terrestrial and any species found in freshwater, brackish or marine environments as 320 aguatic. We found information about the current environment of a species by searching on 321 Google Scholar for publications and also taxa-specific websites, such as AlgaeBase and 322 WoRMs. Where there was only information about ancestral or current environment at a 323 higher taxonomic level (i.e. at the family level but no generic or species information) we 324 assumed it was the same environment for the species in our dataset. We found information 325 on the ancestral environment of each species through broad reviews on the origins of 326 multicellularity including Bonner 1998, Knoll 2011 & Umen 2014 [11,21,28]. It is important 327 to stress that we were interested in the ancestral environment when multicellularity evolved 328 and therefore that was not always the same as the ancestral environment for the whole 329 lineage, including unicellular groups (e.g. for the Fungi, James et al. 2006).

330

Of the 139 species in the dataset, 18 species had a terrestrial ancestral environment and 121 species had an aquatic ancestral environment. For the current environment, 84 species are aquatic, 43 are terrestrial and 12 are unknown.

334

335 Independent transitions to multicellularity

336 Using information from published papers, we identified that within the eukaryotes there 337 have been at least 14 independent transitions to multicellularity (both facultative and 338 obligate) (Table 1, Figure 3a). However, we have most likely underestimated the number of 339 transitions in several groups due to uncertainty about the number of independent transitions 340 within them. For example, it is thought that there have been at least 2 transitions to obligate 341 multicellularity within the Fungi [27,28] and many transitions to facultative multicellularity in 342 the green algae [21] and in the red algae [18]. Therefore, our analyses are conservative 343 and assumed just 1 transition within each group.

344

345 Statistical Methods

346

347 Scaling relationships

348 As a first step in analyzing the data we began with a least square regression to estimate a 349 and b in the scaling equation $\log_{10} y = \log_{10} a + b \log_{10} M$ and describe nature of the 350 dependence of the number of cell types on the total number of cells. We used the R 351 package 'Imodel2', we used reduced major axis (RMA) regression to estimate the intercept 352 and slope in the scaling of log₁₀(cell type) against log₁₀(cell number) across all data, for 353 small species and for large species. RMA is an appropriate line-fitting method in cases 354 when measurement of both Y and X variables are potentially associated with systematic 355 error (e.g. the probability that cell number was precisely measured decreased with 356 increasing body sizes) [42]. RMA (also known as standardized major axis regression) 357 equally weights distances from the regression line in both X and Y directions, with the major 358 axis reflecting the first principal components axis yielded by the covariance matrix, and 359 fitted through the centroid of the data [42]. We then used the package 'segmented' in R [43] 360 to test if there is a 'breakpoint' in the regression - the point at which the shape of the 361 relationship changes dramatically. This allowed us to estimate the different scaling 362 relationships of small versus larger multicellular species.

363

We also noted that the scaling relationships appeared triangular and thus hypothesized that they reflect a constraint function such that total number of cells is necessary, but not sufficient to explain variation in number of cell types [44,45]. To test this hypothesis, we

used least absolute deviation regression to describe scaling for the upper ninetieth
 quantiles of the overall plot and separately for the small and large taxa plots [46,47].

369

370 Bayesian analyses

371 We used the statistical package MCMCgImm [48] to run Bayesian general linear models 372 with Markov Chain Monte Carlo (MCMC) estimation. We fitted three models. Firstly, we 373 tested whether the environment affected the way in which multicellular groups form by fitting 374 a model with group formation as a categorical response variable and the ancestral 375 environment as a categorical explanatory variable (Table S1). Secondly, we tested whether 376 the environment affected the likelihood of obligate or facultative multicellularity by fitting a 377 model with obligate/facultative as a categorical response variable and the ancestral 378 environment as a categorical explanatory variable (Table S3).

379

380 Finally, we tested whether multicellular complexity differed between lineages living on the 381 land versus in aquatic environments by fitting a multi-response model with several 382 explanatory variables using the number of cell types and the logarithm of total number of 383 cells as poisson and Gaussian response variables respectively (Table S4). This allowed us 384 to use both number of cell types and the total number of cells as a combined measure of 385 multicellular complexity, rather than having to run several analyses using different response 386 variables. We fitted several categorical fixed effects: the current environment (aquatic or 387 terrestrial), whether the species is obligately or facultatively multicellular, and the mode of 388 group formation (non-clonal or clonal) to control for the known effects of group formation 389 and obligateness on complexity [4].

390

391 In the first two models, we used uninformative inverse-gamma priors because we had a 392 categorical response variable. We also fixed the residual variance to 1 and specified family 393 = categorical. In the final model, we used uninformative priors because we had a multi-394 response model with both poisson and Gaussian response variables and categorical 395 explanatory variables. We ran the models for 6000000 iterations, with a burn-in of 1000000 396 and a thinning interval of 1000. These were the values that optimised the chain length 397 whilst also allowing our models to converge, which we assessed visually using VCV 398 traceplots. We then ran each model three times and used the Gelman-Rubin diagnostic to

quantitatively check for convergence. We showed our models had converged when the
PSR was < 1.1.

401

We calculated the correlations between the number of cell types and the total number of cells (cov(number of cell types, total number of cells)/sqrt(var(number of cell types)* var(total number of cells)) for species in different environments. We tested if the correlation was significantly different between environments by examining if the 95% credible interval of the difference between the correlations spanned 0, and calculating the % of iterations where the correlation for species living in aquatic environments was greater than that for those living on land.

409

410 **Phylogeny construction**

411 We built the phylogeny for this study using the Open Tree of Life (opentreeoflife.org), which 412 creates synthetic trees built from published phylogenies and taxonomic information. We 413 then used the R package 'rotl' that interacts with the online database and constructs 414 phylogenies (https://cran.r-project.org/web/packages/rotl/index.html). For the majority of 415 species in our dataset, the exact species was also present in a published phylogeny and so 416 we could use phylogenetic information about that species. However, for a few species that 417 were not present in the Open Tree of Life dataset, we had to assign instead a closely 418 related species in the same genera or use a family-level classification. Due to the fact that 419 most species in our dataset represent distant groups on the eukaryotic tree and our 420 phylogeny does not include branch lengths, we were confident this compromise did not 421 affect our statistical analysis.

422

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426

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431

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