The evolution of nesting behaviour in *Peromyscus* mice

Caitlin L. Lewarch a, b, c, Hopi E. Hoekstra a, b, c, *

a Department of Organismic and Evolutionary Biology, Harvard University, Cambridge, MA, U.S.A.
b Department of Molecular and Cellular Biology, Harvard University, Cambridge, MA, U.S.A.
c Museum of Comparative Zoology, Howard Hughes Medical Institute, Harvard University, Cambridge, MA, U.S.A.

Structures built by animals, such as nests, often can be considered extended phenotypes that facilitate the study of animal behaviour. For rodents, nest building is both an important form of behavioural thermoregulation and a critical component of parental care. Changes in nest structure or the prioritization of nesting behaviour are therefore likely to have consequences for survival and reproduction, and both biotic and abiotic environmental factors are likely to influence the adaptive value of such differences. Here we first develop a novel assay to investigate interspecific variation in the nesting behaviour of deer mice (genus *Peromyscus*). Using this assay, we find that, while there is some variation in the complexity of the nests built by *Peromyscus* mice, differences in the latency to begin nest construction are more striking. Four of the seven taxa examined here build nests within an hour of being given nesting material, but this latency to nest is not related to ultimate differences in nest structure, suggesting that the ability to nest is relatively conserved within the genus, but species differ in their prioritization of nesting behaviour. We also find that latency to nest is not correlated with body size, climate or the construction of burrows that create microclimates. However, the four taxa with short nesting latencies all have monogamous mating systems, suggesting that differences in nesting latency may be related to social environment. This detailed characterization of nesting behaviour within the genus provides an important foundation for future studies of the genetic and neurobiological mechanisms that contribute to the evolution of behaviour.

Animal architectures — from the webs spun by spiders to the dams built by beavers — can both facilitate the study of behaviour and provide insight into the selective forces that act on behavioural variation (Hansell, 1984, 2005). Such structures can be considered ‘extended phenotypes’, or traits influenced by genetics but extended outside the body of the individual organism (Dawkins, 1982). Building behaviours are often innate and species specific; for example, the resulting structures have been used for classification purposes in insects and some birds (Hansell, 1984; Knerr & Atwood, 2012; Schmidt, 1964; Winkler & Sheldon, 1983). These structures reflect stereotyped patterns of behaviour and the neural circuits that generate these motor patterns, allowing us to study behaviour and the nervous system by proxy. Moreover, the structures themselves serve important functions and can confer readily quantifiable fitness benefits on the animals that construct them (Hayward, 1965; Mainwaring, Hartley, Lambrechts, & Deeming, 2014; Sealander, 1952).

A widespread and important type of building behaviour is the collection and processing of environmental materials to produce a nest. Nests serve a wide variety of purposes for the animals that construct them. For small-bodied animals, such as rodents, nests provide insulation and reduce the energy expended on the maintenance of body temperature (Pearson, 1960; Sealander, 1952; Vogt & Lynch, 1982). In animals with altricial young, like many birds and rodents, nests are especially critical to protect offspring from heat loss and predation (Bult & Lynch, 1997; Collias, 1964; Lynch & Possidente, 1978; Southwick, 1955). The nest may even serve as a catalyst for social behaviour — nest and bower construction can be integral to courtship in birds (Mainwaring et al., 2014), and investment in elaborate nests likely has been instrumental in the evolution of eusociality in insects (Hansell, 2005). Depending on the species in question and the environment in which they live, nests may be built in trees, in pre-existing cavities or in burrow systems that are also constructed by the animal (Collias, 1964; Dooley & Dueser, 1990; Weber & Hoekstra, 2009). While the...
excavation of burrows is itself a type of animal architecture, nests are often separate structures, made by collecting and processing vegetation and other material from an animal’s environment.

Both the structure of a completed nest and the timing of nest building may be relevant traits for natural selection, and each has distinct implications for the proximate and ultimate factors that contribute to behavioural differences among taxa. Variation in nest structure, as is observed in birds, suggests that animals may differ either in their ability to construct nests or in the desired properties of their nests (Mainwaring et al., 2014). At the level of proximate mechanism, variation could result from morphological differences in the animals, fundamental changes in their stereotyped motor patterns or changes in a more abstract encoding of the animal’s target structure. Moreover, variation in nest structures suggests that the characteristics of the nest itself have fitness consequences. Prime examples of such relationships include the pendulous entrances of some weaverbird nests, which are protective against snake predation (Collias, 1964; Crook, 1963), or the increased size and weight of robin, warbler and finch nests built at colder northern versus southern latitudes (Crossman, Rohwer, & Martin, 2011). Variation in the timing of nesting behaviour, on the other hand, implies that animals differ in their motivation to engage in otherwise conserved behavioural patterns, and suggests that the prioritization of nesting relative to other elements of the animal’s behavioural repertoire is relevant for selection. Prioritization can occur at different scales, from time invested over the course of a single night to relative time spent on the behaviour during different seasons. As the collection of nesting material can be energetically costly and expose the animal to predation (Collias, 1964; Mainwaring et al., 2014), it may be beneficial for an animal to prioritize other behaviours in environmental conditions where heat loss, for example, is not a pressing concern. While population differences in nest size have been studied within and between species of rodents (King, Maas, & Weisman, 1964; Lynch, 1992), we do not know how the prioritization of nesting behaviour has evolved.

To determine how and why these features of nesting behaviour evolve, we focused on deer mice (genus Peromyscus), which have adapted to a wide range of habitats and microhabitats across North America (Bedford & Hoekstra, 2015; Blair, 1950; Dewey & Dawson, 2001). Specifically, deer mice live in climates with pronounced differences in winter temperatures (King et al., 1964), vary in body size, a trait associated with adaptation to cold in other rodents (Lynch, 1992), and have distinct social behaviour and parental care (Jasarević et al., 2013; Turner et al., 2010), all of which may affect nest-building behaviour. Importantly, while these species have evolved in different environments, laboratory colonies allow us to perform behavioural experiments under carefully controlled conditions using animals that share a common environment (Bedford & Hoekstra, 2015). This is therefore an opportunity to explore the evolutionary consequences of different environmental parameters on heritable variation in nest-building behaviour.

Here we develop a novel behavioural assay to evaluate natural variation in both ability and motivation to nest in seven species and subspecies of Peromyscus mice. This detailed characterization of thermoregulatory nesting behaviour then provides a foundation to understand the evolution of this behaviour in natural populations.

**METHODS**

**Ethical Note**

All experimental procedures were approved by the Harvard University Institutional Animal Care and Use Committee (IACUC protocol no. protocol 27-15). The animal housing facility in which these tests were performed maintains full AAALAC accreditation.

**Experimental Cohort**

We selected adult, reproductively inexperienced animals of both sexes from seven laboratory colonies of Peromyscus, representing five species, with well-characterized ecology and social systems (Table 1). While these colonies were isolated from natural populations (brought in from the wild between 2 and 71 years ago, depending on strain; Table 1), all animals in this study were born in captivity.

**Animal Husbandry**

All animals were bred and maintained under the same controlled conditions. We kept the animal housing rooms on a 16:8 h light:dark cycle at 22 °C. We housed animals in ventilated polysulfone mouse cages (Allentown, NJ, U.S.A.) of standard size (19.7 × 30.5 cm and 18.5 cm high), with the exception of the Peromyscus californicus animals, which were housed in rat cages (28.6 × 39.4 cm and 19.3 cm high) due to their large body size (Allentown, NJ). For ordinary housing, we provided all cages with 2.5 g of compressed cotton ‘Nestlet’ (Ancare, Bellmore, NY, U.S.A.), 8–10 g folded paper ‘Enviro-Dri’ nesting material (Shepherd Specialty Papers, Watertown, TN, U.S.A.), a 0.6 cm layer of Anderson’s Bed-o-cob (The Andersons, Inc., Maumee, OH, U.S.A.) and enrichment consisting of a red polycarbonate (9.5 × 4.8 cm and 7.6 cm high) mouse hut (BioServ, Flemington, NJ) or a 15.2 × 7.6 cm inside diameter rat tunnel for the large P. californicus animals (BioServ, Flemington, NJ). All animals had ad libitum access to water and irradiated LabDiet Prolab Isopro RMH 3000 5P75 (LabDiet, St Louis, MO, U.S.A.). We socially housed animals in groups of two to five, by species and sex, after weaning (23 days for most species, 30 days for P. californicus), then tested them as adult virgins, averaging 2–6 months old (Table 1).

**Behavioural Paradigm**

**Standard behavioural assay**

Nesting behaviour in rodents is often assessed by measuring the weight of nesting material an animal uses over 24 h (Hartung & Dewsbury, 1979; King et al., 1964; Layne, 1969; Lynch & Hegmann, 1973), which is readily quantifiable but can obscure variation in the timing of the behaviour or the structure of the nests the animals construct. To measure these aspects of nesting behaviour, we designed a novel assay that consists of an overnight habituation period followed by three consecutive days of testing. On the day before a trial began, we weighed and singly housed adult virgin animals in new mouse cages (including P. californicus) with 5 g of compressed cotton nesting material (or two ‘nestlets’, see above), 0.6 cm layer of Anderson’s Bed-o-cob and a red polycarbonate mouse hut. On the morning following habituation to the novel cage, we took photos of the nest from up to three angles (top and two side views), then removed the mouse hut and replaced all cotton nesting material with 5 g of fresh compressed cotton nestlet. The replacement of nesting material during these trials always occurred between 4.5 and 6.5 h after the lights came on. At 1 h after the replacement of nesting material, we again took photographs of the nest from multiple angles and added the mouse hut back to the cage. We repeated this process on the following two mornings for a total of three sets of photographs (day 1, day 2, day 3) at each of the two time points (1 h and overnight). Research assistants blinded
to the species and sex of the animal later scored these nest photographs according to a standardized scale (Fig. 1, Appendix, Table A1). Scores ranged from 0 (no visible shredding) to 4 (a full ‘dome’ nest with overhead coverage) with only full and half scores given.

**Increased nesting material**

To examine whether the amount of nesting material influenced nest scores in the largest species (*P. californicus*, body mass approximately 42 g, on average), we modified the nesting experiment in two ways. First, we singly housed an independent cohort of 21 adult *P. californicus* animals as above, but provided them with an increasing amount of cotton nesting material on four consecutive days: 5 g on day 1, 10 g on day 2, 15 g on day 3 and 20 g on day 4. We photographed nests and exchanged cotton nesting material once every 24 h, and a research assistant blind to experimental conditions scored these photographs as above to establish whether this increase was sufficient to alter overnight nest scores. Based on the results of these experiments, we then assayed an independent group of 23 adult *P. californicus* animals to evaluate their overnight nesting behaviour using 20 g of cotton nesting material in an otherwise standard nesting assay (see Standard Behavioural Assay above).

**Climate Data**

We drew average winter (December/January/February) temperature data from National Oceanic and Atmospheric Administration (NOAA) 30-year climate normals (Arguez et al., 2010) and averaged these data by state or county of origin for each colony (Table 2).

**Data Analysis**

We performed statistical analyses in R using nonparametric methods for the ordinal nest scores. We summarized an animal’s

---

**Table 1**

<table>
<thead>
<tr>
<th>Species (common name)</th>
<th>County isolated</th>
<th>Year in captivity*</th>
<th>Sample size: total (males, females)</th>
<th>Avg. weight, males (g±SD)</th>
<th>Avg. weight, females (g±SD)</th>
<th>Avg. age (days±SD)</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>P. m. nubiterrae</em></td>
<td>Westmoreland County, PA 2010</td>
<td>47 (31, 16)</td>
<td>18.7±2.3</td>
<td>164±184</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. m. bairdii</em></td>
<td>Washtenaw County, MI 1946–1948</td>
<td>95 (62, 33)</td>
<td>20.3±3.5</td>
<td>106±50</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. p. subgriseus</em></td>
<td>Marion County, FL 1952</td>
<td>130 (80, 50)</td>
<td>14.3±1.9</td>
<td>107±57</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. p. leucocephalus</em></td>
<td>Okaloosa County, FL 2015</td>
<td>37 (23, 14)</td>
<td>14.2±1.1</td>
<td>71±17</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. leucopus</em></td>
<td>Avery County, NC 1982–1985</td>
<td>35 (22, 13)</td>
<td>21.7±4.1</td>
<td>66±7</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. gossypinus</em></td>
<td>Jackson County, FL 2009</td>
<td>27 (19, 8)</td>
<td>25.2±7.0</td>
<td>72±9</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. californicus</em></td>
<td>Ventura County, CA 1979–1987</td>
<td>48 (25, 23)</td>
<td>42.1±3.2</td>
<td>126±30</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

* Some species were brought into captivity multiple times over several years (see Bedford & Hoekstra, 2015). Female *P. m. bairdii* give birth to their first litter when they are approximately 3 months old (Bedford & Hoekstra, 2015), and generation times for other species are similar in the laboratory.
behaviour across the three trial days by its median score (to reflect central tendency) or its maximum score (to represent best effort) at each time point after the replacement of nesting material. To test whether short nesting latency is associated with any environmental variables, we assayed in-situ factors such as the state of either mating system (monogamous versus promiscuous), the presence or absence of burrow complexity, and the temperature of the location. To test whether nesting material affects the tendency to build nests, we utilized the ‘fitPagel’ package in R (Pagel, 1994; Revell, 2012). For this test, we utilized the ‘fitMK’ method, allowed all rates of change to be estimated by the model (model = ‘ARD’), and set nesting latency (short versus intermediate/long) to be dependent on the state of the mating system (monogamous versus promiscuous) or burrow complexity (complex versus simple/absent). As there are no laboratory data on burrowing behaviour in Peromyscus gossypinus, this species was excluded from the latter analysis.

### RESULTS

#### Interspecific Variation in Nesting Latency

To measure an animal’s motivation to nest, we assayed individuals from seven Peromyscus taxa with known evolutionary relationships (Fig. 2a). First, we analysed the median of the three scores an animal received 1 h after the replacement of nesting material, which reflects the tendency of the animal to begin nesting shortly after their nest is disturbed. Scores at 1 h were significantly correlated across the 3 days in the full data set (Spearman rank correlations: day 1 versus day 2: \( r_S = 0.75; \) day 1 versus day 3: \( r_S = 0.68; \) day 2 versus day 3: \( r_S = 0.78; N = 419, P < 2.2 \times 10^{-16} \) for each), and species comparisons were largely the same whether 3-day medians or maxima were used (see below).

In the standard model, the taxa with known evolutionary relationships were used to determine the presence of a significant difference in nesting latency between taxa. We performed phylogenetic generalized least squares (PGLS) analysis using the ‘ape’ package in R (Paradis, Claude, Strimmer, 2004; Pinheiro, Bates, DebRoy, Sarkar, & Core Team, 2017; Symonds & Blomberg, 2014). Covariance due to relatedness was modelled by Brownian motion using the corBrownian function in ape. The covariance was then included as a correlation parameter in the generalized least squares analyses in nlme. The effect of each environmental variable on 1 h nest scores was tested independently. To test whether nesting material affects the tendency to build nests, we utilized the ‘fitPagel’ package in R (Pagel, 1994; Revell, 2012). For this test, we utilized the ‘fitMK’ method, allowed all rates of change to be estimated by the model (model = ‘ARD’), and set nesting latency (short versus intermediate/long) to be dependent on the state of the mating system (monogamous versus promiscuous) or burrow complexity (complex versus simple/absent). As there are no laboratory data on burrowing behaviour in Peromyscus gossypinus, this species was excluded from the latter analysis.
in the first hour and suggest that they are relatively slow to initiate nest construction. Ranking of each taxon’s performance largely followed the same pattern whether maximum or median nest scores were used (Appendix, Fig. A1). The only exception was one species difference: while the median nest scores of *P. gossypinus* and *P. m. bairdii* animals at 1 h were indistinguishable, *P. gossypinus* were slightly more likely to shred the nesting material on at least one of the trial days and therefore had slightly,
but significantly, higher maximum scores (Wilcoxon two-sample test: $W = 839$, $N_1 = 27$, $N_2 = 95$, Bonferroni-corrected $P = 0.049$). We note that taxa differ in the variance of their nest scores: this likely results from within-species variation in nest-building efficiency, time spent nesting and/or the precise initiation time during the first hour. In summary, based on our analysis of 1 h median scores, we identified three main groups of nest builders in our assay: those with short, intermediate or long latencies to nest.

**Interspecific Differences in Nesting Ability**

We next asked whether these taxa differed in their overall ability to construct a three-dimensional nest. To establish the highest-scoring nest that an animal was capable of producing, we used the maximum score achieved over the individual's three overnight time points, which represents the animal's best effort during the longest interval of the trial. Maximum overnight scores varied significantly among taxa (Fig. 2c; Kruskal–Wallis test: $H_N = 127.21, P < 2.2 \times 10^{-16}$), although most animals built full or partial domes. The highest scoring nests were consistently constructed by *P. m. nubiterrae*, *P. p. leucopus* and *P. gossypinus*, which tended to build statistically indistinguishable full domes (Appendix, Table A2; Wilcoxon two-sample test, Bonferroni-corrected $P > 0.05$ for each pairwise comparison). Three taxa — *P. m. bairdii*, *P. p. subgriseus* and *P. leucopus* — had equivalently high maximum scores (Appendix, Table A2; Wilcoxon two-sample test, Bonferroni-corrected $P > 0.05$ for each pairwise comparison), and *P. m. bairdii* and *P. p. subgriseus* which tended to build domes with only partial cover, were significantly different from all but *P leucopus* animals (Appendix, Table A2; Wilcoxon two-sample test, Bonferroni-corrected $P < 0.05$ for each pairwise comparison). Finally, *P. californicus* tended to build nests with walls but without overhead cover, and had significantly lower maximum nest scores than all other species tested (Appendix, Table A2; Wilcoxon two-sample test, Bonferroni-corrected $P < 0.05$ for each pairwise comparison). Notably, we found that all species had at least one individual who constructed a domed nest with full cover (maximum nest score, ‘4’) during the assay, suggesting that all species are capable of building a ‘complete’ nest if given enough time. However, some species showed a large variance in nest scores, and *P. californicus* tended to have lower maximum scores than the other species.

**Nest-building Behaviour in the Large *P. californicus* Mice**

*Peromyscus californicus* animals are much larger than the other taxa included in this study (Table 1), and therefore might require more material to construct a dome nest with overhead cover. To test the possibility that these animals built lower-scoring nests because 5 g of nestlet was an insufficient amount of nesting material, we conducted two additional experiments. First, we gave a group of *P. californicus* increasing amounts of nesting material on four consecutive days and evaluated the nests they produced in each 24 h interval. We found that increasing nesting material from 5 g to 20 g could increase overnight nesting scores (Appendix, Fig. A2; Friedman test: $\chi^2_3 = 13.468, P = 0.004$). However, when we provided an independent group of *P. californicus* with 20 g of nesting material during a 3-day trial (Fig. 2d), there was no difference in overnight maximum scores between those *P. californicus* given 5 g of nestlet and those given 20 g (Wilcoxon two-sample test: $W = 609$, $N_1 = 23$, $N_2 = 48$, $P = 0.47$). Moreover, the maximum overnight nest scores for *P. californicus* given 20 g of nestlet remained significantly lower than the maximum nest scores for all other species (Appendix, Table A2; Wilcoxon two-sample test, Bonferroni-corrected $P < 0.05$ for each pairwise comparison). Thus, the poor nest construction of *P. californicus* in this assay cannot be attributed simply to insufficient nesting material relative to its large body size.

**Sex Differences in Nesting**

We next investigated whether there were any sex differences in the nest scores produced by each species and subspecies. Only two taxa showed evidence of sexual dimorphism in nesting (Fig. 3). Both male *P. m. nubiterrae* and male *P. p. subgriseus* built higher-scoring nests than their female counterparts 1 h after the start of the assay (Appendix, Table A3; Wilcoxon two-sample test, Bonferroni-corrected $P = 0.03$ and 0.002, respectively), and *P. p. subgriseus* males also built higher-scoring nests at the overnight time point (Appendix, Table A3, Fig. A3; Wilcoxon two-sample test, Bonferroni-corrected $P = 0.008$). No other species showed evidence of sex differences in nest scores at either time point (Appendix, Table A3, Wilcoxon two-sample test, Bonferroni-corrected $P > 0.05$ for each pairwise comparison). Therefore, while there was no sexual dimorphism in nesting behaviour for most taxa, in both instances when sex differences were observed, males constructed higher-scoring nests than the females.

**Association between Body Size and Nest Building**

To determine whether body size had an effect on nest-building behaviour, we tested for correlations between body weight and performance in the nesting assay. We found that body weight significantly varied by species, sex and species-by-sex interactions in our experimental cohort (two-way ANOVA, main effect of species: $F_{6,352} = 365.9, P < 2 \times 10^{-16}$; main effect of sex: $F_{1,352} = 5.3, P = 0.02$; interaction: $F_{6,352} = 4.1, P = 0.0005$). However, there was no evidence that species-level average weights altered median 1 h nest scores (Fig. 4a; phylogenetic generalized least squares, median 1 h nesting score by average weight: coefficient $= -0.01$, $SE = 0.06$, $t = -0.15$, $P = 0.89$). Likewise, when we divided the animals by species and sex, we found no correlation between body weight and median nest score at 1 h (Appendix, Table A4; Spearman’s rank correlations, Bonferroni-corrected $P > 0.05$) or maximum overnight nest score (Appendix, Table A4, Spearman’s rank correlations, Bonferroni-corrected $P > 0.05$) within any of the species—sex groups. Thus, while average weight varied almost three-fold among species, body weight was not associated with nesting behaviour in our assay.

**Association between Environment and Nest Construction**

We next asked whether there was an association between performance in the nesting assay and several additional environmental covariates, including latitude and average winter temperature of origin, burrow construction and mating system. Neither latitude nor average winter temperatures were significantly associated with median 1 h scores in these species (Table 2, Fig. 4b; phylogenetic generalized least squares, median 1 h nest score by latitude: coefficient $= -0.05$, $SE = 0.08$, $t = -0.71$, $P = 0.51$; median 1 h nest score by average winter temperature: coefficient $= 0.03$, $SE = 0.05$, $t = 0.69$, $P = 0.52$). Moreover, nesting latency did not appear to be influenced by burrowing behaviour: a model in which short nesting latency was dependent on building complex burrows did not fit the data significantly better than a model where the two traits were
independent (Table 2, Fig. 4b; Pagel's binary character correlation test: AIC (independent model) = 21.97, AIC (dependent model) = 23.88, likelihood ratio = 2.09, \( P = 0.35 \)). However, a model in which short nesting latency depended on mating system fit the observed data significantly better than a model assuming the two traits were independent (Table 2, Fig. 4c; Pagel's binary character correlation test: AIC (independent model) = 26.42, AIC (dependent model) = 21.21, likelihood ratio = 9.21, \( P = 0.01 \)). With the caveat that the sample size for comparisons among taxa is small, these data suggest that mating system is correlated with nesting latency but the other abiotic environmental factors we examined are not.

**DISCUSSION**

Nesting is important for survival in rodents, but it is not clear how this behaviour varies among species or which evolutionary pressures drive these changes. Here we designed a novel high-throughput phenotyping paradigm to evaluate variation in both nest structure and the timing of nesting behaviour in closely related species of deer mice. We found that *Peromyscus* mice are generally able to construct dome-shaped nests, but vary strikingly in their latency to do so. Because nesting latency is not simply correlated with phylogeny, this raises the possibility that natural selection may contribute to intertaxon variation. When we tested for correlations between latency to nest and several abiotic and biotic variables, we found that mating system, but surprisingly not climate or body size, was correlated with nesting behaviour.

Nesting has been well studied in laboratory models (e.g. Lisk, Pretlow, & Friedman, 1969; Lynch, 1980). However, the majority of these nesting experiments, including some studies in *Peromyscus*, measure the amount of nesting material that an animal pulls into its cage or the final nest structure achieved over a 24 h period (Hartung & Dewsbury, 1979; King et al., 1964; Layne, 1969; Lynch & Hegmann, 1973). By contrast, we focused on both the timing of the behaviour and the final nest structure. By evaluating nests just 1 h after the replacement of nesting material, we were able to assess whether the animals differed in their latency to begin nest construction—what might be considered a baseline motivation to nest. This was complemented by a second measurement at the more permissive overnight time point, which allowed us to evaluate whether...
animals varied in their overall ability to build three-dimensional structures. This novel phenotyping paradigm therefore allowed us to distinguish between animals that differed in their motivation to construct nests of similar shape from those that differed in their ability to construct nests.

Using this approach, we found that even closely related *Peromyscus* species vary dramatically in their latency to begin nesting, while variation in final nest structure is much more modest. This is in contrast to studies of nesting in birds and insects, where the structures of complete, species-typical nests are highly variable (Collias, 1964; Healy, Walsh, & Hansell, 2008; Knerer & Atwood, 2012; Price & Griffith, 2017; Schmidt, 1964), or even burrow construction in *Peromyscus*, where species excavate cavities that significantly differ in size and shape (Hu & Hoekstra, 2017; Weber & Hoekstra, 2009). The relative conservation of nest structure implies that the ability to produce dome-shaped nests is important for most animals in the genus. However, variation in latency to begin nesting suggests that prioritization of the behaviour varies among taxa. These patterns also imply that variation in nesting in these mice is likely due to altered motivation rather than changes in stereotyped motor patterns, morphology or target nest structure.

All animals were acclimated to and tested in a common environment, specifically at 22 °C, which is below the preferred temperatures (Ogilvie & Stinson, 1966) and thermoneutral zones (Glaser & Lustick, 1975; Hayward, 1965; Layne & Dolan, 1975) of many *Peromyscus* species. While these taxa may differ in their behavioural response to this thermal environment due to differences in basal metabolic rate or thermoneutral zone, we found no evidence of a correlation between nesting scores at either time point and body weight, a trait strongly related to both metabolic parameters in *Peromyscus* (Hayward, 1965; Hill, 1983). It is worth noting that the positive relationship between body weight and nest weight observed in previous studies of rodent nesting (King et al., 1964; Lynch, 1992; Wolfe, 1970) might be at least partially explained by larger animals requiring more material to build equivalently shaped structures. By focusing on the structure of the nest rather than the weight of nesting material used to construct it, we minimized this confounding factor.

Body size aside, it is reasonable to hypothesize that climate could alter this thermoregulatory behaviour. Other studies have suggested that climate (King et al., 1964; Lynch, 1992) and microclimate (Wolfe, 1970) alter the amount of nesting material used by rodents in natural populations. However, we found only modest variation in final nest shape and no evidence for a relationship between nesting latency and average winter temperatures or latitude of origin, which is frequently used as a proxy for temperature. Nor did we find evidence for an association between nesting latency and the construction of elaborate burrows, which function as microclimates and buffer the animals from changes in ambient temperature (Hayward, 1965; Sealander, 1952; Weber & Hoekstra, 2009). Although these colonies have experienced reduced selective pressure while bred in laboratory settings, there was still no evidence of an association between climate/microclimate and latency when we considered only the colonies founded within the past 10 years (Table 1; *P. gossypinus, P. p. leucocephalus, P. m. californicus*, *P. p. subgriseus, P. m. nubiterrae, P. m. bairdii*, and *P. n. argenteus*).

---

**Figure 4.** Environmental factors and nesting behaviour. (a) Species median 1 h nest scores and average body weight are plotted with bars indicating interquartile range (nest score) and standard deviation (body weight). (b) Sites of colony origin (on USA map), burrow shape (by symbol) and nesting latency (by colour) are indicated for each taxon (following legend). Map colours represent average winter temperatures by state (Arguez et al., 2010). Nesting latency category (short, intermediate, long) was determined by the significant species groups depicted in Fig. 2b. (c) Association between a taxon’s mating system and nesting latency.
nubiterrae). Given that most of the variation we observed took the form of prioritization differences rather than changes in nest size or shape, it may be especially necessary to consider the broader behavioural repertoire of these taxa, including biotic factors that could contribute to the motivation to nest.

While we did not observe a simple relationship between nesting behaviour and any of the abiotic factors we examined, we did find an intriguing correlation between social environment and nesting latency. With the caveats that our sample size was relatively low and that the classification of species as monogamous or promiscuous relied on incomplete evidence, our results suggest that mating system and nesting latency are not independent, with all putatively monogamous species having short latencies to nest. It is possible that this reflects a tendency to invest in a home territory that is more beneficial for monogamous animals than for promiscuous ones (Gaulin & FitzGerald, 1988), or that selection for increased paternal care, a hallmark of monogamous mating systems (Kleiman, 1977), might result in increased motivation to nest even in virgin animals. This potential relationship between social behaviour and the prioritization of nesting behaviour underscores the importance of considering both biotic and abiotic environment when investigating the causes of behavioural evolution.

Conclusion

Measurement of extended phenotypes such as nests allows us to study how behaviours evolve within and between species. Here we showed that the ability to nest is relatively conserved in the genus Peromyscus, but latency to begin nest construction is highly variable, even between sister species. This suggests that evolution of nesting behaviour in these animals is characterized by differences in the prioritization of an otherwise conserved behavioural pattern. Intriguingly, while abiotic environment cannot explain these species differences in nesting behaviour, we found a correlation between latency to nest and mating system, with monogamous species prioritizing nesting. Finally, as the innate differences in nesting behaviour in Peromyscus appear to be largely changes in the motivation to nest, future studies in this system may elucidate genetic and neurobiological mechanisms that lead to differences in motivation to engage in particular behaviours, a topic with implications far beyond nesting behaviour.

Acknowledgments

We thank Harvard undergraduates A. Bialkowski, M. Charifson, E. D’Agostino, M. Noriega and J. Rhodes for blinded scoring of behavioural assays. N. Bedford and C. Hu assisted with the figures. L. Revell and O. Lapedra provided advice on comparative methods. N. Bedford, E. Hager, D. Haig, C. Hu, B. König, M. Noriega, K. Pritchett-Corning, C. Reed, K. Turner and two anonymous referees provided helpful feedback on the manuscript. We also thank the Harvard OAR staff for their assistance with animal husbandry. C.L.L. was supported by a Morris E. Zuckerman Fellowship, a Smith Family Graduate Science and Engineering Fellowship and the Harvard Molecules, Cells, and Organisms PhD Program. C.L.L. received a Harvard Mind, Brain, and Behaviour Student Award. H.E.H. is an Investigator of the Howard Hughes Medical Institute. This work was supported, in part, by a National Science Foundation Doctoral Dissertation Improvement Grant (DDIG IOS 1701805).

References


Appendix

Figure A1. Maximum nest score achieved 1 h after receiving new nesting material over the 3 trial days. Letters indicate species groups that did not significantly differ from another, while all other pairwise comparisons were significant (Wilcoxon two-sample test: Bonferroni-corrected $P < 0.05$; compare with median scores in Fig. 2b, Table A2). Box plots show the 25% and 75% quartiles (boxes), medians (lines inside boxes), and outermost values within 1.5 times the interquartile range of their respective quartiles (whiskers). Sample sizes are provided for each species.

<table>
<thead>
<tr>
<th>Species</th>
<th>manicolors</th>
<th>bairdii</th>
<th>subterrane</th>
<th>leucopus</th>
<th>gosspinus</th>
<th>californica</th>
</tr>
</thead>
<tbody>
<tr>
<td>N</td>
<td>47</td>
<td>95</td>
<td>130</td>
<td>35</td>
<td>27</td>
<td>48</td>
</tr>
</tbody>
</table>

Figure A2. Effect of increasing nesting material in *P. californicus*. Adult animals (*N* = 21) were given increasing amounts of nesting material (5 g, 10 g, 15 g and 20 g) on 4 sequential days (Friedman repeated measures test: $P = 0.003$). Box plots as in Fig. A1.


Figure A3. Sex differences in maximum overnight nest scores (Wilcoxon two-sample test, Bonferroni-corrected $P$). Sample sizes are provided for females and males of each species. Box plots as in Fig. A1.
Figure A4. Effect of body weight on nesting behaviour by species and sex. (a) Median nest scores at 1 h (Spearman correlation, Bonferroni-corrected $P > 0.05$). (b) Overnight maximum nest scores (Spearman correlation, Bonferroni-corrected $P > 0.05$). Sample sizes are provided by sex and were the same for both time points. Box plots as in Fig. A1.

Table A1
Detailed nest scoring criteria

<table>
<thead>
<tr>
<th>Score</th>
<th>Score description</th>
<th>Shredding</th>
<th>Nest site$^a$</th>
<th>Walls$^b$</th>
<th>Overhead cover</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>No manipulation</td>
<td>None</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>0.5</td>
<td>Minor shredding</td>
<td>Minor: ≤ top of 1 nestlet</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>1</td>
<td>Extensive shredding</td>
<td>Extensive: &gt; top of 1 nestlet</td>
<td>No: neither a nor b is true</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td>1.5</td>
<td>Ambiguous nest site</td>
<td>Extensive</td>
<td>Unclear: either a or b is true</td>
<td>No</td>
<td>–</td>
</tr>
<tr>
<td>2</td>
<td>Platform nest</td>
<td>–</td>
<td>Yes: both a and b are true</td>
<td>No: ≤½ sphere height for ≤½ circumference</td>
<td>–</td>
</tr>
<tr>
<td>2.5</td>
<td>Partial cup nest</td>
<td>–</td>
<td>Yes</td>
<td>Partial: ≤½ sphere height for &gt;½ circumference OR ≥½ sphere height for &lt;½ of circumference</td>
<td>–</td>
</tr>
<tr>
<td>3</td>
<td>Cup nest</td>
<td>–</td>
<td>Yes</td>
<td>Yes: ≥½ sphere height for ≥½ circumference</td>
<td>No overhead cover</td>
</tr>
<tr>
<td>3.5</td>
<td>Partial dome nest</td>
<td>–</td>
<td>Yes</td>
<td>Yes</td>
<td>Partial: ≤50% of the sphere is covered or there are multiple entrance holes and there is at most one entrance hole</td>
</tr>
<tr>
<td>4</td>
<td>Full dome nest</td>
<td>–</td>
<td>Yes</td>
<td>Yes</td>
<td>Yes: ≥50% of the sphere is covered</td>
</tr>
</tbody>
</table>

$^a$ A nest site is defined according to two criteria: (a) there is a ‘contiguous concentration’ of nestlet around a ‘central point’ consisting of ≥90% of any material the animal has shredded; (b) the shape of the putative nest site is defined by the ‘shredded material’ (≥50% of the nestlet at the nest site is shredded).

$^b$ To evaluate walls, imagine that the nest cavity is filled by a sphere (sensu Hess et al., 2008). The walls are compared to the height of the sphere within the cavity, and the proportion of the circumference of the sphere that is surrounded by walls is noted.
Table A2

Pairwise species comparisons of 1 h median and overnight maximum scores for Peromyscus mice

<table>
<thead>
<tr>
<th>Species</th>
<th>P. m. nubiterrae</th>
<th>P. m. bairdii</th>
<th>P. p. subgriseus</th>
<th>P. p. leucocephalus</th>
<th>P. leucopus</th>
<th>P. gossypinus</th>
<th>P. californicus (5 g)</th>
<th>P. californicus (20 g)</th>
</tr>
</thead>
<tbody>
<tr>
<td>N.</td>
<td>47</td>
<td>95</td>
<td>130</td>
<td>37</td>
<td>35</td>
<td>27</td>
<td>48</td>
<td>23</td>
</tr>
<tr>
<td>P. m.</td>
<td>W=3224.5</td>
<td>W=4124.5</td>
<td>W=819.5</td>
<td>W=905</td>
<td>W=540</td>
<td>W=2102</td>
<td>W=974.5</td>
<td>W=76.10^-8</td>
</tr>
<tr>
<td><em>P. nubiterrae</em></td>
<td><em>P=9.3</em>10^-5</td>
<td>*P=0.0026</td>
<td>*P=1</td>
<td>*P=1</td>
<td>*P=1</td>
<td><em>P=2.8</em>10^-5</td>
<td><em>P=3.8</em>10^-6</td>
<td>*P=0.027</td>
</tr>
<tr>
<td>P. m.</td>
<td>W=4046</td>
<td>W=5306.5</td>
<td>W=846</td>
<td>W=1133</td>
<td>W=545</td>
<td>W=3473</td>
<td>W=1553.5</td>
<td>W=2428.5</td>
</tr>
<tr>
<td><em>P. m. Bairdii</em></td>
<td><em>P=2.2</em>10^-16</td>
<td>*P=1</td>
<td><em>P=1.8</em>10^-5</td>
<td>*P=0.073</td>
<td><em>P=2.8</em>10^-5</td>
<td><em>P=3.8</em>10^-6</td>
<td>*P=0.027</td>
<td><em>P=1.3</em>10^-5</td>
</tr>
<tr>
<td>P. p.</td>
<td>W=2565</td>
<td>W=734</td>
<td>W=1535.5</td>
<td>W=1778</td>
<td>W=866</td>
<td>W=5435.5</td>
<td>W=2428.5</td>
<td></td>
</tr>
<tr>
<td><em>P. p. Subgriseus</em></td>
<td>*P=1</td>
<td>*P=0.0025</td>
<td>*P=0.72</td>
<td>*P=0.00017</td>
<td><em>P=0.6</em>10^-4</td>
<td><em>P=1.3</em>10^-5</td>
<td>*P=0.0024</td>
<td><em>P=4.9</em>10^-7</td>
</tr>
<tr>
<td>P. p. leucocephalus</td>
<td>W=924.5</td>
<td>W=256.6</td>
<td>W=3088.5</td>
<td>W=746.5</td>
<td>W=447.5</td>
<td>W=1603</td>
<td>W=791</td>
<td></td>
</tr>
<tr>
<td><em>P. leucocephalus</em></td>
<td>*P=1</td>
<td>*P=0.16</td>
<td>*P=1</td>
<td>*P=1</td>
<td><em>P=2.9</em>10^-12</td>
<td><em>P=3.9</em>10^-8</td>
<td>*P=0.027</td>
<td><em>P=3.9</em>10^-8</td>
</tr>
<tr>
<td>P. gossypinus</td>
<td>W=1159.5</td>
<td>W=375</td>
<td>W=3593.5</td>
<td>W=929.5</td>
<td>W=362</td>
<td>W=1438.5</td>
<td>W=666</td>
<td></td>
</tr>
<tr>
<td><em>P. gossypinus</em></td>
<td>*P=0.027</td>
<td>*P=0.0023</td>
<td>*P=0.59</td>
<td><em>P=2.9</em>10^-7</td>
<td>*P=0.00024</td>
<td><em>P=1.2</em>10^-9</td>
<td><em>P=4.9</em>10^-7</td>
<td></td>
</tr>
<tr>
<td>P. californicus</td>
<td>W=990.5</td>
<td>W=103</td>
<td>W=3492</td>
<td>W=678.5</td>
<td>W=281</td>
<td>W=74.5</td>
<td></td>
<td></td>
</tr>
<tr>
<td><em>P. californicus</em></td>
<td><em>P=2.2</em>10^-16</td>
<td>*P=1</td>
<td>*P=1</td>
<td><em>P=3.1</em>10^-6</td>
<td><em>P=2.4</em>10^-9</td>
<td></td>
<td></td>
<td></td>
</tr>
</tbody>
</table>

The results of pairwise Wilcoxon two-sample tests for species differences in median 1 h scores (below diagonal) or maximum overnight scores (above diagonal). For each comparison, test statistics (W) and Bonferroni-corrected P values are reported; significant results (P < 0.05) are in bold.

Table A3

Sex differences in nest scores of Peromyscus mice

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Males</th>
<th>Females</th>
<th>1 h median score</th>
<th>Maximum overnight score</th>
</tr>
</thead>
<tbody>
<tr>
<td>P. m. nubiterrae</td>
<td>31</td>
<td>16</td>
<td>W=1195.5 P=0.03</td>
<td>W=266 P=1</td>
</tr>
<tr>
<td>P. m. bairdii</td>
<td>62</td>
<td>33</td>
<td>W=858 P=0.11</td>
<td>W=1173 P=1</td>
</tr>
<tr>
<td>P. p. subgriseus</td>
<td>80</td>
<td>50</td>
<td>W=1262.5 P=0.002</td>
<td>W=1361 P=0.008</td>
</tr>
<tr>
<td>P. p. leucocephalus</td>
<td>23</td>
<td>14</td>
<td>W=1635.5 P=1</td>
<td>W=133 P=1</td>
</tr>
<tr>
<td>P. leucopus</td>
<td>22</td>
<td>13</td>
<td>W=125 P=1</td>
<td>W=152 P=1</td>
</tr>
<tr>
<td>P. gossypinus</td>
<td>19</td>
<td>8</td>
<td>W=87 P=1</td>
<td>W=84 P=1</td>
</tr>
<tr>
<td>P. californicus</td>
<td>25</td>
<td>23</td>
<td>W=2195 P=1</td>
<td>W=238 P=1</td>
</tr>
</tbody>
</table>

The results of pairwise Wilcoxon two-sample tests for sex differences in 1 h median nest scores or maximum overnight nest scores. Sample sizes, test statistics (W), and Bonferroni-corrected P values are reported; significant results (P < 0.05) are in bold.

Table A4

Spearman correlations between weight and nest scores within species/sex groups of Peromyscus mice

<table>
<thead>
<tr>
<th>Taxon</th>
<th>Weight vs 1 h median score</th>
<th>Weight vs maximum overnight score</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Males</td>
<td>Females</td>
</tr>
<tr>
<td>P. m. nubiterrae</td>
<td>$r_s=-0.24$, N=7, P=1</td>
<td>$r_s=-0.09$, N=6, P=1</td>
</tr>
<tr>
<td>P. m. bairdii</td>
<td>$r_s=-0.22$, N=58, P=1</td>
<td>N=29</td>
</tr>
<tr>
<td>P. p. subgriseus</td>
<td>$r_s=-0.01$, N=76, P=1</td>
<td>$r_s=-0.04$, N=48, P=1</td>
</tr>
<tr>
<td>P. p. leucocephalus</td>
<td>$r_s=-0.03$, N=23, P=1</td>
<td>$r_s=-0.10$, N=14, P=1</td>
</tr>
<tr>
<td>P. leucopus</td>
<td>$r_s=-0.41$, N=22, P=0.82</td>
<td>$r_s=0.22$, N=13, P=1</td>
</tr>
<tr>
<td>P. gossypinus</td>
<td>$r_s=-0.25$, N=17, P=1</td>
<td>$r_s=-0.24$, N=7, P=1</td>
</tr>
<tr>
<td>P. californicus</td>
<td>$r_s=-0.08$, N=23, P=1</td>
<td>$r_s=-0.14$, N=23, P=1</td>
</tr>
</tbody>
</table>

Sample sizes, Spearman correlation coefficient ($r_s$), and Bonferroni-corrected P values are reported. Sample sizes are smaller than for other tests due to missing weight data.\*

\*We were unable to perform correlations between 1 h scores and weight within P. m. bairdii females because all 29 animals received a median score of 0 at 1 h.

\*Similarly, all female P. gossypinus produced maximum scores of 4 at the overnight time point.