Discrete genetic modules are responsible for complex burrow evolution in *Peromyscus* mice

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Relative to morphological traits, we know little about how genetics influence the evolution of complex behavioural differences in nature\(^1\). It is unclear how the environment influences natural variation in heritable behaviour\(^2\), and whether complex behavioural differences evolve through few genetic changes, each affecting many aspects of behaviour, or through the accumulation of several genetic changes that, when combined, give rise to behavioural complexity\(^3\). Here we show that in nature, oldfield mice (*Peromyscus polionotus*) build complex burrows with long entrance and escape tunnels, and that burrow length is consistent across populations, although burrow depth varies with soil composition. This burrow architecture is in contrast with the small, simple burrows of its sister species, deer mice (*P. maniculatus*). When investigated under laboratory conditions, both species recapitulate their natural burrowing behaviour. Genetic crosses between the two species reveal that the derived burrows of oldfield mice are dominant and evolved through the addition of multiple genetic changes. In burrows built by first-generation backcross mice, entrance-tunnel length and the presence of an escape tunnel can be uncoupled, suggesting that these traits are modular. Quantitative trait locus analysis also indicates that tunnel length segregates as a complex trait, affected by at least three independent genetic regions, whereas the presence of an escape tunnel is associated with only a single locus. Together, these results suggest that complex behaviours—in this case, a classic ‘extended phenotype’\(^4\)—can evolve through multiple genetic changes each affecting distinct behaviour modules.

Animal architectures, such as beehives, bird nests, spider webs, termite mounds and rodent burrows, are remarkably diverse traits that can evolve through natural selection. Despite their great diversity, these extended phenotypes have similarities: they seem to be constructed through largely unlearned motor patterns; they are often consistent within a species (or population); and, when architectures differ, these differences reflect important fitness-related functions in the wild\(^5\). Thus, genetic changes are predicted to contribute to the evolution of different architectures, even between closely related species; however, biologists have long questioned how genetic changes can lead to the evolution of distinct behaviours\(^6\). Because animal architectures can be precisely measured—similarly to morphological traits, which have been successfully dissected genetically\(^7,8\)—these are excellent traits for the genetic analysis of behaviour.

Many species of *Peromyscus* mice build burrows, which were initially described by natural historians working in the field\(^9-14\). However, these burrowing behaviours can also be studied in the laboratory\(^5,14\), and our previous work showed that burrow differences among *Peromyscus* mice have a strong genetic component\(^15\). In particular, *P. polionotus* and *P. maniculatus* are recently diverged and interfertile sister species that build distinct burrows. *P. polionotus* is an open-field specialist and is restricted to the southeastern United States (Fig. 1a and Supplementary Fig. 1a), whereas *P. maniculatus* is a generalist, which inhabits prairie and forest habitats across much of North America (Supplementary Fig. 1c). Whereas most *Peromyscus* species build small, single-tunnel burrows (Supplementary Fig. 1d) or no burrows at all, *P. polionotus* construct unique burrows characterized by a long entrance tunnel that leads into a nest cavity, and a secondary tunnel that emanates from the nest and terminates just below the soil surface (Fig. 1b and Supplementary Fig. 1b)\(^9-11\). This secondary tunnel may serve several functions\(^11\), most notably its use as an escape tunnel when predators (for example, snakes) invade the entrance tunnel (see Supplementary Video). Overall, the complex burrow architecture in *P. polionotus* is derived\(^12\), and probably associated with adaptation to living in an exposed, open habitat\(^5\).

To examine natural variation in burrow complexity, we measured burrow dimensions and soil composition (that is, percentage silt, sand and clay) across the range of *P. polionotus* (Fig. 1a and Supplementary Figure 1d). The correlation between silt composition of soil and burrow depth (asterisk, Correlation between silt composition of soil and burrow depth (asterisk, $r = -0.48$, $P = 0.01$). Each point represents a burrow, and shapes represent the eight different sampling sites (from a). The number of burrows measured at each site is shown in parentheses.

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Figure 1 | Natural variation in *P. polionotus* burrows. a, Sampling of burrows at eight sites in the southeastern United States from across the range of *P. polionotus* (grey area). Average percentage of soil silt at each sampling site is provided. b, Diagram of a typical *P. polionotus* burrow showing the measures for entrance-tunnel length, total length and burrow depth, as well as a typical escape tunnel. c, Variation in total burrow length among sites (mean ± range; from a), which are ordered by increasing percentage of silt (left to right). d, Correlation between silt composition of soil and burrow depth (asterisk indicates Spearman correlation, $r = -0.48$, $S = 6632$, $P = 0.01$). Each point represents a burrow, and shapes represent the eight different sampling sites (from a). The number of burrows measured at each site is shown in parentheses.
Table 1; see Methods for details). Burrows generally had two long tunnels—an entrance and an escape tunnel (Fig. 1b)—and a mean total length of 181 cm (standard deviation (s.d.) of ±53). Of the three soil variables measured, the percentage of silt was the most likely to influence soil compaction and thereby influence burrow dimensions18. Despite large differences in the percentage of silt among locations (Fig. 1c; Kruskal–Wallis (K–W) test, chi-squared = 19.42, degrees of freedom (df) = 7, P = 0.01), burrow length did not differ significantly among locations (K–W test, chi-squared = 6.59, df = 7, P = 0.47), nor did burrow length correlate with the percentage of silt measured at each burrow (Spearman test, rho = −0.16, S = 5,226, P = 0.39). In contrast, there was a significant, negative correlation between the percentage of silt in the soil at each burrow and burrow depth (Fig. 1d; Spearman test, rho = −0.48, S = 6,632, P = 0.01), suggesting that the burrows are shallower, but not shorter, when constructed in compact, silty soil11 (Fig. 1d). Overall, natural burrow shape and length are remarkably conserved in wild P. polionotus, yet variation in soil composition affects burrow depth. We therefore focused our genetic analyses on burrow length.

To measure the genetic component of complex burrowing, we first assayed the burrowing behaviour of P. polionotus and P. maniculatus in a single soil type under controlled laboratory conditions (see Methods). Briefly, each assay involved placing a mouse in a large, sand-filled enclosure for 46 h (two full-night activity periods). Then, we removed the mouse and made a polyurethane cast of the burrow (Supplementary Fig. 1c, d), which we then measured (Fig. 1b). We tested each animal in three consecutive assays. Captive-reared mice, which had never been exposed to sand or allowed to burrow, recapitulated their natural burrowing behaviour in our assays. We found no significant effects of assay number, sex, age, mass or enclosure on total burrow length (data not shown). Statistical comparisons of P. polionotus and P. maniculatus burrows revealed significant differences in total burrow length (Welch’s two-tailed t-test, t = 3.24, df = 20.98, P < 0.01), consistent with previous results15,16. Moreover, total burrow length is composed of two main parts: entrance-tunnel length, which differed significantly between species (Fig. 2b; Welch’s two-tailed t-test, t = 6.72, df = 24.39, P < 5 × 10−3), and escape-tunnel length—only P. polionotus constructed escape tunnels (Fig. 2c). Thus, despite having no previous experience with either sand substrate or our enclosures, these species built consistently distinct burrows in the laboratory.

To determine the inheritance patterns of this behavioural variation, we characterized the genetic architecture of burrowing differences observed between P. polionotus and P. maniculatus. First, we crossed the two species and assayed the burrows of their hybrid offspring (Supplementary Methods and Fig. 2a). First-generation (F1) hybrids built entrance tunnels that were significantly longer (after Bonferroni correction, z = 1.67 × 10−2) than P. maniculatus (Welch’s two-tailed t-test, t = 5.34, df = 20.7, P = 2 × 10−4). But F1 entrance lengths did not differ from the P. polionotus parent (one sample t-test, μ = 14 cm, t = −0.14, df = 12, P = 0.89) or from the P. polionotus population mean (Fig. 2b; Welch’s two-tailed t-test, t = 2.35, df = 22.7, P = 0.03), and all F1 animals constructed escape tunnels (Fig. 2c). Thus, the alleles contributing to burrow size and shape seem to segregate in a dominant fashion.

We next crossed F1 animals with P. maniculatus to create a recombinant backcross (BC) generation (n = 272). BC mice constructed entrance tunnels that varied continuously in length between the parental extremes, but approximately one of eight (n = 36) of the BC mice built P. polionotus-length tunnels (Fig. 2b; >14 cm in length), suggesting that only a few loci are necessary to generate this behaviour. In contrast, half (46%) of the BC mice built escape tunnels (Fig. 2c). This inheritance pattern is consistent with the action of either a single major-effect locus or of multiple loci that interact to create a threshold effect19, such that only some loci need to be co-inherited to cause the expression of a trait. Finally, tunnel lengths and the presence of escape tunnels (that is, tunnel number) are only weakly correlated in BC mice (Supplementary Table 2). Therefore, the complex burrows of P. polionotus comprise at least two separate behavioural modules, one for tunnel length and one for the presence of an escape tunnel.

To identify the chromosomal locations involved and quantify the phenotypic effects of loci that influence these behavioural modules, we used a quantitative trait locus (QTL) mapping approach. We first identified 526 diagnostic single nucleotide polymorphisms (SNPs) using a double-digest restriction-site-associated-DNA (ddRAD) method20, and genotyped the 272 BC mice for which we measured burrowing behaviour (that is, entrance-tunnel length and the presence of an escape tunnel; see Methods for details). Consistent with simple inheritance patterns predicted by burrow phenotypes in hybrid animals, we identified three genomic regions that contributed to variation in entrance-tunnel length, and a single region associated with escape-tunnel construction (Fig. 3a and Supplementary Fig. 5). All four QTLs are unlinked and segregate on separate chromosomes. The entrance-tunnel-length QTLs have similar effect sizes, interact additively (as determined by both a lack of epistasis between loci and graphical comparisons; Fig. 3b and Supplementary Methods), and together explain ~15% of tunnel-length variation (Table 1). Because approximately 24% of the variation in burrow length is likely to have a
 genetic basis (as measured by the repeatability of burrow length in *P. polionotus*)\(^\text{17}\), the QTLs that we report may explain more than half of the genetic variation for this trait (but see ref. 21). Even more notable, each *P. polionotus* allele increases entrance-tunnel length by, on average, 3 cm (Fig. 3b). Furthermore, BC individuals with all three *P. maniculatus* alleles versus those with one *P. polionotus* allele at each locus (that is, heterozygotes) nearly recapitulate the difference in entrance-tunnel length between the two species (that is, the mean phenotype of the parents used in the cross). Although it is possible that each QTL may harbour multiple, possibly interacting, mutations, these results are consistent with only a few genes (as few as three) being needed to explain the continuous tunnel-length variation observed in BC mice.

By contrast, QTL analysis identified only a single locus for the presence of escape tunnels on linkage group 5. Although this QTL explains only 6% of the escape-tunnel variance, BC individuals that inherited one *P. polionotus* allele at this locus were 30% more likely on average to build an escape tunnel than individuals with only *P. maniculatus* alleles (Fig. 3c). Given that the QTL is located at the end of a linkage group, it is possible that the causative mutation(s) is still far from our nearest marker. If this is the case, then we will have underestimated the phenotypic effect of this QTL, and it alone could explain the presence/absence of escape tunnels. An alternative explanation is that escape-tunnel construction is a threshold trait, but only a high-powered experiment would detect additional loci with either additive or epistatic effects on this behaviour. In either case, any additional QTLs are likely to have small phenotypic effects.

Our results show that QTLs are specific to either entrance-tunnel length or escape-tunnel presence, highlighting the modular nature of the complex *P. polionotus* burrowing behaviour. Moreover, all alleles derived from the *P. polionotus* parent cause an increase in trait value (that is, longer entrance tunnels or the addition of an escape tunnel; Fig. 3b), consistent with the role of natural selection in driving the evolution of the complex burrow architecture. Together, these results show that two aspects of burrow architecture evolved independently, as they are genetically discrete, suggesting that entrance-tunnel length and the presence of an escape tunnel are both ecologically important components of the complex *P. polionotus* burrows.

Although progress has been made towards understanding the genetic underpinnings of innate behaviours, most studies have been restricted to a few laboratory-based model systems\(^\text{22}\) or candidate gene approaches in more diverse taxa\(^\text{23}\). By taking advantage of variation in an extended phenotype, in this case burrow architecture in *Peromyscus*, we have gained insight into how a seemingly complex behaviour evolves in the wild. First, we show that the complex, derived burrowing behaviour of *P. polionotus* largely results from evolution at a surprisingly small number of loci. Second, we demonstrate that burrowing behaviour is composed of distinct behavioural modules, which are controlled by independent genetic loci. This is consistent with a century-old observation that elaborate animal behaviour, such as the courtship rituals of grebes, may evolve by integrating simple behaviours\(^\text{24}\). Although genetic modularity has been shown to be important for the evolution of complex morphological traits\(^\text{25}\) and laboratory-based behaviours\(^\text{26}\), our results extend this pattern to wild

Table 1 | Models for individual and combined QTL effects

<table>
<thead>
<tr>
<th>Trait</th>
<th>Chromosome</th>
<th>Position</th>
<th>Genotype completeness (%)</th>
<th>lod score</th>
<th>PVE</th>
<th>Additive effect</th>
</tr>
</thead>
<tbody>
<tr>
<td>Log10 (average entrance-tunnel length)</td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>75.9</td>
<td>77.6</td>
<td>2.48</td>
<td>3.6</td>
<td>0.063</td>
<td></td>
</tr>
<tr>
<td>2</td>
<td>25.1</td>
<td>73.9</td>
<td>3.34</td>
<td>4.9</td>
<td>0.072</td>
<td></td>
</tr>
<tr>
<td>20</td>
<td>59.6</td>
<td>58.8</td>
<td>2.47</td>
<td>3.6</td>
<td>0.064</td>
<td></td>
</tr>
<tr>
<td>Full entrance-tunnel-length model</td>
<td></td>
<td></td>
<td>9.55</td>
<td>14.8</td>
<td>-</td>
<td></td>
</tr>
<tr>
<td>Escape-tunnel presence (binary)</td>
<td>5</td>
<td>60.3</td>
<td>3.38</td>
<td>6.2</td>
<td>0.943</td>
<td></td>
</tr>
</tbody>
</table>

Full model includes all three QTLs. lod, log odds ratio; PVE, percentage of phenotypic variance explained.
behaviours. Thus, we suggest that the behavioural diversity observed in nature may often evolve by accumulating and combining alleles, each with modular effects.

**METHODS SUMMARY**

**Experimental design.** We originally obtained outbred stocks of *Peromyscus* from the Peromyscus Genetic Stock Center. To start the genetic cross, we mated a single female *P. maniculatus bairdii* with a male *P. polionotus subgriseus* to generate 13 F₁ hybrids, which were then backcrossed to *P. maniculatus* to produce 272 BC₁ generation hybrids. We assayed burrowing behaviour following our previously described methods³⁴. In brief, we placed a single, virgin mouse (60–90 days of age) in an 1.2 × 1.5 × 1.1 m enclosure filled with food, water, nesting material and 700 kg of hard-packed sand under constant temperature and light cycle. Each trial lasted approximately 46 h, and at the end of each trial, we made a polyurethane cast of the resulting burrow¹⁶,²⁵. From each cast, we measured the length of the entrance tunnel (from the surface to the nest chamber; Fig. 1b) and recorded the presence/absence of an escape tunnel. All mice were assayed in three, consecutive 2-day trials, each time in a new, randomly assigned enclosure.

**Genotyping.** We extracted DNA using a high-throughput automated phenol-chloroform method (Autogen). We genotyped all individuals using a ddRAD-sequencing approach⁻²⁰, which identified 526 SNPs with fixed nucleotide differences between the parental species. We used the genotypes of hybrids to estimate genetic linkage among markers. This produced a map containing 24 linkage groups, corresponding to the 24 chromosomes in *P. maniculatus* and *P. polionotus*, with a total map length of 1825.5 cM (Supplementary Fig. 2).

**Data analysis.** Burrow lengths were log-normalized before correlation and QTL analyses. We performed all statistical tests in the R statistical package²⁸. QTLs associated with burrow phenotypes were identified using r/QTL²⁹. Specifically, we sequentially performed Haley–Knot regressions, interval mapping, and interval mapping with imputation of missing genotypes, and report QTLs that are consistent across all three methods. We used permutation tests to determine false discovery rates. Linkage groups were assigned to chromosomes using comparative mapping information derived from the *P. polionotus* mitochondrial genome and the *P. maniculatus* nuclear genome. To test whether sex differences were present, we performed sex-specific analyses.

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**Supplementary Information** is available in the online version of the paper.

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**Author Contributions** J.N.W. and H.E.H. conceived and designed the experiments. B.K.P. and J.N.W. generated the ddRAD genotypes. J.N.W. performed the behaviour experiments and analysed the genetic and behavioural data. J.N.W. and H.E.H. wrote the paper.

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METHODS

Field observations and soil analysis. We cast and excavated *P. polionotus* burrows at eight sites, distributed broadly across the species range. First, we removed mice from active burrows by inserting plastic tubing into a burrow entrance tunnel while slowly excavating the tunnel (see Supplementary Video for the method conducted in a laboratory environment). The mice generally exited burrows through their escape tunnels when our excavation neared the nest chamber. Once the mice were removed, we filled the remaining burrow cavity with expansive polyurethane filling foam (either from Hilti, or Great Stuff, Dow Chemicals)27. After the foam hardened, we excavated the cast and measured the entrance tunnel, the total burrow length, as well as the depth of the lowest point in the nest from the surface (Supplementary Table 1).

In addition to burrow measurements, we collected and analysed samples of undisturbed soil that immediately surrounded the nest chambers of excavated burrows. Before analysing the soil, we sifted each sample through a standardized sieve (2 mm diameter) and then oven dried the remaining soil. We then performed a gravimetric particle analysis on the dried soils (using a hydrometer method28) to quantify clay, sand and silt content (Supplementary Table 1).

Statistical analyses of soil and burrow variation. Initially, we log_{10} transformed all continuous variables to make them normally distributed. For the soil and natural burrow measurements, we used conservative, non-parametric tests to analyse these data because we had small sample sizes. Specifically, we used Spearman tests for analyses of correlation and Kruskal–Wallis tests for differences in variation among groups. For the data collected in the laboratory, we calculated Pearson correlation coefficients among all behavioural variables, and also tested for correlations between burrow measures and soil moisture, animal mass and animal age. We found no significant correlations between total burrow length and moisture, mass or age (*P* > 0.05). Similarly, an analysis of variance (ANOVA) showed no differences in the total length of burrows constructed in different enclosures (*P* > 0.05). We performed all statistical tests using R

Genetic cross. We performed experiments using outbred mice originally obtained from the Peromyscus Genetic Stock Center (PGSC). We initially crossed a single female *P. maniculatus bairdii* with a male *P. polionotus subgriseus* to generate 13 first-generation (F1) hybrids, which we then backcrossed to *P. maniculatus* to produce 272 BC-generation hybrids. We followed standard husbandry protocols to maintain constant soil and moisture levels (5–9% and 17–22% water content on the upper and lower surfaces, respectively). We initially identified 1,700 SNPs, each fixed within but different among species of *P. maniculatus* and *P. polionotus*, using a modified RAD-tag approach30.

Briefly, for each individual, we digested ~1 µg of genomic DNA with two restriction endonucleases (100 units of EcoR1-HF and 20 units of Msp1). We ligated the resulting fragments to sequencing adapters containing unique barcodes for each sample. We then pooled ligation products among individuals and isolated fragments in the size range of 280–320 bp using a Pippin Prep electrophoresis platform (Sage BioSciences). Finally, we amplified the remaining fragments using a Phusion High Fidelity PCR Kiti (ThermoFisher Scientific) and sequenced the resulting libraries on a Genome Analyzer II (Illumina). For a more complete description of our computational methods used to analyse short-read sequences and to determine genotypes, see ref. 20.

Linkage map construction. We constructed a linkage map in R/qtl31 using genotypes from 1,700 markers scored in BC-generation individuals. Initially, we calculated the fraction of recombination events and lod score between all marker pairs. We identified 97 markers with genotype information identical to another marker and removed them from the map. We then grouped markers by varying the maximum fraction of recombination events and minimum lod score allowed between markers on the same linkage group. Because the karyotypes of both species are known (*n* = 24 chromosomes), we varied recombination parameters until we recovered a map with 24 linkage groups, each comprising at least 30 markers. This map also contained linkage groups with fewer than 10 markers; we removed the markers in small linkage groups. We ordered the remaining markers by individually scanning linkage groups in overlapping windows of 4–8 markers and minimizing the frequency of recombination events between markers in each window. Next, we removed all markers genotyped in fewer than 100 individuals (Supplementary Fig. 2) as well as all markers with high error rates30. Finally, we evaluated our genotyping error rate by comparing the likelihood of our marker data, given our estimated linkage map, under different error regimes.

QTL mapping. We performed Haley–Knott regressions and interval mapping analyses sequentially in R/qtl31 to identify QTLs contributing to burrowing differences. Using permutation tests, we determined the genome-wide significance level for association between markers and phenotypes (*z* = 0.05, *n* = 2,000 permutations) as lod = 3.05 and 3.08 for average entrance-tunnel length and binary escape-tunnel presence, respectively32. For mapping of escape-tunnel presence, we excluded 29 animals that had unclear escape-tunnel phenotypes (criteria: built an escape tunnel in only one out of three trials and the escape tunnel was ≤4 cm long). Lod scores across all linkage groups for both entrance-tunnel length and escape-tunnel presence are shown in Supplementary Fig. 4. With 272 animals in a backcross design, we had 80% power to detect QTLs with effect sizes of >0.3 s.d., assuming the trait is ~70% heritable32. We also scanned for pairwise interactions between loci across all linkage groups (using R/qtl31), as well as for QTLs that segregated differentially among our BC families due to the outbred nature of our parental stocks (using QTLrel33,34). We found no significant evidence for epistasis or for family differences in QTLs. To determine the additive effect of having different genotypes at markers under QTL peaks, we used the fitql() and refitqtl() functions in R/qtl35 to construct models that estimated the percentage of phenotypic variance explained by each QTL (Table 1).