

# Genome-enabled discovery of evolutionary divergence in brains and behavior

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## Research Article

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28 development, behavior

29 **Abstract**

30 Lake Malawi cichlid fishes exhibit extensive divergence in form and function built from a  
31 relatively small number of genetic changes. We compared the genomes of rock- and  
32 sand-dwelling species and asked which genetic variants differed among the groups. We  
33 found that 96% of differentiated variants reside in non-coding sequence but these non-  
34 coding diverged variants are evolutionarily conserved. Genome regions near  
35 differentiated variants are enriched for craniofacial, neural and behavioral categories.  
36 Following leads from genome sequence, we used rock- vs. sand- species and their  
37 hybrids to (i) delineate the push-pull roles of BMP signaling and *irx1b* in the specification  
38 of forebrain territories during gastrulation and (ii) reveal striking context-dependent brain  
39 gene expression during adult social behavior. Our results demonstrate how divergent  
40 genome sequences can predict differences in key evolutionary traits. We highlight the  
41 promise of evolutionary reverse genetics – the inference of divergence in phenotype from  
42 genome sequencing in natural populations.

## 43 **Introduction**

44 Our understanding of how the genome encodes natural variation in form and function is  
45 still limited. This is the case for almost any trait, from height to behavior to complex  
46 disease <sup>1</sup>. The reasons for this are manifold, but they include an underappreciated role of  
47 non-coding genetic variants linked to differences in traits. This is apparent in our  
48 assumptions and in syntheses of data. For instance, only 25 years ago, experts thought  
49 that the human genome might contain 70,000 to over 100,000 genes to account for our  
50 complexity <sup>2</sup>. More recently, it has been estimated that upwards of 93% of human disease  
51 related variants – traits for which we have the most data from genome wide association  
52 studies (GWAS) – reside in noncoding DNA sequence <sup>3</sup>. Many of these noncoding  
53 variants are regulatory, that is, they affect the expression of genes <sup>4</sup>. Therefore, despite  
54 a refined understanding of how single genes work in controlled cellular environments, it  
55 remains unclear how the genome is activated to produce natural phenotypes, and this  
56 may be particularly vexing for context-dependent processes like development or  
57 behavior.

58  
59 Over the past two decades, systems have been developed to identify the genetic basis  
60 of traits from nature <sup>5</sup>. Amongst vertebrate animals, these traits include body armor <sup>6</sup>,  
61 color <sup>7</sup>, head and jaw shape <sup>8-10</sup>, parental care <sup>11,12</sup>, song <sup>13</sup> and coordinated movement  
62 <sup>14</sup>. The take home message from this work has been that a small number of genes from  
63 recognizable pathways explain a considerable proportion of phenotypic variance. Yet,  
64 these studies may be biased in interpretation and limited in inference space. The focus is  
65 typically on one or two species and one trait at a time, often using hybrid pedigrees  
66 founded by a small number of individuals, and candidate gene or QTL approaches. Here,  
67 we explored a different strategy in a system of many species with many divergent traits.  
68 We sought to determine the genetic differences between closely related groups of species  
69 and then to focus experiments on leads from genome divergence. In essence, we've  
70 asked the genome which traits to follow.

71  
72 The Malawi cichlid system is an apposite one for our research aims. The assemblage  
73 comprises hundreds of closely related species that have diversified in the last 500,000 to

74 one million years <sup>15</sup>, such that the genomes of individuals across species boundaries  
75 remain highly similar <sup>16</sup>. An appreciable fraction of genetic polymorphism identified in  
76 Malawi species is shared with cichlid lineages from throughout East Africa -- suggesting  
77 that ancient genetic variation fuels diversification of the Malawi flock <sup>17</sup>. Set against this  
78 background of genome similarity, Malawi cichlids exhibit staggering diversity in  
79 phenotypes including pigmentation <sup>18</sup>, sex determination <sup>19,20</sup>, craniofacial and brain  
80 patterning <sup>8,21,22</sup> and social behavior <sup>23-25</sup>. Previous work has focused on the genomic and  
81 early developmental underpinnings of this diversity, in rock- vs. sand-dwelling species  
82 <sup>16,21,22,26</sup>.

83  
84 Rock- vs. sand- species form ecologically distinct groups similar to other ecotypes in well-  
85 known adaptive radiations (marine vs. freshwater sticklebacks; tree vs. ground finches  
86 and anoles) <sup>27</sup>. The main difference in this case is that each of the rock- and sand- groups  
87 contains more than 200 species. Recent divergence, rapid speciation and meta-  
88 population dynamics synergistically lead to the broad sharing of polymorphism across the  
89 rock-sand speciation continuum <sup>17,28</sup>. Malawi rock-dwellers tend to be strongly territorial  
90 and aggressive; they breed and feed at high density in complex rock-reef habitats. Most  
91 eat algae from the substratum with strongly reinforced jaws packed with teeth. Adult rock-  
92 dweller brains exhibit enlarged anterior components, telencephala and olfactory bulbs.  
93 Sand-dwellers are less site-specific and less aggressive. They often breed on communal  
94 leks where males build sand 'bowers' to attract females <sup>29</sup>. Many capture small prey using  
95 acute vision and fast-moving gracile jaws; their brains and sensory apparatus are  
96 elaborated for more posterior structures optic tecta, thalamus and eyes (Supplementary  
97 Figure 1). We aimed to understand evolutionary divergence between rock- and sand-  
98 dwelling lineages by identifying the number, type and spectrum of genetic variants that  
99 separate these groups.

100  
101 To target this primary axis of evolutionary divergence in the Lake Malawi species  
102 assemblage <sup>27</sup>, we compared whole genomes of one male individual each from 8 rock-  
103 dwelling and from 14 sand-dwelling species (Supplementary Table 1), to an average of  
104 25X coverage per individual. Species were chosen to represent the diversity present

105 within each of the rock- and sand- groups (Figure 1A), in terms of body size, color pattern,  
106 ecology and phylogenetically defined lineages within the sand- species group <sup>28</sup>.

## 107 **Results**

### 108 *The genomic signature of rock-sand divergence*

109 We compared the genomes of 8 rock dwellers and 14 sand dwellers to uncover the  
110 genomic signature of rock- versus sand- evolutionary diversification. We aligned  
111 sequence data to a reference genome of nearly 1 gigabase<sup>30</sup> and identified  
112 approximately 22 million Single Nucleotide Polymorphisms (SNPs) and 200,000  
113 Insertion-Deletions (InDels). We calculated  $F_{ST}$  per variant, and averaged across 10kb  
114 windows, to quantify divergence between rock and sand species. We found that 0.06%  
115 of SNPs and 0.44% of InDels are alternately fixed between rock- and sand- groups. When  
116 these divergent variants and genome regions (2.5% FDR) are mapped to linkage groups  
117 (chromosomes), it is apparent that the signature of rock- vs. sand- divergence is  
118 distributed relatively evenly across the chromosomes (Figure 1B). Among fixed variants,  
119 3.5% were found in coding regions and 96.5% were predicted to be non-coding; ~17% in  
120 intergenic regions, 38% in introns, 38% in flanking regions (within 25kb up- or  
121 downstream of a gene), and 3% in annotated UTRs. Rock vs. sand fixed coding variants  
122 were more likely to be missense/loss-of-function (72.6%) than silent (27.3%).

123

124 We next generated whole-genome alignments of five published cichlid reference  
125 genomes from across East Africa<sup>31</sup> and estimated an evolutionary conservation score for  
126 each nucleotide position. Akin to phylogenetic footprinting, this approach allows inference  
127 of function for regions that are slower to change than others due to the long-term effect  
128 of purifying selection. For both coding and non-coding portions of the genome, we found  
129 that rock-sand divergence correlates positively with evolutionary conservation scores  
130 (Figure 1C), suggesting that differentiated rock-sand variants, including many non-coding  
131 variants, are enriched for function.

132

133 A total of 4,484 genes lie within 25 kb of either an alternately fixed variant or a highly  
134 divergent 10kb window (2.5%FDR). Pathway enrichment analysis<sup>32</sup> of human  
135 homologs/analogs for these genes reveals categories spanning early embryonic  
136 development, craniofacial morphogenesis, brain development, synaptic transmission and

137 neuronal function (Supplementary Table 2). In particular, rock-sand divergent genes are  
138 enriched for GO Biological Process terms ‘telencephalon development’ ( $p < 1.7e-18$ ),  
139 ‘adult behavior’ ( $p < 2e-14$ ), ‘synaptic plasticity’ ( $p < 1.4e-12$ ), ‘odontogenesis’ ( $p < 3.7e-$   
140 11), ‘response to BMP’ ( $p < 3.2e-09$ ), ‘gastrulation’ ( $p < 5.6e-06$ ), ‘face morphogenesis’  
141 ( $p < 8.9e-08$ ), ‘neural crest cell differentiation’ ( $p < 4.3e-13$ ), and ‘eye development’ ( $p <$   
142 1.3e-15). Over-represented gene families included nuclear hormone receptors ( $p < 3.0e-$   
143 08), HOXL subclass homeoboxes ( $p < 1.4e-07$ ), TALE class homeoboxes ( $p < 4.2e-04$ )  
144 and Forkhead boxes ( $p < 9.33e-04$ ; for novel expression domains in cichlid *foxp2* see  
145 Supplementary Figure 2). We observed enrichment for the mouse phenotypes ‘abnormal  
146 cognition’ ( $p < 3.3e-15$ ), ‘abnormal learning and memory’ ( $p < 2.6e-15$ ), ‘abnormal  
147 craniofacial morphology’ ( $p < 4.9e-11$ ) and ‘abnormal social/conspecific interaction’ ( $p <$   
148 7.1e-14). We used the list of differentiated genes to query an Allen Brain Atlas dataset  
149 that reports gene expression in hundreds of brain regions<sup>33</sup>. Rock- sand- divergent genes  
150 were enriched for the basomedial nucleus of the amygdala, a sub-region of the  
151 telencephalon ( $p_{\text{adj}} = 0.001$ ) that regulates fear, anxiety, and physiological responses to  
152 territorial intruders in rodents<sup>34,35</sup>, and has been linked to Social Anxiety Disorder in  
153 humans<sup>36</sup>. Finally, we queried databases of genes involved in human disease. Genes  
154 near divergent variants are significantly enriched for factors implicated in neurological  
155 disease like Autism Spectrum Disorder (SFARI<sup>37</sup>, Fisher’s exact test  $p$  value  $< 2e-16$ )  
156 and disorders related to the neural crest<sup>38</sup>, (Fisher’s exact test  $p$  value  $< 2e-16$ ).

157

158 Given the prevalence of evolutionarily conserved, non-coding, divergent rock-sand  
159 variants and genome-wide enrichment for craniofacial and neural crest biology, we  
160 examined overlap with published datasets of mammalian neural crest and craniofacial  
161 enhancers<sup>39,40</sup>. These data allow us to identify craniofacial and cranial neural crest cell  
162 (CNCC) enhancers conserved between mammals and cichlids and fixed variants  
163 between rock and sand species within these conserved regulatory elements. A total of  
164 275 craniofacial enhancer elements and 234 human CNCC enhancers are evolutionarily  
165 conserved between mammals and cichlids. We found divergent rock-sand mutations  
166 within the enhancer elements of key genes integral to CNCC specification and migration  
167 (Supplementary Table 2). Notably, from both datasets, fixed rock-sand variants were

168 found within the enhancer region of the gene *nr2f2*, a nuclear receptor and master neural  
169 crest regulator<sup>41</sup>. Rock-sand divergent variants were similarly located within craniofacial  
170 enhancers of three genes (*yap1*, *fat4*, *rere*) that function in the Hippo signaling pathway,  
171 as well as within enhancers of *irx3* and *axin2*. These data linking rock-sand fixed  
172 SNP/InDels to evolutionarily conserved, experimentally verified enhancers further  
173 underscore the importance of non-coding variation in the craniofacial evolution of rock-  
174 and sand- lineages<sup>42</sup>.

175

176 Genome-wide divergence between rock vs. sand Malawi cichlids involves a relatively  
177 small percentage of genetic variants. Divergent variants are (a) predominantly non-  
178 coding, (b) in long-term evolutionarily conserved loci (c) enriched for genes and pathways  
179 involved in embryonic development, brain development, brain function and behavior, and  
180 craniofacial morphogenesis. Given these strong patterns of enrichment, we used the  
181 experimental power of the Malawi cichlid system to interrogate features of early  
182 development and adult behavior that differ between rock- and sand- groups.

183

#### 184 *A gastrula-stage map of forebrain diversification*

185 Rock- vs. sand-dwelling Malawi cichlids exhibit divergence in or near genes enriched for  
186 BMP signalling, gastrulation, eye and telencephalon development, as well as the TALE  
187 (*Ir*x) gene family. To explore the developmental consequences of this differentiation, we  
188 investigated early forebrain specification in rock- and sand- embryos, building upon our  
189 previous studies and interest in *irx1b* and early brain development<sup>21,22</sup>. During  
190 development, the complexity of the vertebrate brain is first laid out in the neural plate, a  
191 single-cell thick sheet of cells that forms between non-neural ectoderm and the germ ring  
192 at gastrulation. *Ir*x genes act as transcriptional repressors of BMP signal in gastrulation,  
193 and function to specify the neural plate<sup>43</sup>. BMPs, in turn, are protective of the anterior-  
194 most region of the neural plate, which will ultimately give rise to the telencephalon, and  
195 suppress the eye field<sup>44</sup>. Given alternatively fixed variants in the *irx1b* gene, expected  
196 interactions between *Ir*x and BMP signaling in the early embryo and known telencephalon

197 vs. eye size differences between rock- vs. sand- species <sup>21,22</sup>, we examined and  
198 quantified the early activity of *irx1b* and BMP in rock- vs. sand- embryos.

199  
200 We used a custom device to orient and image cichlid embryos *in toto* at gastrula and  
201 neurula stages <sup>45</sup>. In early gastrula (EG), *irx1b* (red) and BMP signal (green, PSMAD)  
202 delineate complementary dorsal and ventral domains of the embryo (Figure 2A). By mid-  
203 gastrula (MG), *irx1b* shows two expression domains, one in the posterior portion of the  
204 developing neural plate (np) and the second co-expressed with PSMAD activity around  
205 its anterior border (white arrowheads). By late gastrula (LG), the domains of *irx1b*  
206 expression and PSMAD activity sharpen around the leading edge of the neural plate but  
207 remain overlapping around the periphery. Notably, *irx1b* expression is expanded in the  
208 anterior domain of sand-dwellers (S) compared to rock-dwellers at EG and MG, and then  
209 defines the boundary of the neural plate earlier in sand-dwellers (S) in LG (arrowheads).  
210 As a consequence, BMP signal should have a longer-lasting influence on the neural plate  
211 in rock-dwelling species. Based upon manipulative experiments in zebrafish <sup>44</sup>, this is  
212 predicted to result in a relatively larger presumptive telencephalon and smaller eye field.

213  
214 We developed a panel of rock- x sand- hybrid crosses to formally evaluate the role of  
215 *irx1b* in forebrain diversification. First, we used quantitative RT-PCR to measure allele-  
216 specific expression (ASE) in heterozygous rock- x sand- F<sub>2</sub> hybrids, across the stages of  
217 gastrulation. We observed that the sand- *irx1b* allele was expressed at significantly higher  
218 levels (average of 2.5-fold;  $p = 4.5e-13$ ; Student's t-test) and that this difference was  
219 largely confined to MG (Figure 2B). Next, we used hybrid embryos to chart the  
220 development of the telencephalon and the eye field. Rock- x sand- F<sub>2</sub> hybrids, indexed  
221 for *irx1b* genotype, were raised to neurula and somitogenesis stages and we examined  
222 the expression of *shh* (which induces *foxd1* and the ventral forebrain), *foxd1* (a marker of  
223 the telencephalon), and *rx3* (a marker of the eye field), by in situ hybridization. F<sub>2</sub>  
224 individuals homozygous for rock- *irx1b* alleles exhibited a larger and more rostral domain  
225 of *shh* expression, an earlier and larger domain of *foxd1* and a smaller *rx3* domain (Figure  
226 2C, D). These differences between rock- vs. sand- *irx1b* genotypes match expression  
227 divergence observed amongst rock- vs. sand- species <sup>21,22</sup>. Finally, when we compared

228 the relative size of the telencephalon among *irx1b* genotypes, individuals homozygous  
229 for rock- alleles exhibited larger telencephala (Supplementary Figure 3). We conclude  
230 that genetic variants in and around the *irx1b* gene contribute to divergent specification of  
231 the Malawi cichlid forebrain, likely via spatial, temporal and quantitative variation in the  
232 expression of *irx1b* itself.

233  
234 Our genome sequencing revealed a near-fixed InDel in the 3' UTR of the Malawi cichlid  
235 *irx1b* gene (Supplementary Figure 4). Rock- species possess an 85bp insertion,  
236 compared to cichlid species from outside of the Malawi lineage. Sand-dwellers largely  
237 lack the insertion and exhibit a 6bp deletion compared to outgroups. The insertion shows  
238 strong genetic similarity to a fragment from the Rex1 family of non-LTR retrotransposons  
239 <sup>46</sup>. Given the likelihood that *Astatotilapia calliptera* populations surrounding Lake Malawi  
240 may have seeded the Malawi evolutionary radiation and contributed to rock- and sand-  
241 dwelling lineages <sup>17,28</sup>, we explored the presence/absence of this InDel in *Astatotilapia*  
242 samples. We found that most *Astatotilapia* individuals and populations had the rock- *irx1b*  
243 allele (the insertion), but that an individual from Chizumulu Island was fixed for the sand-  
244 allele and two individuals sampled from Itupi were heterozygous. Because the 85bp  
245 insertion in rock- species is a partial Rex1 fragment, and sand- species carry a 6bp  
246 deletion compared to outgroups, we speculate that the current rock- and sand- divergent  
247 alleles were generated by at least two imperfect excision events of an element that  
248 invaded the genome of the Malawi + *Astatotilapia* ancestor. Rex1/Babar retrotransposons  
249 have been active in African cichlid genomes, and are known to influence gene expression  
250 when inserted in 5' and 3' UTRs <sup>31</sup>. Future experiments will determine whether this Rex1  
251 insertion causes the differences in *irx1b* gene expression and forebrain specification we  
252 observed.

253

#### 254 *Genomics of divergent social challenge and opportunity*

255 Rock- and sand-dwelling Malawi cichlids live in strikingly different social and physical  
256 environments. Rock-dwelling males tend to be more aggressive than sand-dwellers <sup>23</sup>  
257 and defend territories year-round as sites for feeding and breeding. By contrast, sand-

258 dwellers are more exploratory than rock-dwellers <sup>25</sup> and only breeding males tend to be  
259 territorial, often building sand bowers to attract females and mitigate male-male  
260 aggression. Given these observations and genome-wide enrichment for categories  
261 related to adult behavior and social interaction, we designed an experimental paradigm  
262 to investigate brain gene expression profiles associated with divergent rock- vs. sand-  
263 social behaviors.

264  
265 We evaluated social challenge and opportunity amongst males using a large tank with a  
266 'rock' habitat at one end and 'sand' at the other, separated by glass bottom (Figure 3A).  
267 When parental rock- species are placed in this tank paradigm, males court females on  
268 the rock side of the tank. Males of sand- species court females over sand and construct  
269 species-appropriate bowers. When single hybrid rock- x sand- F<sub>1</sub> males are placed in this  
270 arena with hybrid F<sub>1</sub> females, males invariably court females over the 'rock' habitat.  
271 However, when two rock- x sand- hybrid F<sub>1</sub> males (brothers) were allowed to compete for  
272 gravid hybrid F<sub>1</sub> females in this tank paradigm, we observed something different. One  
273 male, typically the larger, courted females over the rock habitat, and the other  
274 simultaneously constructed bowers to court females over the sand. We found no  
275 difference in gonadal-somatic index (GSI), an established biological metric of  
276 reproductive status and maturity, between F<sub>1</sub> males behaving as 'socially rock' vs.  
277 'socially sand.' (Supplementary Figure 5). Our observation of divergent behavior between  
278 F<sub>1</sub> brothers in the same tank suggests an interaction between the genome and the social  
279 environment.

280  
281 We used RNA-seq to investigate gene expression profiles associated with behavior of  
282 rock- x sand- F<sub>1</sub> hybrid males that were actively courting females over rock vs. sand.  
283 Whole brains of F<sub>1</sub> males tested singly (n=2 lone) as well as F<sub>1</sub> brothers assayed in dyads  
284 (n=4 dyads) were collected during courtship, and interrogated by RNA-seq. Strikingly,  
285 gene expression profiles clustered not by fraternal relatedness, but rather by behavioral  
286 context (Figure 3B). Males from dyads that courted females over rocks had expression  
287 profiles similar to single males (who also courted over rocks) but distinct from their  
288 brothers that built bowers and courted females over sand in the same tank. Genes were

289 considered significantly differentially expressed between 'social rock' and 'social sand'  
290 brains if they exhibited both a fold change  $\geq 2$  and crossed the threshold of  $p_{\text{adj}} < 0.05$ .  
291 Based on this criterion, we found 832 genes differentially expressed between rock- vs.  
292 sand-behaving males (Figure 3B, Supplementary Table 3). Among differentially  
293 expressed genes, we observed significant functional enrichment for GO Biological  
294 Process categories 'synaptic signaling' ( $p < 2.3e-21$ ), 'synaptic plasticity' ( $p < 3.6e-09$ ),  
295 'visual behavior' ( $p < 2.09e-06$ ); mouse phenotypes 'abnormal  
296 learning/memory/conditioning' ( $p < 5.9e-07$ ), 'abnormal telencephalon morphology' ( $p <$   
297  $3.95e-07$ ), 'abnormal spatial learning' ( $p < 9.9e-07$ ) and pathways 'axon guidance' ( $p <$   
298  $3.3e-05$ ), 'oxytocin signaling' ( $p < 5.2e-05$ ) and 'estrogen signaling' ( $p < 1.8e-04$ )  
299 (Supplementary Table 3). Matches against the Allen Brain Atlas database of gene  
300 expression yielded enrichment for exclusively sub-regions of the telencephalon: CA3,  
301 hippocampus ( $p_{\text{adj}} = 4.6e-6$ ), CA2, hippocampus ( $p_{\text{adj}} = 7.3e-5$ ), CA4, hippocampus ( $p_{\text{adj}}$   
302  $= 0.001$ ), claustrum ( $p_{\text{adj}} = 0.001$ ), subiculum ( $p_{\text{adj}} = 0.003$ ), dentate gyrus ( $p_{\text{adj}} = 0.03$ )  
303 and the basomedial nucleus of the amygdala ( $p_{\text{adj}} = 0.03$ ). The hippocampus encodes  
304 episodic memory and spatial representations of the environment <sup>47</sup>, and more recently its  
305 subregions have been shown to play critical roles in anxiety, social interaction, and social  
306 memory formation <sup>48-50</sup>. Roughly 38% of differentially expressed genes also contained  
307 genetically differentiated SNP/InDels between rock- and sand- species ( $p$ -value  $< 2e-6$ ,  
308 Fisher's exact test), implying considerable cis-acting genetic variation. Enrichment of  
309 numerous categories related to brain function and synaptic plasticity showed greater  
310 overlap than expected (Figure 3C). These context-dependent differences suggest  
311 concerted changes in brain gene expression as males experienced and responded to  
312 different social challenges and opportunities <sup>24,51</sup>.

313 **Discussion**

314 A fundamental problem in evolutionary biology is understanding the cellular,  
315 developmental and genetic basis of how traits change. This is a challenge because we  
316 lack sufficient information about how genes work in outbred genomes from nature and we  
317 do not fully comprehend the causal role of noncoding variation in specifying form and  
318 function. This problem is especially difficult for traits that are only observed in particular  
319 contexts, like development and behaviour. To make progress, we and others have  
320 focused on study systems exhibiting abundant phenotypic diversity built from a relatively  
321 small number of genetic changes. Here we identify and characterize the genetic variants  
322 that demarcate one of the deepest evolutionary splits amongst Lake Malawi cichlid  
323 groups, that between rock- and sand-dwelling species thought to have diverged in the  
324 last one million years. We found a small percentage (less than 0.1%) of genetic variants  
325 to be differentiated between rock- and sand- groups, and that the majority of differentiated  
326 variants (>96%) were noncoding. Differentiated non-coding variants were more likely to  
327 be in an evolutionarily conserved locus as a function of genetic differentiation, suggesting  
328 that divergent rock- vs. sand- noncoding changes are functional. To support this idea, we  
329 identified alternately fixed rock- vs. sand- noncoding variants within experimentally  
330 verified, vertebrate-conserved craniofacial and cranial neural crest cell enhancers. The  
331 latter observation is similar in type to the discovery of human-specific deletions within  
332 mammal-conserved regulatory sequence <sup>52</sup>.

333

334 Recently we surveyed genome-wide divergence between sand-dweller sub-groups that  
335 construct pit vs. castle bowers, sand-made structures to attract females for mating <sup>24</sup>.  
336 Mapping those variants to the same genome reference, we expected distinct patterns of  
337 diversification because rock-sand and pit-castle divergence likely occurred at different  
338 times, along different trait axes, under the control of different evolutionary forces <sup>27</sup>.  
339 Consistent with expectation, there is clear clustering of genome divergence on  
340 chromosome 11 for the pit-castle comparison (Supplementary Figure 6), while all  
341 chromosomes carry the signature of rock-sand diversification (Figure 1B). However,  
342 contrary to our expectation, rock- vs. sand- and pit- vs. castle- radiations have diverged  
343 in similar gene sets. Out of 3070 genes identified near 10kb high  $F_{ST}$  regions in the rock-

344 vs. sand- comparison, 483 overlap with 1090 genes identified near high  $F_{ST}$  regions in  
345 the pit- vs. castle- comparison (p-value < 2e-9, Fisher's exact test, Supplementary Table  
346 4). This result may imply that evolutionary diversification in Lake Malawi is limited, or  
347 constrained, by chromosomal location.

348

349 Overall, genes in proximity to rock-sand divergent variants were enriched for functional  
350 categories related to early forebrain and craniofacial development, neuronal function and  
351 social behavior. This list of variants, coupled with consistent patterns of functional and  
352 pathway enrichment, motivated follow up experiments focused on early brain  
353 development and adult social behavior (Figure 4). It is apparent from our work here and  
354 previously <sup>21,22</sup>, that Malawi cichlid brains and nervous systems begin to differ during  
355 gastrulation in pathways that can be predicted from divergent genome sequences. This  
356 is interesting for at least two reasons. First, this observation runs counter to the 'late  
357 equals large' textbook example <sup>53</sup> of how brains evolve differences in relative proportions  
358 of their parts <sup>22</sup>. Similarly, such early variation in development is not thought to be a driving  
359 force in evolution, precisely because early changes can have global and ramifying effects.  
360 Collectively, our findings provide a partial description of the conditions wherein variation  
361 during the earliest stages of development can contribute to evolutionary diversification. In  
362 each case we have examined, variation in gene expression is quantitative, heterochronic  
363 and limited to a precise stage or time period.

364

365 Sydney Brenner recognized the relationship between the genetic specification of nervous  
366 systems and the behavioral output of the brain <sup>54</sup>. However, because these events take  
367 place so far apart in the lifespan of a vertebrate, they are rarely studied simultaneously.  
368 Here, the genome connects the two phenomena: rock- vs. sand- divergent gene sets  
369 indicating that both brain development and social behavior have been under divergent  
370 selection during the evolutionary diversification of these groups. To evaluate social  
371 behavior in rock- vs. sand-dwelling Malawi cichlids, we constructed a social context  
372 arena. The presence of sand and simulated rocky caves was sufficient to elicit species-  
373 appropriate male behavior when rock- or sand- males were tested with rock- or sand-  
374 gravid females. When rock- x sand-  $F_1$  hybrid males were tested, one per tank, with  $F_1$

375 hybrid females, males courted females in the rock quadrant of the tank. Notably, when  
376 dyads of F<sub>1</sub> brothers were tested in this tank paradigm with gravid F<sub>1</sub> females, we  
377 observed simultaneous 'social rock' and 'social sand' behavior. Brain gene expression  
378 profiles from behaving males clustered by social context (social rock vs social sand), and  
379 not by fraternal relationships. Differentially expressed genes were enriched for brain  
380 regions and pathways implicated in social interaction and overlapped significantly with  
381 rock- vs. sand- divergent genetic variants.

382

383 Social context is known to influence the brain. For instance, our clustering results are  
384 similar to those of Whitfield and colleagues (Whitfield et al. 2003) who showed that brain  
385 gene expression in honey bees was predictive of behavior. Likewise, changes in brain  
386 morphology and gene expression predictably accompany the ascent to dominance in the  
387 cichlid fish *Astatotilapia burtoni*<sup>55</sup>. Our data seem not to fit the model of dominant-  
388 subordinate however. In our experiments, the gonado-somatic index (GSI) did not differ  
389 between social rock vs. social sand brothers within dyads. Both males exhibited nuptial  
390 coloration, courted females and in cases with multiple gravid females, both brothers  
391 reproduced. Body size was associated with divergent social rock vs. social sand behavior  
392 of F<sub>1</sub> males; the social rock brother was always larger (mean mass was 26.96g ± 3.4 [SE]  
393 compared to 19.45g ± 2.2).

394

395 Our experiments demonstrate that F<sub>1</sub> hybrid male brains can express both social rock-  
396 and social sand- behavioral programs, and that social context determines which program  
397 is executed. This observation is similar to, but also different than, pit-digging x castle-  
398 building F<sub>1</sub> male Malawi cichlids who carry out parental bower behaviors in a specific  
399 sequence<sup>24</sup>. Notably, in both cases, the hybrid males exhibit one of the two different  
400 parental behaviors at any one time – there is no intermediate behavior. In the pit- vs.  
401 castle- case, we think that the bower structure itself and/or a threshold signal from females  
402 might lock the hybrid male brain into a behavioral state. In the rock- vs sand- case here,  
403 it appears that other social cues (i.e., the presence and size of a rival male) lock the hybrid  
404 male brain into a behavioral state. These context-dependent behaviors, accompanied by  
405 changes in brain gene expression, are compelling examples of interaction between the

406 genome and the social environment. The cellular and genetic basis of these behaviors  
407 and their plasticity deserves further attention. Our comparative genomic and brain gene  
408 expression data, combined with enrichment testing and experimental approaches,  
409 highlight that the Malawi cichlid telencephalon will be central to this future work.

## 410 **Materials and Methods**

### 411 *Genome sequencing*

412 We extracted genomic DNA, from fin clips of 22 male individuals (Qiagen DNeasy, Cat  
413 #69504), from 8 rock dwelling and 14 sand dwelling Lake Malawi species (Supplementary  
414 Table 2) representing broad diversity across the rock and sand lineages in Lake Malawi.  
415 We made libraries using the Illumina Nextera Library prep kit and performed paired-end  
416 sequencing on the Illumina Hi-Seq 2500 at Georgia Tech. The *Metriaclima zebra*  
417 reference genome version MZ\_UMD2a<sup>30</sup> was used for genome alignment, variant  
418 discovery and annotation using standard BWA<sup>56</sup> and GATK practices<sup>57</sup>. The maximum  
419 likelihood tree in Figure 1A was constructed using SNPPhylo<sup>58</sup>, from variant data.

420

### 421 *Genetically Divergent Regions*

422 Vcftools<sup>59</sup> was used to calculate  $F_{ST}$  (--weir-fst-pop) between the 8 rock and 14 sand  
423 species. Variants with  $F_{ST} = 1$  were noted to be alternately fixed between rock and sand  
424 lineages in our dataset.  $F_{ST}$  was also measured across 10kb windows (--fst-window-size).  
425 Significance thresholds were marked using the fdrtool package in R. All variants were  
426 annotated using Snpeff 4.3i<sup>60</sup>. We tested the genes within 25 kb of significantly  
427 differentiated variants for enrichment of functional categories. The cichlid gene names  
428 were converted to human analogs using Treefam based mapping<sup>61</sup> and functional  
429 enrichment was determined using the TOPPFUN web-browser interface<sup>62</sup>.

430

### 431 *PhastCons analysis*

432 Pairwise alignments were generated using lastz v1.02<sup>63</sup>, with the following parameters:  
433 "B=2 C=0 E=150 H=0 K=4500 L=3000 M=254 O=600 Q=human\_chimp.v2.q T=2  
434 Y=15000". This was followed by using UCSC genome utilities  
435 (<https://genome.ucsc.edu/util.html>,  
436 [https://hgdownload.soe.ucsc.edu/admin/exe/linux.x86\\_64/FOOTER](https://hgdownload.soe.ucsc.edu/admin/exe/linux.x86_64/FOOTER)) axtChain tool with -  
437 minScore=5000. Additional tools with default parameters were then used following the

438 UCSC whole-genome alignment paradigm  
439 ([http://genomewiki.ucsc.edu/index.php/Whole\\_genome\\_alignment\\_howto](http://genomewiki.ucsc.edu/index.php/Whole_genome_alignment_howto)) in order to  
440 obtain a contiguous pairwise alignment. Multiple alignments were generated from  
441 pairwise alignments with the multiz v11.2<sup>64</sup> program, using default parameters and the  
442 following pre-determined phylogenetic tree: ((((*M. zebra*, *P. nyererei*), *A. burtoni*), *N.*  
443 *brichardi*), *O. niloticus*) in agreement with Brawand et al.<sup>31</sup>. Sequence conservation  
444 scores were then obtained using PhastCons<sup>65</sup> with a phylogenetic model estimated by  
445 the phyloFit<sup>66</sup> program, both from the PHAST software package (v.1.3). The model fitting  
446 was done using default parameters. PhastCons was run in two iterations, first to obtain  
447 the free parameters of the model (--estimate-trees and --no-post-probs) and then using  
448 the output from this we ran PhastCons again to attain the conservation scores with --  
449 target-coverage 0.3 --expected-length 100.

450

#### 451 *Vertebrate-conserved enhancer elements*

452 A comparative genomic approach was used to identify putative craniofacial and neural  
453 crest CNEs in mammals that segregate SNPs between rock-sand cichlid species.  
454 Experimentally verified and published genome-wide craniofacial and neural crest  
455 enhancers active during early embryonic stages that play a role in shaping the  
456 development of neural crest and craniofacial structures in mammals were identified from  
457 published literature<sup>39,40</sup>. We used the liftOver tool<sup>67</sup>, which maps orthologous genomic  
458 regions between species to convert genomic coordinates from one species to another.  
459 Using a Human to *Oreochromis niloticus* to *Metriaclima zebra* mapping and a Mouse to  
460 *Oreochromis niloticus* to *Metriaclima zebra* mapping, we identified the orthologous  
461 genomic locations of the published craniofacial and neural crest enhancers in cichlids.  
462 We designated any alternately fixed variant (variant with  $F_{ST} = 1$ ) that was also within an  
463 orthologous CNE as putatively involved in the rock-sand divergence (Supplementary  
464 Table 2).

465

#### 466 *Brain region enrichment analysis*

467 We identified 10,391 cichlid genes with human homologues and generated an expression  
468 matrix for each gene across 250 human brain structures spanning telencephalon,  
469 diencephalon, mesencephalon, and metencephalon using adult human brain microarray  
470 data collected by the Allen Brain Institute<sup>33</sup>. Cortical regions and gyri for which fish do  
471 not have putative homologues were excluded from the analysis (100/350, leaving 250  
472 regions for subsequent analysis). The expression matrix was generated using the  
473 `get_expression` function in the `ABAEnrichment` Bioconductor package in R<sup>68</sup>. We then  
474 calculated the specificity of expression for each gene in each of these brain regions using  
475 the `specificity.index` function in the `pSI` package for R. This function calculates a matrix of  
476 gene expression specificity indices, and corresponding p-values, as described previously  
477<sup>69,70</sup>. We then tested whether 1) genes within 25kb of rock vs. sand significantly  
478 differentiated variants (described above under “Genetically Divergent Regions”), and 2)  
479 genes that were differentially expressed between rock- vs. sand-behaving F1 hybrid  
480 males, were enriched for transcriptional markers of specific brain regions using the  
481 *fisher.iteration* function with Benjamini-Hochberg correction, again using the `pSI` package  
482 for R. For enrichment testing of differentially expressed genes, we restricted analysis to  
483 genes that met the following criteria: 1) transcripts for the gene were detected in all eight  
484 paired behaving males, and 2) at least 6 transcripts were detected in each subject.

485

#### 486 *Staging during gastrulation*

487 Cichlid gastrulation was split into three sub stages within the gastrula stage 9<sup>71</sup>.  
488 Gastrulation lasts 8 to 12 hours, depending on the species, and is defined as after the  
489 shield (as described in zebrafish) stage until the presence of the first somite at the  
490 beginning of neurula (stage 10). Embryos were classified as early gastrula (EG) by an  
491 asymmetry in epiboly after shield stage until the formation of a ridge that is analogous to  
492 the anterior neural ridge (ANR) in chick and mouse and the anterior neural border (ANB)  
493 in zebrafish. At that point embryos were classified as mid gastrula (MG). MG lasts until  
494 the formation of the dorsal-ventral axis, defined by further lengthening of one side of the  
495 embryo, which begins to thicken as epiboly progresses. This is the dorsal side of the  
496 embryo, and the side opposite the ANR is classified the ventral side of the embryo. At this

497 point the embryos are defined as late gastrula (LG). LG ends with the specification of the  
498 neural plate, which appears as a portion of the dorsal embryo that is raised relative to  
499 ventral side, usually in line with the ANR.

500

#### 501 *Immunohistochemical staining*

502 Embryos were harvested at 24 hours post fertilization (hpf) from each of the rock -dwelling  
503 cichlids *Metriaclima patricki* and *Metriaclima zebra* and the sand-dwelling cichlid  
504 *Copadichromis borleyi* and *Tramitichromis intermedius*. The embryos were cultured until  
505 they reached gastrula stage, approximately 36 to 40 hpf, then fixed at intervals throughout  
506 gastrula until neurula. The embryos were then treated with auto-fluorescence reducer  
507 (1.55mL 5M NaCl, 250ul Tris-HCl, pH 7.5, and 95mg NaBH<sub>4</sub>) overnight, and 10% 2-  
508 mercaptoethanol for 1 hour. Next, whole mount *in situ* hybridization was done, using a  
509 modification methods we published previously <sup>72</sup>. *irx1b* was visualized using Fast Red  
510 (naphthol chromogen, Roche Diagnostics), which fluoresces at near red wavelengths  
511 (500-650 nm). After *in situ* hybridization, embryos were immunostained for pSMAD 1,5,8  
512 protein, using published protocols <sup>73</sup>. Embryos were then bathed in Vectashield (Vector  
513 Labs) containing DAPI and placed in a specially built mold<sup>45</sup> that accommodates the large  
514 yolk and holds the embryo upright. Embryos were then scanned using a Zeiss LSM 700-  
515 405 confocal microscope and processed using LSM 700 software and Image J.

516

#### 517 *Rock-Sand hybridization and genotyping*

518 Two rock-sand crosses, one between *Copadichromis borleyi* (CB, sand-dweller sire) and  
519 *Metriaclima zebra* (MZ, rock-dweller dam) and another between *Mchenga conophoros*  
520 (MC, sand- sire) and *Petrotilapia sp.* 'thick bar' (PT, rock- dam), were artificially generated  
521 by taking the eggs from the dam just prior to spawning and mixing with sperm from the  
522 sire. The resultant F<sub>1</sub> were grown in tanks and allowed to spawn normally to generate F<sub>2</sub>.  
523 Several F<sub>2</sub> broods were taken from multiple F<sub>1</sub> females for each cross, a total of 355  
524 individuals for the CB x MZ cross and 608 for the MC x PT cross. The embryos were fixed  
525 at every stage starting at gastrula (stage 9) until early pharyngula (stage 14). The F<sub>2</sub>

526 embryos were RNA-extracted at stage 9. DNA extraction was performed by fixing the  
527 embryos (stage 11-14) in 70% ethanol, then removing the tail from each individual and  
528 extracting the DNA using an extraction kit (Qiagen). Following extraction, the F<sub>2</sub> embryos  
529 were genotyped using custom probes (CAAATCTCCC[C/T]CCGCGGC, Taqman custom  
530 probes, Invitrogen) designed to identify a SNP in *irx1b* using RT-PCR. A subset of the  
531 embryos was also sequenced at a 900 bp interval around the *irx1b* SNP to verify the  
532 custom probes.

533

#### 534 *Quantitative F<sub>2</sub> Analysis*

535 We quantified *irx1b* in F<sub>2</sub> at stage 9 and separated by genotypic class. The 74  
536 heterozygous rock X sand F<sub>2</sub> embryos were dissected to remove most of the yolk and the  
537 total RNA was extracted from each individual using an RNA Extraction Kit (Qiagen).  
538 The amount of mRNA specific to each allele of *irx1b* was quantified by using the RNA-to-  
539 Ct kit (Invitrogen) and the custom probes. The delta Ct for each heterozygote was  
540 generated with the equation,  $2^{(\text{allele from dam} - \text{allele from sire})}$ . We tested the data  
541 with an ANOVA, followed by a Tukey's multiple comparison test to determine significance  
542 between genotype classes.

543

#### 544 *Forebrain and eye measurements*

545 The forebrain and eyes were measured by integrating the area of transverse sections in  
546 embryos of rock- and sand-dweller cichlid species, using previously published methods  
547 <sup>21</sup>. The rock-dweller species included *Cynotilapia afra* (CA, planktivore), *Labeotropheus*  
548 *fuelleborni* (LF, algivore) and *Metriaclima zebra* (MZ, generalist); sand-dweller species  
549 included *Aulonocara jacobfreibergi* (AJ, 'sonar' hunter), *Copadichromis borleyi* (CB,  
550 planktivore) and *Mchenga conophoros* (MC, insectivore/generalist). Embryos from each  
551 species, as well as the F<sub>2</sub> individuals, were measured starting from the earliest the  
552 telencephalon can be differentiated from the forebrain (mid-somitogenesis, stage 12) and  
553 at each subsequent stage until the forebrain has defined prosomeres (early pharyngula,  
554 stage 14) <sup>22</sup>. To keep measurements standardized across stages, all measurements were

555 defined by forebrain morphology at the earliest timepoint (stage 12). The 'eye'  
556 measurement remains consistent at all stages, the 'anterior' measurement includes the  
557 telencephalon and presumptive olfactory bulb, and the 'posterior' measurement includes  
558 the diencephalon and each of its constitutive prosomeres (dorsal and ventral thalamus  
559 and hypothalamus). To facilitate measurements, we used gene expression of *rx3* (for  
560 stage 12 embryos) and *pax6* (stage 13 and 14) to identify the different structures of the  
561 forebrain and eye.

562

### 563 *RNA Extraction and Sequencing, Adult Social Behavior*

564

565 Two adult F<sub>1</sub> hybrid males from rock- x sand- interspecific crosses (Supplementary table  
566 5) were introduced to an assay tank containing 3-5 hybrid females of the same cross with  
567 simulated rock habitat on one side and simulated sand habitat on the other side separated  
568 by empty tank space (Figure 3A). Males were observed over a period of four days for  
569 courtship behavior, either similar to a rock parent (courting females over the rocky parts  
570 of the tank), here designated "social rock" --- or to a sand parent (bower building in the  
571 sand side of the tank while courting females), here designated "social sand." When  
572 courtship behavior of both males was observed, we primed experiments the next morning  
573 by resetting the tanks -- knocking down or filling in the sand bower, and adjusting the rock  
574 habitat. Both males were allowed to exhibit courtship behavior over a period of 90  
575 minutes. Males were sacrificed immediately after this 90-minute period by rapid  
576 decapitation and whole brains were immediately stored in RNAlater (Thermo Fisher Cat#  
577 AM7020) within 20 minutes of decapitation. Two tanks with one F<sub>1</sub> male and 3-5 F<sub>1</sub>  
578 females were also observed. These males exhibited social rock behavior and were  
579 designated as "lone" in the behavior assay; their brains were collected in the exact same  
580 way as the other samples.

581

582 Tissues were frozen in liquid nitrogen, homogenized using a mortar and pestle and placed  
583 in trizol. Following standard chloroform extraction, RNeasy mini columns (Qiagen Cat  
584 No./ID: 74104) were utilized to purify RNA for sequencing. Total RNA was quantified

585 using Qubit (Molecular Probes) and quality analyzed using the Agilent 2100 Bioanalyzer  
586 System for RNA library preparation. RNA input was normalized to 1 µg and libraries were  
587 prepared using the TruSeq Stranded mRNA Sample Prep Kit (Illumina- Kit A). Libraries  
588 were again quantified, quality assessed, and normalized for sequencing on the HiSeq  
589 2500 Illumina Sequencing System (Georgia Tech Genomics Core, standard practices).  
590 Experimental design and raw files can be accessed on the NCBI Gene Expression  
591 Omnibus database under the accession number GSE122500.

592

### 593 *Differential Gene Expression Analysis*

594 Raw sequence reads from whole brain transcriptomes were quality controlled using the  
595 NGS QC Toolkit <sup>74</sup>. Raw reads with an average PHRED quality score below 20 were  
596 filtered out. Filtered reads were also trimmed of low-quality bases at the 3' end. High  
597 quality sequence reads were aligned to the *M.zebra* reference genome  
598 MZ\_UMD2a<sup>30</sup> using TopHat v2.0.9 <sup>75</sup>. On average, across all samples, over 95% of reads  
599 mapped to the reference genome. The resulting TopHat2 output bam files were sorted  
600 and converted to sam files using samtools v0.19 <sup>76</sup>. Sorted sam files were used as input  
601 for the HTSeq-count v0.6.1 program to obtain fragment counts for each locus <sup>77</sup>.  
602 Fragment counts were scale-normalized across all samples using the calcNormFactors  
603 function in the edgeR package v3.6.8 <sup>78</sup>. Relative consistency among replicates and  
604 samples was determined via the Multidimensional scaling (MDS) feature within the edgeR  
605 package in R. The native R function *hclust(dist)* used to cluster samples. Scale-  
606 normalized fragment counts were converted into log<sub>2</sub> counts per million reads mapped  
607 (cpm) with precision weights using voom and fit to a linear model using limma v3.20.9 <sup>79</sup>.  
608 Pairwise contrasts were constructed between socially rock and socially sand samples.  
609 After correcting for multiple comparisons using the Benjamini-Hochberg method <sup>80</sup>, genes  
610 were considered differentially expressed between socially rock and socially sand samples  
611 if they exhibited both a fold change  $\geq 2$  and  $P_{adj} < 0.05$ . Using Treefam based mapping <sup>61</sup>  
612 the cichlid gene names were converted to human analogs and functional enrichment was  
613 determined using the TOPPFUN web-browser <sup>62</sup>.

614 *Ethics Statement*

615 All of the animals were handled according to methods approved by the Georgia Tech  
616 Institutional Animal Care and Use Committee (IACUC) protocols (permit A100029) with  
617 every effort made to minimize suffering and is in compliance with the ARRIVE guidelines.

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- 625 1 Boyle, E. A., Li, Y. I. & Pritchard, J. K. An Expanded View of Complex Traits: From  
626 Polygenic to Omnigenic. *Cell* **169**, 1177-1186, doi:10.1016/j.cell.2017.05.038 (2017).
- 627 2 Fields, C., Adams, M. D., White, O. & Venter, J. C. How many genes in the human  
628 genome? *Nature Genetics* **7**, 345-346, doi:10.1038/ng0794-345 (1994).
- 629 3 Maurano, M. T. *et al.* Systematic localization of common disease-associated variation in  
630 regulatory DNA. *Science*, 9718-9723 (2012).
- 631 4 Degner, J. F. *et al.* DNase I sensitivity QTLs are a major determinant of human expression  
632 variation. *Nature* **482**, 390 (2012).
- 633 5 Streelman, J., Peichel, C. L. & Parichy, D. Developmental genetics of adaptation in fishes:  
634 the case for novelty. *Annu. Rev. Ecol. Evol. Syst.* **38**, 655-681 (2007).
- 635 6 Colosimo, P. F. *et al.* The genetic architecture of parallel armor plate reduction in  
636 threespine sticklebacks. *Plos Biol* **2**, E109, doi:10.1371/journal.pbio.0020109 (2004).
- 637 7 Kratochwil, C. F. *et al.* Agouti-related peptide 2 facilitates convergent evolution of stripe  
638 patterns across cichlid fish radiations. *Science* **362**, 457-460, doi:10.1126/science.aao6809  
639 (2018).
- 640 8 Albertson, R. C., Streelman, J. T., Kocher, T. D. & Yelick, P. C. Integration and evolution  
641 of the cichlid mandible: The molecular basis of alternate feeding strategies. *P Natl Acad  
642 Sci USA* **102**, 16287-16292, doi:DOI 10.1073/pnas.0506649102 (2005).
- 643 9 Shapiro, M. D. *et al.* Genomic diversity and evolution of the head crest in the rock pigeon.  
644 *Science* **339**, 1063-1067, doi:10.1126/science.1230422 (2013).
- 645 10 Lamichhaney, S. *et al.* Evolution of Darwin's finches and their beaks revealed by genome  
646 sequencing. *Nature* **518**, 371-375 (2015).
- 647 11 Okhovat, M., Berrio, A., Wallace, G., Ophir, A. G. & Phelps, S. M. Sexual fidelity trade-  
648 offs promote regulatory variation in the prairie vole brain. *Science* **350**, 1371-1374 (2015).
- 649 12 Bendesky, A. *et al.* The genetic basis of parental care evolution in monogamous mice.  
650 *Nature* **544**, 434 (2017).
- 651 13 Pfenning, A. R. *et al.* Convergent transcriptional specializations in the brains of humans  
652 and song-learning birds. *Science* **346**, 1256846 (2014).

- 653 14 Greenwood, A. K., Wark, A. R., Yoshida, K. & Peichel, C. L. Genetic and neural  
654 modularity underlie the evolution of schooling behavior in threespine sticklebacks. *Curr*  
655 *Biol* **23**, 1884-1888, doi:10.1016/j.cub.2013.07.058 (2013).
- 656 15 Kocher, T. D. Adaptive evolution and explosive speciation: The cichlid fish model. *Nat*  
657 *Rev Genet* **5**, 288-298 (2004).
- 658 16 Loh, Y. H. E. *et al.* Comparative analysis reveals signatures of differentiation amid  
659 genomic polymorphism in Lake Malawi cichlids. *Genome Biology* **9** (2008).
- 660 17 Loh, Y. H. E. *et al.* Origins of Shared Genetic Variation in African Cichlids. *Mol Biol Evol*  
661 **30**, 906-917 (2013).
- 662 18 Streelman, J. T., Albertson, R. C. & Kocher, T. D. Genome mapping of the orange blotch  
663 colour pattern in cichlid fishes. *Mol Ecol* **12**, 2465-2471, doi:10.1046/j.1365-  
664 294x.2003.01920.x (2003).
- 665 19 Roberts, R. B., Ser, J. R. & Kocher, T. D. Sexual conflict resolved by invasion of a novel  
666 sex determiner in Lake Malawi cichlid fishes. *Science* **326**, 998-1001 (2009).
- 667 20 Parnell, N. F. & Streelman, J. T. Genetic interactions controlling sex and color establish  
668 the potential for sexual conflict in Lake Malawi cichlid fishes. *Heredity* **110**, 239-246  
669 (2013).
- 670 21 Sylvester, J. B. *et al.* Competing signals drive telencephalon diversity. *Nat Commun* **4**, 4  
671 (2013).
- 672 22 Sylvester, J. B. *et al.* Brain diversity evolves via differences in patterning. *Proceedings of*  
673 *the National Academy of Sciences* **107**, 9718-9723 (2010).
- 674 23 Baran, N. M. & Streelman, J. T. Ecotype differences in aggression, neural activity and  
675 behaviorally relevant gene expression in cichlid fish. *Genes Brain Behav* **19**, e12657,  
676 doi:10.1111/gbb.12657 (2020).
- 677 24 York, R. A. *et al.* Behavior-dependent cis regulation reveals genes and pathways associated  
678 with bower building in cichlid fishes. *Proceedings of the National Academy of Sciences*  
679 **115**, E11081-e11090, doi:10.1073/pnas.1810140115 (2018).
- 680 25 Johnson, Z. V. *et al.* Exploratory behaviour is associated with microhabitat and  
681 evolutionary radiation in Lake Malawi cichlids. *Animal Behaviour* **160**, 121-134,  
682 doi:https://doi.org/10.1016/j.anbehav.2019.11.006 (2020).
- 683 26 Fraser, G. J. *et al.* An ancient gene network is co-opted for teeth on old and new jaws. *Plos*  
684 *Biol* **7**, e1000031 (2009).
- 685 27 Streelman, J. T. & Danley, P. D. The stages of vertebrate evolutionary radiation. *Trends*  
686 *Ecol Evol* **18**, 126-131 (2003).

- 687 28 Malinsky, M. *et al.* Whole-genome sequences of Malawi cichlids reveal multiple radiations  
688 interconnected by gene flow. *Nat Ecol Evol* **2**, 1940-1955, doi:10.1038/s41559-018-0717-  
689 x (2018).
- 690 29 McKaye, K. R., Louda, S. M. & Jay R. Stauffer, J. Bower Size and Male Reproductive  
691 Success in a Cichlid Fish Lek. *The American Naturalist* **135**, 597-613, doi:10.1086/285064  
692 (1990).
- 693 30 Conte, M. A. *et al.* Chromosome-scale assemblies reveal the structural evolution of African  
694 cichlid genomes. *GigaScience* **8**, doi:10.1093/gigascience/giz030 (2019).
- 695 31 Brawand, D. *et al.* The genomic substrate for adaptive radiation in African cichlid fish.  
696 *Nature* **513**, 375-381 (2014).
- 697 32 Ben-Ari Fuchs, S. *et al.* GeneAnalytics: An Integrative Gene Set Analysis Tool for Next  
698 Generation Sequencing, RNAseq and Microarray Data. *Omics* **20**, 139-151 (2016).
- 699 33 Hawrylycz, M. J. *et al.* An anatomically comprehensive atlas of the adult human brain  
700 transcriptome. *Nature* **489**, 391-399, doi:10.1038/nature11405 (2012).
- 701 34 Adhikari, A. *et al.* Basomedial amygdala mediates top-down control of anxiety and fear.  
702 *Nature* **527**, 179-185, doi:10.1038/nature15698 (2015).
- 703 35 Mesquita, L. T. *et al.* New insights on amygdala: Basomedial amygdala regulates the  
704 physiological response to social novelty. *Neuroscience* **330**, 181-190,  
705 doi:https://doi.org/10.1016/j.neuroscience.2016.05.053 (2016).
- 706 36 Carvalho, F. R., Nóbrega, C. D. R. & Martins, A. T. Mapping gene expression in social  
707 anxiety reveals the main brain structures involved in this disorder. *Behavioural Brain*  
708 *Research* **394**, 112808, doi:https://doi.org/10.1016/j.bbr.2020.112808 (2020).
- 709 37 Abrahams, B. S. *et al.* SFARI Gene 2.0: a community-driven knowledgebase for the autism  
710 spectrum disorders (ASDs). *Mol Autism* **4**, 36, doi:10.1186/2040-2392-4-36 (2013).
- 711 38 Piñero, J. *et al.* DisGeNET: a comprehensive platform integrating information on human  
712 disease-associated genes and variants. *Nucleic Acids Research* **45**, D833-D839,  
713 doi:10.1093/nar/gkw943 (2017).
- 714 39 Attanasio, C. *et al.* Fine Tuning of Craniofacial Morphology by Distant-Acting Enhancers.  
715 *Science* **342** (2013).
- 716 40 Rada-Iglesias, A. *et al.* Epigenomic Annotation of Enhancers Predicts Transcriptional  
717 Regulators of Human Neural Crest. *Cell Stem Cell* **11**, 633-648 (2012).
- 718 41 Simoes-Costa, M. & Bronner, M. E. Establishing neural crest identity: a gene regulatory  
719 recipe. *Development* **142**, 242-257, doi:10.1242/dev.105445 (2015).

- 720 42 Roberts, R. B., Hu, Y., Albertson, R. C. & Kocher, T. D. Craniofacial divergence and  
721 ongoing adaptation via the hedgehog pathway. *Proc Natl Acad Sci U S A* **108**, 13194-  
722 13199, doi:10.1073/pnas.1018456108 (2011).
- 723 43 Cavodeassi, F., Modolell, J. & Gómez-Skarmeta, J. L. The Iroquois family of genes: from  
724 body building to neural patterning. *Development* **128**, 2847-2855 (2001).
- 725 44 Bielen, H. & Houart, C. BMP signaling protects telencephalic fate by repressing eye  
726 identity and its Cxcr4-dependent morphogenesis. *Developmental cell* **23**, 812-822 (2012).
- 727 45 White, D. E. *et al.* Quantitative multivariate analysis of dynamic multicellular morphogenic  
728 trajectories. *Integrative Biology* **7**, 825-833, doi:10.1039/c5ib00072f (2015).
- 729 46 Volff, J. N., Korting, C. & Schartl, M. Multiple lineages of the non-LTR retrotransposon  
730 Rex1 with varying success in invading fish genomes. *Mol Biol Evol* **17**, 1673-1684,  
731 doi:10.1093/oxfordjournals.molbev.a026266 (2000).
- 732 47 Olton, D. S., Becker, J. T. & Handelmann, G. E. Hippocampus, space, and memory.  
733 *Behavioral and Brain sciences* **2**, 313-322 (1979).
- 734 48 Hitti, F. L. & Siegelbaum, S. A. The hippocampal CA2 region is essential for social  
735 memory. *Nature* **508**, 88-92 (2014).
- 736 49 Chiang, M.-C., Huang, A. J., Wintzer, M. E., Ohshima, T. & McHugh, T. J. A role for CA3  
737 in social recognition memory. *Behavioural brain research* **354**, 22-30 (2018).
- 738 50 Zou, D. *et al.* DREADD in parvalbumin interneurons of the dentate gyrus modulates  
739 anxiety, social interaction and memory extinction. *Current Molecular Medicine* **16**, 91-102  
740 (2016).
- 741 51 O'Connell, L. A. & Hofmann, H. A. Evolution of a vertebrate social decision-making  
742 network. *Science* **336**, 1154-1157 (2012).
- 743 52 McLean, C. Y. *et al.* Human-specific loss of regulatory DNA and the evolution of human-  
744 specific traits. *Nature* **471**, 216-219, doi:10.1038/nature09774 (2011).
- 745 53 Finlay, B. L. & Darlington, R. B. Linked regularities in the development and evolution of  
746 mammalian brains. *Science* **268**, 1578-1584, doi:10.1126/science.7777856 (1995).
- 747 54 Brenner, S. Genetics of *Caenorhabditis-Elegans*. *Genetics* **77**, 71-94 (1974).
- 748 55 Fernald, R. D. & Maruska, K. P. Social information changes the brain. *Proc Natl Acad Sci*  
749 *U S A* **109 Suppl 2**, 17194-17199, doi:10.1073/pnas.1202552109 (2012).
- 750 56 Li, H. & Durbin, R. Fast and accurate short read alignment with Burrows-Wheeler  
751 transform. *Bioinformatics* **25**, 1754-1760, doi:10.1093/bioinformatics/btp324 (2009).

- 752 57 Van der Auwera, G. A. *et al.* From FastQ data to high confidence variant calls: the Genome  
753 Analysis Toolkit best practices pipeline. *Current protocols in bioinformatics* **43**, 11.10.11-  
754 33, doi:10.1002/0471250953.bi1110s43 (2013).
- 755 58 Lee, T. H., Guo, H., Wang, X. Y., Kim, C. & Paterson, A. H. SNPhylo: a pipeline to  
756 construct a phylogenetic tree from huge SNP data. *Bmc Genomics* **15** (2014).
- 757 59 Danecek, P. *et al.* The variant call format and VCFtools. *Bioinformatics* **27**, 2156-2158  
758 (2011).
- 759 60 Cingolani, P. *et al.* A program for annotating and predicting the effects of single nucleotide  
760 polymorphisms, SnpEff: SNPs in the genome of *Drosophila melanogaster* strain w(1118);  
761 iso-2; iso-3. *Fly* **6**, 80-92 (2012).
- 762 61 Ramakrishnan Varadarajan, A., Mopuri, R., Streelman, J. T. & McGrath, P. T. Genome-  
763 wide protein phylogenies for four African cichlid species. *BMC Evol Biol* **18**, 1,  
764 doi:10.1186/s12862-017-1072-2 (2018).
- 765 62 Chen, J., Bardes, E. E., Aronow, B. J. & Jegga, A. G. ToppGene Suite for gene list  
766 enrichment analysis and candidate gene prioritization. *Nucleic Acids Res* **37**, W305-W311,  
767 doi:10.1093/nar/gkp427 (2009).
- 768 63 Harris, R. S. *Improved Pairwise Alignment of Genomic DNA* Doctor of Philosophy thesis,  
769 The Pennsylvania State University, (2007).
- 770 64 Blanchette, M. *et al.* Aligning multiple genomic sequences with the threaded blockset  
771 aligner. *Genome research* **14**, 708-715 (2004).
- 772 65 Siepel, A. *et al.* Evolutionarily conserved elements in vertebrate, insect, worm, and yeast  
773 genomes. *Genome Res* **15**, 1034-1050, doi:10.1101/gr.3715005 (2005).
- 774 66 Siepel, A. & Haussler, D. Phylogenetic estimation of context-dependent substitution rates  
775 by maximum likelihood. *Mol Biol Evol* **21**, 468-488 (2004).
- 776 67 Kent, W. J. *et al.* The human genome browser at UCSC. *Genome research* **12**, 996-1006  
777 (2002).
- 778 68 Grote, S., Prufer, K., Kelso, J. & Dannemann, M. ABAEnrichment: an R package to test  
779 for gene set expression enrichment in the adult and developing human brain.  
780 *Bioinformatics* **32**, 3201-3203, doi:10.1093/bioinformatics/btw392 (2016).
- 781 69 Dougherty, J. D., Schmidt, E. F., Nakajima, M. & Heintz, N. Analytical approaches to  
782 RNA profiling data for the identification of genes enriched in specific cells. *Nucleic Acids*  
783 *Res* **38**, 4218-4230, doi:10.1093/nar/gkq130 (2010).
- 784 70 Xu, X., Wells, A. B., O'Brien, D. R., Nehorai, A. & Dougherty, J. D. Cell type-specific  
785 expression analysis to identify putative cellular mechanisms for neurogenetic disorders. *J*  
786 *Neurosci* **34**, 1420-1431, doi:10.1523/JNEUROSCI.4488-13.2014 (2014).

787 71 Murata, Y. *et al.* Allometric growth of the trunk leads to the rostral shift of the pelvic fin  
788 in teleost fishes. *Dev Biol* **347**, 236-245, doi:10.1016/j.ydbio.2010.07.034 (2010).

789 72 Fraser, G. J., Bloomquist, R. F. & Streelman, J. T. A periodic pattern generator for dental  
790 diversity. *Bmc Biol* **6** (2008).

791 73 Tucker, J. A., Mintzer, K. A. & Mullins, M. C. The BMP signaling gradient patterns  
792 dorsoventral tissues in a temporally progressive manner along the anteroposterior axis. *Dev*  
793 *Cell* **14**, 108-119, doi:10.1016/j.devcel.2007.11.004 (2008).

794 74 Patel, R. K. & Jain, M. NGS QC Toolkit: A Toolkit for Quality Control of Next Generation  
795 Sequencing Data. *PLoS One* **7**, 7 (2012).

796 75 Trapnell, C., Pachter, L. & Salzberg, S. L. TopHat: discovering splice junctions with RNA-  
797 Seq. *Bioinformatics* **25**, 1105-1111 (2009).

798 76 Li, H. *et al.* The Sequence Alignment/Map format and SAMtools. *Bioinformatics* **25**, 2078-  
799 2079 (2009).

800 77 Anders, S., Pyl, P. T. & Huber, W. HTSeq--a Python framework to work with high-  
801 throughput sequencing data. *Bioinformatics* **31**, 166-169 (2015).

802 78 Robinson, M. D., McCarthy, D. J. & Smyth, G. K. edgeR: a Bioconductor package for  
803 differential expression analysis of digital gene expression data. *Bioinformatics* **26**, 139-140  
804 (2010).

805 79 Ritchie, M. E. *et al.* limma powers differential expression analyses for RNA-sequencing  
806 and microarray studies. *Nucleic Acids Res* **43**, e47 (2015).

807 80 Hochberg, Y. & Benjamini, Y. More powerful procedures for multiple significance testing.  
808 *Stat Med* **9**, 811-818 (1990).

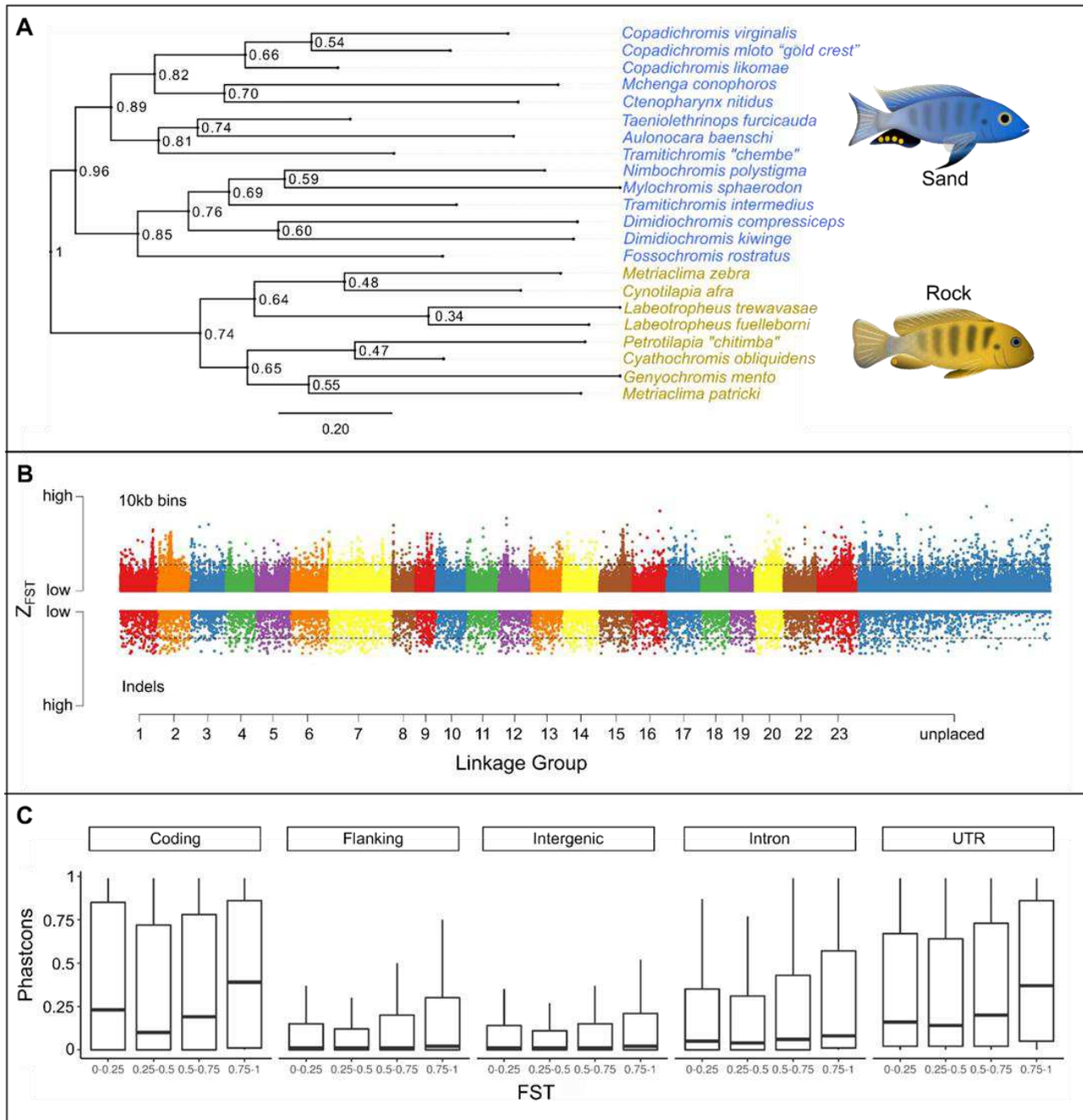
809 81 Bloomquist, R. F., Fowler, T. E., Sylvester, J. B., Miro, R. J. & Streelman, J. T. A  
810 compendium of developmental gene expression in Lake Malawi cichlid fishes. *BMC Dev*  
811 *Biol* **17**, 3, doi:10.1186/s12861-017-0146-0 (2017).

812 82 Bonkowsky, J. L. *et al.* Domain-specific regulation of foxP2 CNS expression by lef1. *BMC*  
813 *Dev Biol* **8**, 103, doi:10.1186/1471-213X-8-103 (2008).

814 83 Shi, Z. *et al.* miR-9 and miR-140-5p target FoxP2 and are regulated as a function of the  
815 social context of singing behavior in zebra finches. *J Neurosci* **33**, 16510-16521,  
816 doi:10.1523/JNEUROSCI.0838-13.2013 (2013).

817  
818

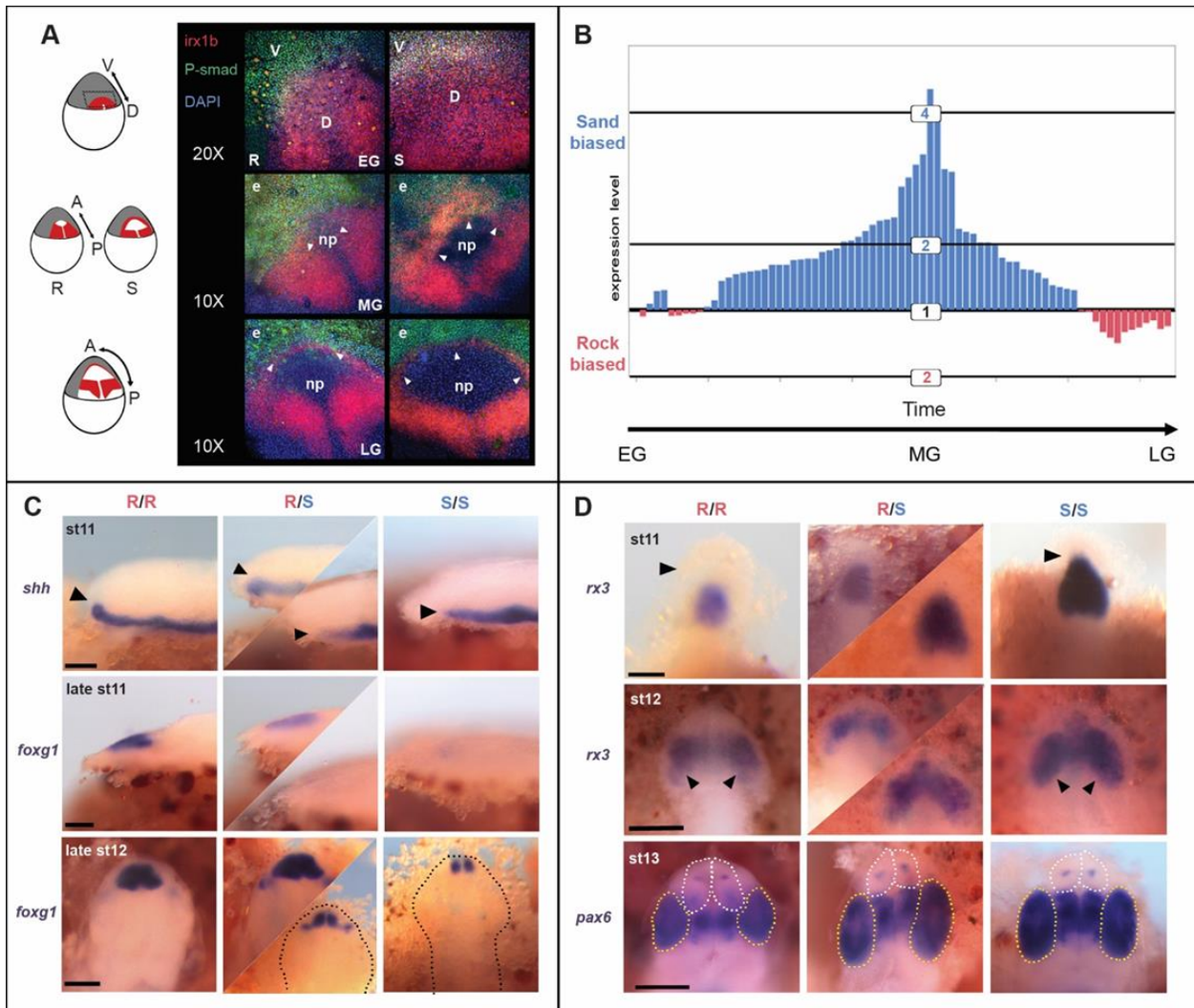
819 **Author Contributions:** CP, JBS, KA and JTS designed research; CP, JBS, KA, MWN, KP, RFB  
820 performed research; CP, KA, MM, ZVJ and PTM analyzed data; CP, JBS, ZVJ and JTS wrote the paper.  
821  
822 **Competing Interest Statement:** The Authors declare no competing interests.  
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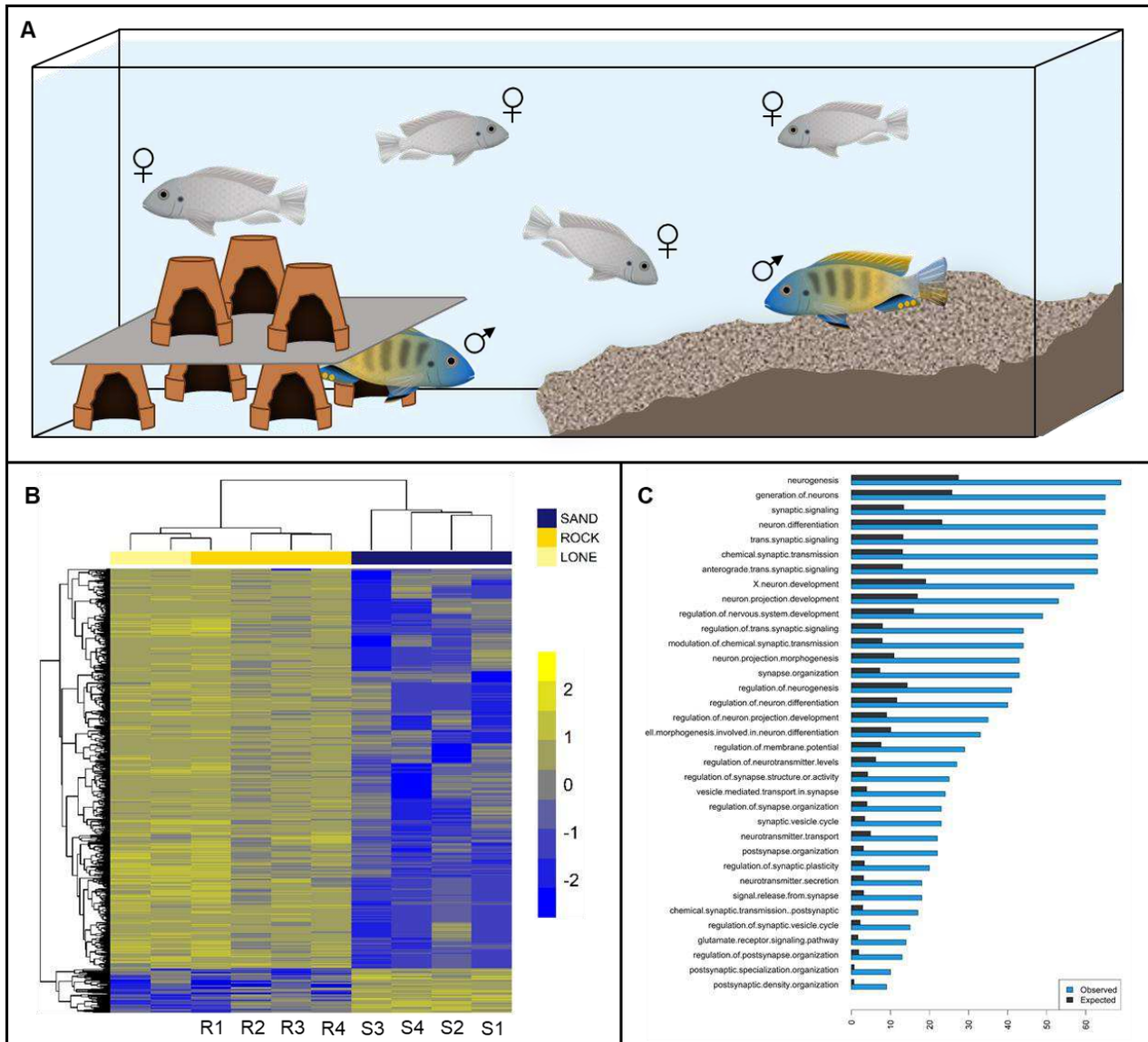
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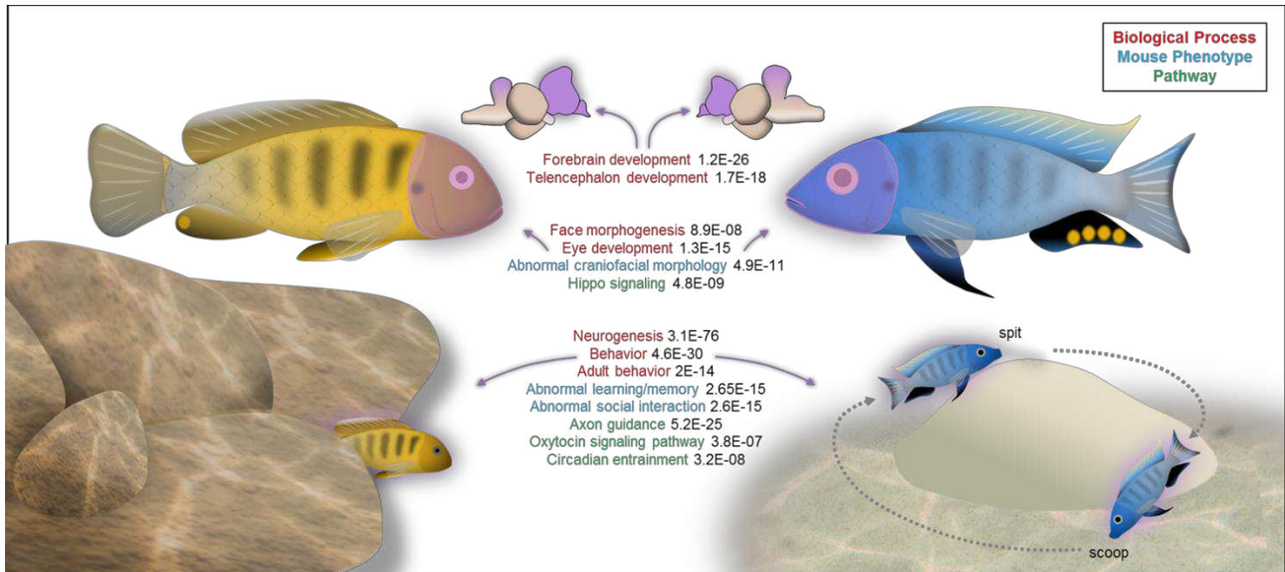


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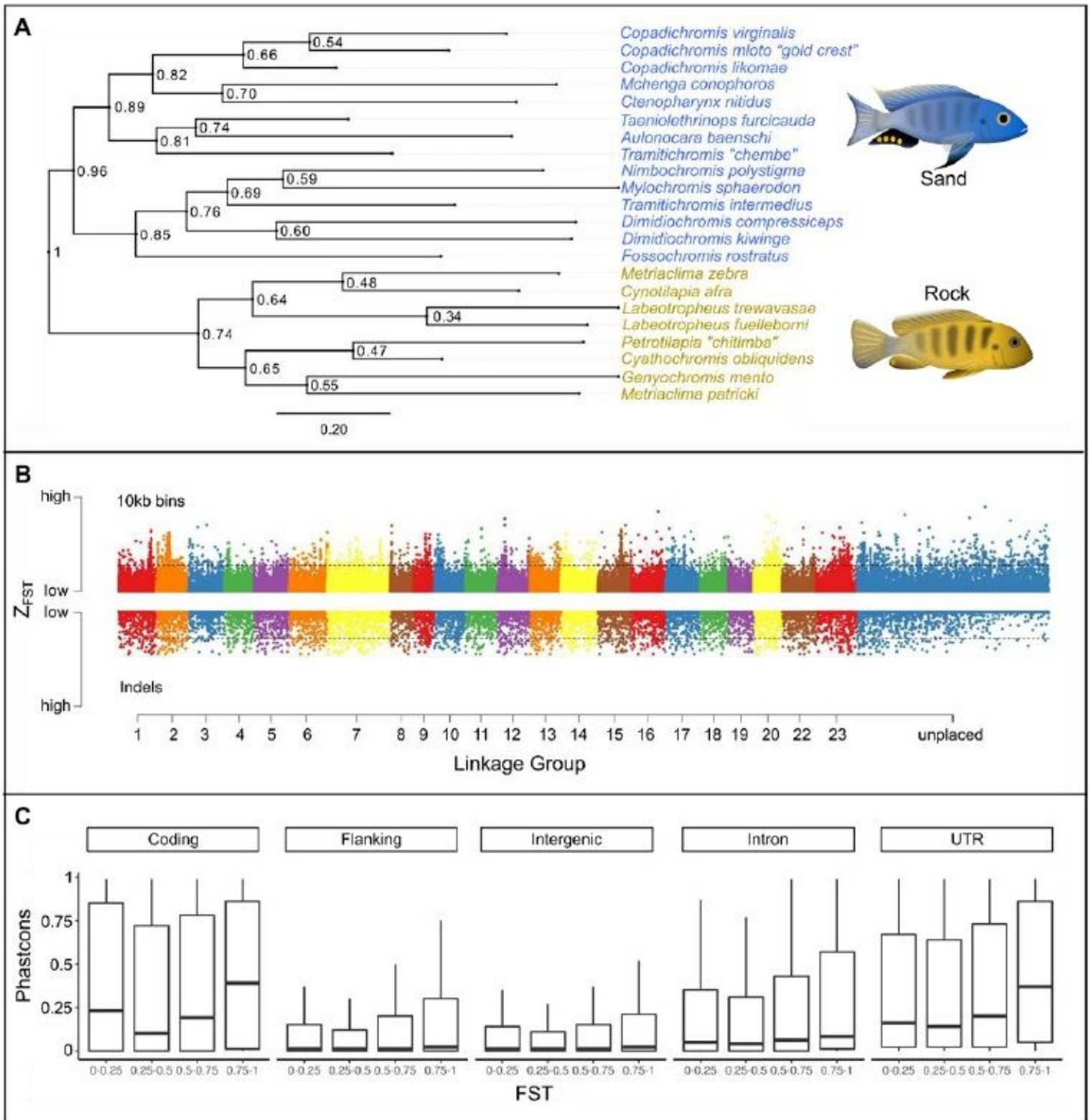
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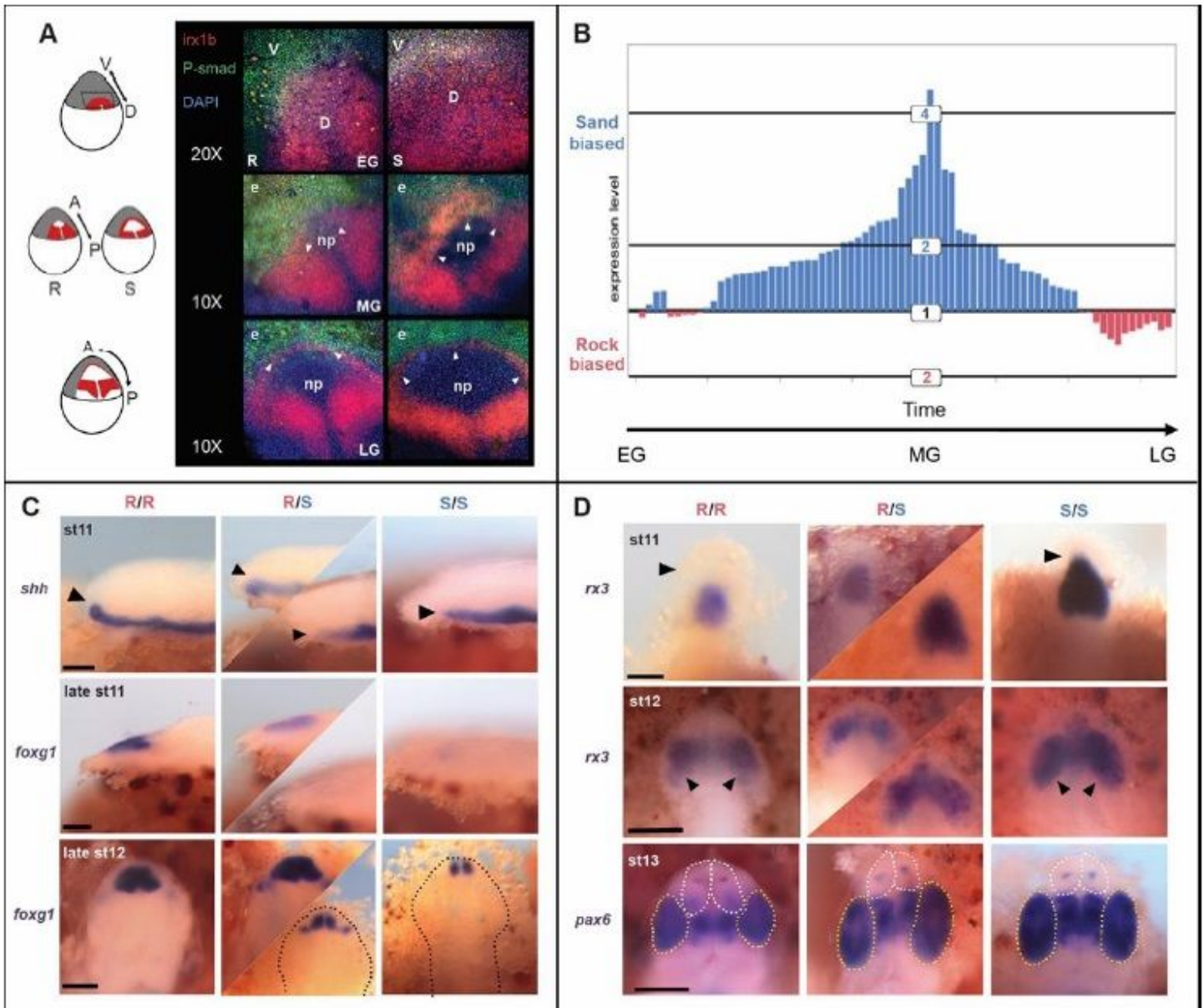
# Figures



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**Figure 2**

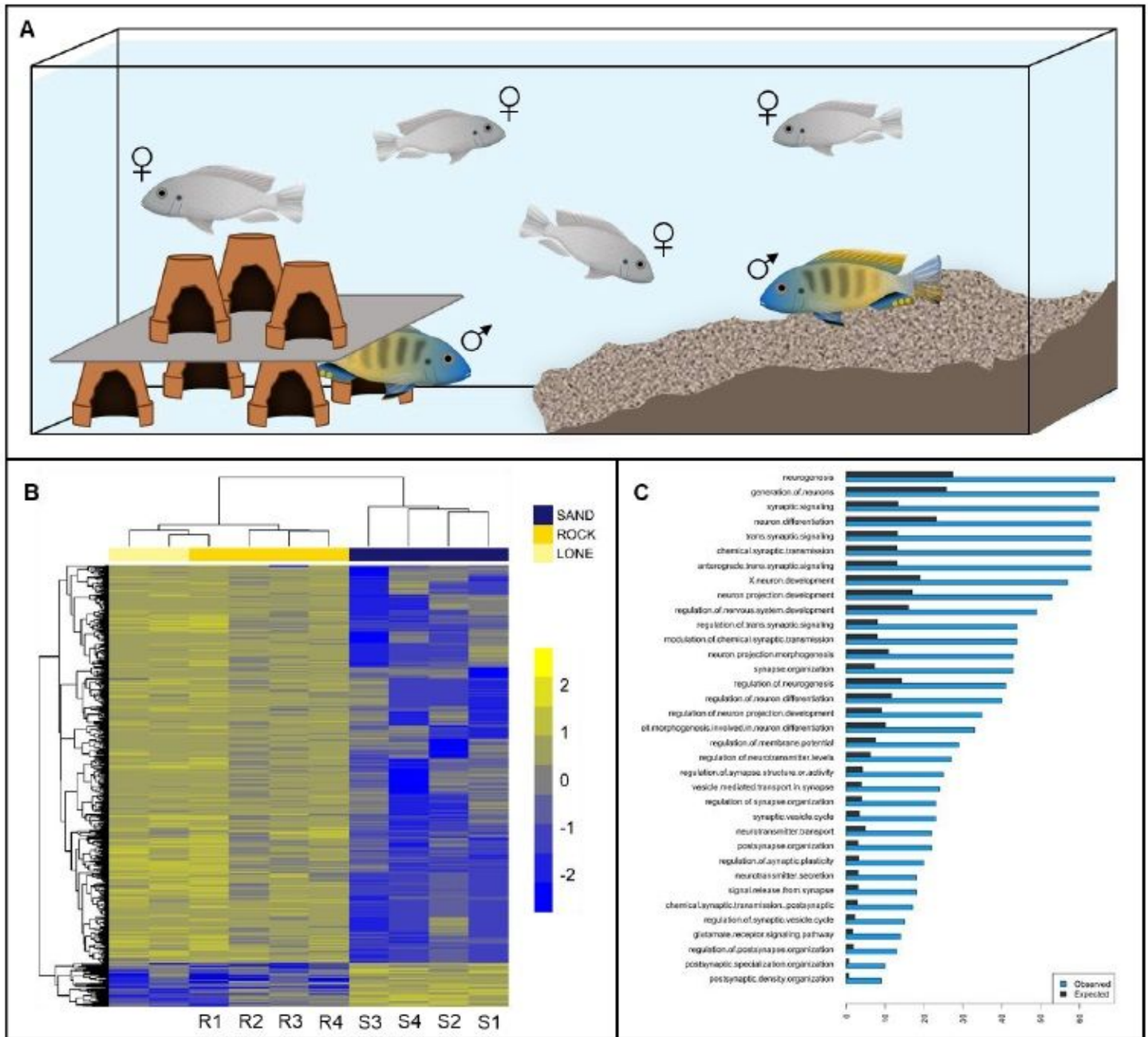
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(C) *in situ* hybridization of *shh* and *foxg1*, during neurula stages, showing development of the telencephalon in rock- X sand- F2 embryos, indexed for *irx1b* genotype. F2 individuals homozygous for rock- *irx1b* alleles (R/R) show a more dorsal progression of *shh* expression (black arrowheads), an earlier and a larger expression domain of the telencephalon marker *foxg1*. The top two rows are lateral views; bottom row is a dorsal view. Dotted lines demarcate the outline of the embryo in dorsal view. Heterozygous individuals exhibit greater variation in expression domains (middle columns), indicating that genetic factors other than variants in *irx1b* contribute to this phenotype.

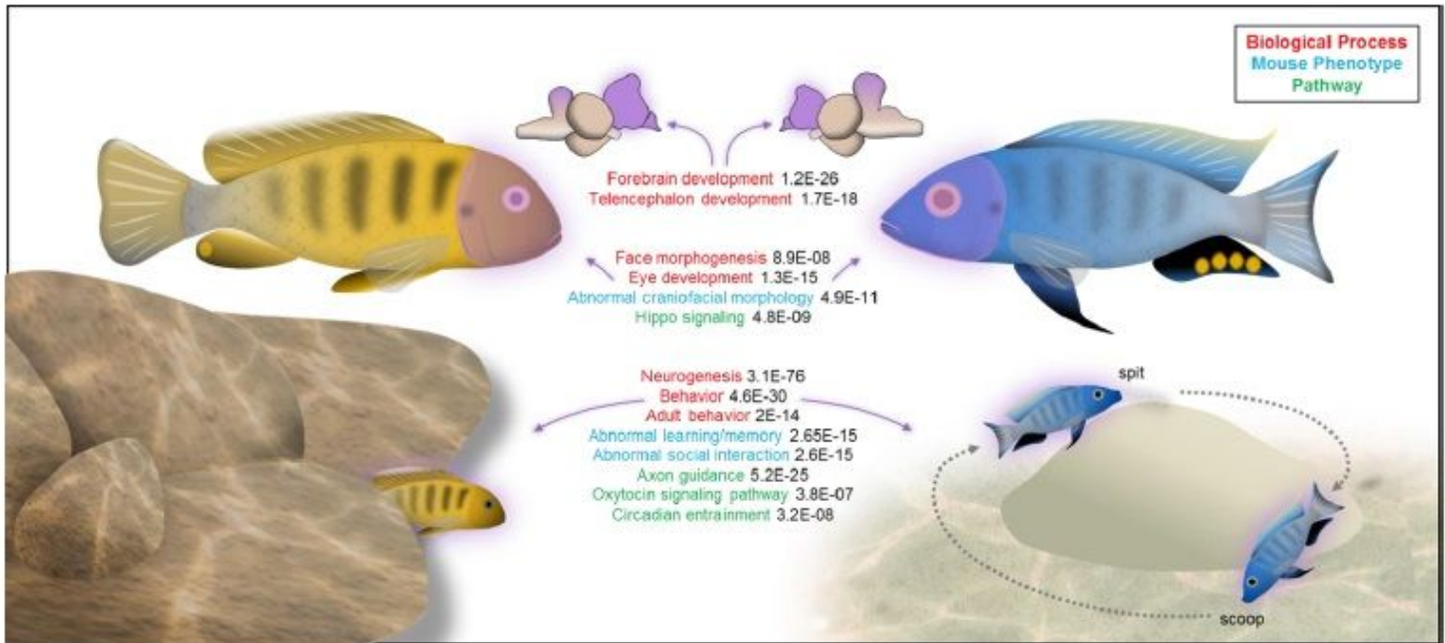
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**Figure 4**

Genome-enabled discovery of evolutionary change in morphology and behavior. | Summary cartoon synthesizing significant enrichment categories that differentiate the genomes of rock- vs. sand-dwelling Malawi cichlids. Strong and consistent enrichment of craniofacial, neural and behavioral categories motivated follow-on experiments in early brain development (Figure 2) and adult social behavior (Figure 3).

## Supplementary Files

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- [SupplementaryTable4rocksandversuspitcastle.xlsx](#)
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