Fitness, risk taking, and spatial behavior covary with boldness in experimental vole populations

Jana A. Eccard1 | Antje Herde1,2 | Andrea C. Schuster1,3 | Thilo Liesenjohann1,4 | Tatjana Knopp1 | Gerald Heckel3 | Melanie Dammhahn1

Abstract
Individuals of a population may vary along a pace-of-life syndrome from highly fecund, short-lived, bold, dispersive “fast” types at one end of the spectrum to less fecund, long-lived, shy, plastic “slow” types at the other end. Risk-taking behavior might mediate the underlying life history trade-off, but empirical evidence supporting this hypothesis is still ambiguous. Using experimentally created populations of common voles (Microtus arvalis)—a species with distinct seasonal life history trajectories—we aimed to test whether individual differences in boldness behavior covary with risk taking, space use, and fitness. We quantified risk taking, space use (via automated tracking), survival, and reproductive success (via genetic parentage analysis) in 8 to 14 experimental, mixed-sex populations of 113 common voles of known boldness type in large grassland enclosures over a significant part of their adult life span and two reproductive events. Populations were assorted to contain extreme boldness types (bold or shy) of both sexes. Bolder individuals took more risks than shyer ones, which did not affect survival. Bolder males but not females produced more offspring than shy conspecifics. Daily home range and core area sizes, based on 95% and 50% Kernel density estimates (20 ± 10 per individual, n = 54 individuals), were highly repeatable over time. Individual space use unfolded differently for sex-boldness type combinations over the course of the experiment. While day ranges decreased for shy females, they increased for bold females and all males. Space use trajectories may, hence, indicate differences in coping styles when confronted with a novel social and physical environment. Thus, interindividual differences in boldness predict risk taking under near-natural conditions and have consequences for fitness in males, which have a higher reproductive potential than females. Given extreme inter- and intra-annual fluctuations in population density in the study species and its short life span, density-dependent fluctuating selection operating differently on the sexes might maintain (co) variation in boldness, risk taking, and pace-of-life.

KEYWORDS
animal personality, automated radio telemetry, behavioral type, fitness, home range, Microtus arvalis, parentage, reproductive success
1 INTRODUCTION

Risk-reward trade-offs may favor the coexistence of different behavioral types in populations (Sih & Del Giudice, 2012). Bolder animals may be rewarded for taking higher risks by producing more offspring, and shyer animals may have an increased life span but lower reproductive output per time unit. In species that are highly depredated, however, the fitness gains must clearly outweigh the survival costs of boldness to maintain bold behavior. Alternatively, different behavioral phenotypes can be favored under different environmental conditions, which may lead to similar fitness between behavioral phenotypes and fluctuations of phenotype frequencies in populations (Bergeron et al., 2013; Dingemanse et al., 2004; Nicolaus et al., 2016; Roth et al., 2021).

Similarly, selection related to fluctuations in population density can maintain variation in life history trajectories (Sæther et al., 2016), linking ecological dynamics to evolutionary processes. Life history trajectories may be related to favorable physiological and behavioral components, forming an extended pace-of-life syndrome (POLS, e.g., Careau et al., 2008; Dammhahn et al., 2018; Réale et al., 2010) with fast POL individuals increasing their fitness by higher risk taking, and slow POL individuals by avoiding risks (Wolf et al., 2007a; Wolf et al., 2007b; Wright et al., 2019). Thus, both variation in life histories and the associated among-individual differences in behavior could potentially be explained through their eco-evolutionary dynamics with fluctuations in population density (Milles et al., 2022; Wright et al., 2019), group size, or composition of personality in groups (Roth et al., 2019).

Small rodents offer a suitable study system to assess whether and how among-individual differences translate into variation in risk taking and space use and have consequences for fitness components. Despite extreme predation pressure (Norrdahl & Korpimäki, 1998), consistent individual differences in risk taking have been observed in several small rodent species (Eccard et al., 2020; Herde & Eccard, 2013; Lantová et al., 2011; Mazza et al., 2018). Small mammals in temperate environments follow very distinct life history trajectories within populations, with some individuals—born early or in the middle of the productive season—reproducing immediately and repeatedly in the season of birth and other individuals—born late in the productive season—having to delay maturity, survive the unproductive season and, wait for the next productive season (Eccard & Herde, 2013). These trajectories are flexible and triggered by density-dependent processes (Prévot-Julliard et al., 1999), allowing the parallel existence of very different life history trajectories, possibly connected to behavioral differentiation into pace-of-life syndromes and maintained by frequency-dependent selection during density fluctuations (Wright et al., 2019).

Consistent among-individual differences in behavior may contribute to variation in individual spatiotemporal distribution and might, thus, influence individuals’ interactions with biotic and abiotic components of their environment (Bolnick et al., 2011; Wolf & Weissing, 2010). Whether feedback between space use and individual differences in behavior exists and how this potential feedback drives and/or maintains intraspecific (co)variation in these traits under heterogeneous environmental conditions is matter of current debate (Spiegel et al., 2017). In order to start illuminating these aspects, we need studies quantifying among-individual differences in behavior and space use independently from each other (e.g., birds; Arvidsson et al., 2017). More ideally, proxies of fitness components, such as reproductive success and survival, would allow assessing the consequences of interindividual differences in behavior. The main aim of this study was to investigate whether between-individual differences in risk taking and activity behavior that are measured in the laboratory are linked to space use in the field. The second aim was to investigate whether the laboratory measurements can be used to predict survival and reproductive success under field conditions.

We focused on common voles (Microtus arvalis), a common microtine rodent, characterized by a high reproductive potential balancing strong predation pressure, a promiscuous mating system (Borkowska & Ratkiewicz, 2010; Fink et al., 2006), larger home ranges of males than females (as other Microtus species: Borowski & Owadowska, 2010; Gliwicz, 1997; Solomon & Jacquot, 2011), which are overlapping (Madison, 1980; Spritzer et al., 2006), and male-biased dispersal (Hahne et al., 2011). Common voles can be concurrently pregnant and lactating and produce litters of 1–8 offspring (median 4–5), depending on mothers’ age (Migula, 1969; Tkadlec & Krejčová, 2001), every 18 days. Parental care is provided by the female alone. As all vole species, common voles are highly depredated by avian and mammalian predators (Halle, 1988; Norrdahl & Korpimäki, 1998; Norrdahl & Korpimäki, 1995).

To quantify among-individual variation in two behavioral traits, we conducted two repeated laboratory tests. We measured boldness and activity (Réale et al., 2007), which are highly positively correlated at the phenotypic level in common voles; that is, bolder individuals are more active (Eccard & Herde, 2013; Gracceva et al., 2014; Herde & Eccard, 2013; Lantová et al., 2011). Subsequently, we ecologically validated these personality traits by quantifying space use in a grassland, the natural habitat of common voles, and tested the consequences of among-individual differences on survival and reproductive success in large outdoor enclosures in experimental populations, which also provided a social environment to the animals. Space use of animals was monitored with automated radio telemetry (ART, e.g., Hoffmann et al., 2018; Kays et al., 2011; Schirmer et al., 2019), and risk taking via radio frequency identification (RFID) systems placed at risky locations. The combination of different methods
should allow to complement their respective limitations in temporal or spatial accuracy and detection biases. We tested for relationships between among-individual differences in boldness on differences in survival probability, risk taking, space use, and reproductive success over 5 weeks under near-natural conditions. The study period of 5 weeks covers a substantial proportion of an average adult vole’s life span of weeks to months (Halle & Stenseth, 2000).

We predicted that individual differences in boldness and activity, quantified in standardized tests in the laboratory, translate into behavioral differences in space use and risk taking under near-natural conditions. Specifically, we predicted that bold/active individuals occupy larger home ranges and core areas than shy/inactive individuals, as shown for other taxa including birds (Minderman et al., 2010) and small mammals (Boon et al., 2008; Schirmer et al., 2020). Further, we predicted that bold/active individuals—in contrast to shy/inactive individuals—use unsafe open areas at the edge of the suitable habitat patches in large outdoor enclosures (i) with a higher propensity, (ii) a higher frequency, and (iii) longer duration because boldness predicts risk taking (Dammhahn & Almeling, 2012) and dispersal propensity (Cooper et al., 2017) in other small mammals.

We further expected lower survival of bold/active individuals compared to shy/inactive conspecifics because high levels of risk taking and activity may lead to increased predation (meta-analysis: Smith & Blumstein, 2008; but see Moiron et al., 2020). Elevated exploration and activity might pose a high predation risk, but may result in more encounters with the other sex, or increase attractiveness, and, thus, result in reproductive gains (Ophir et al., 2008; Smith & Blumstein, 2008; Sih et al., 2014; but see Araya-Ajoy et al., 2016). Contrarily, high activity levels could also be advantageous if they are connected to the speed of exploration, like in eastern chipmunks (Tamias striatus) where fast explorers had lower mortality compared to slow explorers, probably because they had increased information about the environment (Bergeron et al., 2013). Our study period included two reproductive cycles of common voles, and we quantified the number of offspring produced during these cycles via genetic parentage analysis. We expected bold/active males to sire more offspring than shy/inactive males. Similarly, we expected bold/active females to have higher reproductive success because they might occupy larger (or better quality) ranges (Schirmer et al., 2019). Moreover, since boldness and exploration correlate in common voles (Herde & Eccard, 2013), bold/active females might provision more food to their offspring than shy/inactive individuals as shown for more explorative blue tit (Mutzel et al., 2013).

### Table 1 Schedule of experimental and biological events for common vole subjects in the laboratory and in outdoor grassland enclosures; (a–c) refer to accompanying sections in the methods part

| Experimental time in days | Experimental events | Location |
|---------------------------|---------------------|----------|
| -50                       | (a) Capture and 3–6 weeks of acclimatization | Laboratory |
| -7 to -2                  | Behavioral testing in cohorts of 24 animals | |
| -1                        | Assembly of experimental populations | |
| (b) Application of PITs and radio collars for ART, tissue sampling | |
| -2 to 0                   | Release of four males each into grassland enclosures | Enclosure |
| 0 to 4                    | Release of four females each into grassland enclosures | |
| 3 to 15                   | Exploration phase (Exp) | |
| 16 to 20                  | 1st pregnancy of females (Grav1) | |
| 20 to 35                  | Parturition of 1st litters, postpartum estrus of females and 2nd mating (Mate) | |
| From 35 on                | 2nd pregnancy of females while nursing a litter (Grav2) | |
| -2 to 38                  | Weaning of first litters | |
| (c) Start of removal from enclosures, tissue sampling of first litter (weanlings) for parentage assignment | |
| Continuous measurements of movement and risk-taking behavior | |
| 38 to 40                  | Parturition of 2nd litters | Laboratory |
| 56                        | Weaning of 2nd litters, tissue sampling | |
| 70                        | Release to capture locations | |
2.1.1  | (a) Composing experimental populations

We captured a total of 168 adult common voles between April and August 2010 (N = 120) and 2011 (N = 48), using baited live traps (Ugglan special No2, Grahnab, Sweden; with shrew exits, Eccard & Klemme, 2013) from meadows around Potsdam, Germany (52°26’21.83”N, 13°00’44.14”O). Animals were housed singly at room temperature 18–23°C and natural seasonal photoperiod in standard rodent cages (Ehet GmbH, Germany, Makrolon Type III: 42 cm x 27 cm x 16 cm). Cages contained pellet food, potatoes, and hay ad libitum, plus wood shavings and paper rolls for shelter. Capture and housing were conducted under permission of the Landesumweltamt Brandenburg (ref. RW-7.1 24.01.01.10), and experiments were performed under the permission of the Landesamt für Umwelt, Gesundheit, und Verbraucherschutz Brandenburg (LUGV ref. V3-2347-44-2011).

All captured individuals were subjected to a battery of repeated behavioral tests to assess the correlational structure of behavioral variables and temporal consistency of among-individual differences, that is, animal personality; these results are presented elsewhere (Herde & Eccard, 2013). To compose experimental populations of extreme behavioral types, we used a subset of these individuals and based our selection (see below for more details) on among-individual differences in activity and boldness quantified in two behavioral tests, the barrier test and the open field test. All details are in Herde & Eccard, 2013; here, we only briefly describe these test. Both test types were performed in the housing rooms, in small arenas, lasted 5 min each, and were conducted within days (not on the same day). We repeated both test types after two weeks. In the barrier-test, animals were transferred to one compartment of a two-compartment plastic box and we measured two behavioral variables: (i) the latency to cross a 2.5 cm barrier into the unknown compartment, and (ii) the frequency of crossing the barrier (expressed as crossings per minute, subtracting the latency to first move). The open field test was performed in a round arena of 1 m diameter, and we measured two behavioral variables: (i) the latency to leave the safe wall zone (defined as a zone of 10 cm width along the wall) and enter the unsafe center zone and (ii) overall activity. In a focal observation of the individual, we recorded via instantaneous sampling and a sampling interval of 10 s whether an individual was active (defined as all types of movement except for cleaning) or inactive, yielding 30 sampling intervals over 5-min test duration. As detailed in Herde & Eccard, 2013, all behavioral variables were repeatable over time; latencies and activity variables (frequency of crossing and activity in the open field), respectively, were correlated across the two test types; and combined scores for activity and boldness were correlated at the phenotypic level, across all animals tested in the laboratory, male voles were more active than females but sexes did not differ in latencies. The individual time in captivity before the first behavioral test was performed did not explain variation in boldness nor activity (R² < .03, for all four variables).

Behavioral testing started 3–6 weeks after the animals were captured (to ensure that females were not pregnant, and that pregnant females were able to give birth, raise, and wean the litter), and as soon as a cohort of 24 animals had been collected (12 individuals per sex). For logistical reasons, we had to base the selection of individuals for experimental populations on subsequent cohorts of captured individuals. From each test cohort of 24 individuals, we subsampled 16 animals for two experimental populations run in parallel in two enclosures. To select individuals, we ranked same-sex animals based on the values of behavioral variables obtained in the first test round, that is, according to their latencies (shortest = lowest rank) and activities (most active = lowest rank). From the four behavioral variables, we calculated a raw rank for each individual within its test cohort. The four males and four females with the lowest mean ranks (thereafter called bold) and the four males and females with the highest mean ranks (thereafter called shy) were used for this experiment and assigned alternatingly by rank to two populations, each consisting of two bold males, two bold females, two shy males, and two shy females. Animals with medium ranks were released at their capture location. Binning into a bold and a shy category was used to be able to compare extreme phenotypes (bold/shy) in replicated populations, reflecting our original hypotheses. Since we had removed intermediate phenotypes from the setup, we refrained from correlating obtained variables to original behavioral values, personality scores, or ranks. This approach potentially limits our ability to detect gradual effects of behavioral phenotypes or specifics of extreme versus intermediate behavioral phenotypes.

Across all test cohorts, absolute values of behavioral variables from individuals classified as bold (n = 56) differed from those of individuals classified as shy (n = 56 + 1 additional shy animal released to the first enclosure by mistake. Student’s t tests for all behavioral variables 2.4 < t < 7.8; all p < .02, Tables S1 and S2). Also body weights differed among types, with animals classified as shy being 10% heavier than animals classified as bold (t < 2.0, p < .049, Table S1).

2.1.2  | (b) Experimental populations in large near-natural grassland enclosures

Experimental populations were kept for five weeks between June and November in one of six large grassland enclosures of 50 m x 50 m each. Each enclosure was fenced with a galvanized metal wall (1 m below and 0.5 m above surface). Enclosures were protected against ground predators by a veterinary fence (2 m height) and an electrical fence, but were open for natural avian predation. Ugglan live traps were set in a regular five by five grid with 10 m distance to recapture individuals at the end of the experiment. Two to four experimental populations were operated in parallel, resulting in 14 population replicates (ten in 2010, four in 2011). Enclosures had a built-in automated radio telemetry system (ART, Figure 1a), consisting of pairs of four-element Yagi antennae (Winkel-Spezialantennen, Germany) mounted on a stand of 3.2 m height in each corner, connected via subterraneous cables to an eight-channel automated receiving unit for each enclosure (ARU; JDJC Corp., Sparrow systems, Illinois).
Vegetation along the inside of the enclosure walls was mowed in a strip of 1.5 m width to prevent animals from climbing (Figure 1a). The strip was kept short by regular hand mowing, and intervals depended on local vegetation height and rain patterns. Mowing did not kill any of the collared voles. In the mowed area, the perceived avian predation risk is high for a ground dwelling mammal (Jacob & Brown, 2000).

2.1.3 | (c) Behavior in the field

Before the release into grassland enclosures, each animal was marked with a unique passive integrated transponder (PIT; 0.1 g; Trovan ID-100, Euro ID, Germany), placed subcutaneously in the scapular region, for individual recognition at RFID readers. Animals were fitted with a radio telemetry collar (Biotrack, UK; 1.0 g including cable tie, <5% of mean body mass before release) with an individual radio frequency.

We placed barriers (Figure 1b) perpendicular to the enclosure wall into the vegetation free strip, guiding a passing animal through a passage (diameter 4.5 cm) monitored by an RFID ring antenna (diameter 5.5 cm) with a detection range at ca. 1 cm before and after the ring, and a storing unit (LID 650 Euro ID, Germany). By integrating this system into a drift fence, we assumed to detect individuals that moved in the open areas and along enclosure walls, while short excursions into the open area were not detected. The system recorded each individual RFID code with a time stamp every 0.1 sec. Animals were classified as "visitors" (at least one reading in the edge zone at one of four antennae in 35 days) and "nonvisitors" (no reading). For visitors, we further analyzed number of visits (two visits were separated by a minimum of 5 min between two readings) and mean duration of visits. The definition of visit bout length was based on a pilot study filming vole behavior at the barriers.

Males were released two days prior to females to display potential differences in exploration behavior and avoid an immediate associate with locations of females (Im schau, 1988). Automated Receiving Units (ARU) logged signal strength for each radio tag's frequency on each antenna. In 2010, we logged frequencies in parallel (each frequency once every 2 min) and calculated a location integrating 48 signals across the antenna array over 12 min. After learning from the RFID data in 2010 that individuals could potentially (one animal) cover the length of the enclosure in the duration between two location fixes, we changed the logging rhythm in 2011 to sequential logging; integrating the same number of signals over 1.5 min per animal before switching to the next frequency, while keeping the logging interval of 12-min constant between years. Within each antennae pair, we converted the distribution of median signal strengths from 12 signals into a bearing. Locations were calculated with triangulation of these bearings (see Hoffmann et al., 2018). We removed data after receiver failures (three populations) or antenna failures (two populations), and during rainy periods within replicates (poor transmission through wet vegetation). Overall, we obtained $N = 1004$ telemetry days with 90–120 locations per day for 58 individuals in nine populations (mean ± SD: 20 ± 10.4 per individual, min-max range: 3–35 day ranges). We conducted calibration, precision, and maintenance checks on the telemetry grids before, in the middle, and after each replicate, using stationary experimental tags. Calibrations lasted ca. 1 h leaving sufficient time to collect locations.
for the respective day ranges. We found that the absolute day range sizes varied greatly between populations (ANOVA of 95% Kernels: 
\( F = 135, p < .001, df = 8; \) population means from 348 m² to 1168 m²; data were square-root-transformed beforehand), and also the precision of single locations varied between experimental populations (5 to 15 m) due to seasonal changes in vegetation height, wind, or moisture. To account for differences among replicates, we obtained z-scores within populations, relating the individual day range to the respective population mean (in: percent of population mean). We calculated day ranges (i.e., 95% density Kernels) and core area of day ranges (i.e., 50% density Kernels) as estimates of individual vole movement and mobility (Worton, 1987, 1995). Since all analyses conducted with both kernel sizes yielded very similar statistics (both were based on the same location data set), we present only the statistical results on day ranges here. We hypothesized that behavior in the field also depends on the social interactions in the experimental population. Particularly, mobility of males may vary with the availability of mating partners, which would be low during synchronized pregnancy phases of females. Further, we hypothesized that boldness types may cope differently with being released to the unknown habitat (Veerbek et al., 1994). Therefore, we included the following experimental phases based on the species’ life history into the analysis of location data (Table 1): (1) Expl: exploration phase (days 0–3), when animals explore the habitat and unknown conspecifics and mating occurs (females were introduced nongravid) resulting in synchronous reproduction cycles; (2) Grav1: the first pregnancy when females were synchronously gravid (days 4–15); (3) Mate: a phase during which females give birth and mate again postpartum (days 15–20) with males presumably increasing mobility to roam between females; and (4) Grav2: a second pregnancy phase when females were supposedly gravid again (days 20–35).

2.1.4 | (d) Fitness

To remove adults and their offspring from the enclosures, we set live traps after 35 days (Table 1). It took up to 5 days until animals were removed. Individuals without a mobile radio tracking signal that were not recaptured during removal trapping or did not reappear in a later replicate (6 cases) were considered to be dead. To estimate reproductive success, we collected small tissue samples from the ears of adult voles before release to the enclosures, from offspring born in and captured from the enclosures (\( N = 335 \) juveniles), and from offspring born to females kept in cage cages after the experiment (\( N = 85 \) juveniles). Laboratory procedures for genotyping followed Braaker and Heckel (2009; Table S3). Microsatellite alleles were determined using Genemapper® Software, version 3.7 (Applied Biosystems). The number of alleles ranged between two and 32 (mean = 13.9) per microsatellite locus. We used the software CERVUS 3.0.3 (Kalinowski et al., 2007; Marshall et al., 1998) and individual parental candidate exclusion for parentage identifications (details in Table S3).

From the analysis, we obtained 421 parental assignments for 276 offspring and identified both parent candidates for 64% of offspring and one parent candidate for an additional 17.4% of offspring. To obtain robust estimates of relative individual reproductive success of a parent within a population, we included replicates with 16 parentages assigned (8 replicates). The excluded replicates had <8 parental assignments, and we excluded assignments to parents outside their original replicates. With these limitations we were able to use data of 346 assignments for 258 offspring to 57 parental candidates. There were six animals among those candidates where no offspring was assigned, but which had been able to potentially sire offspring, as indicated either by their recapture after the experiment (\( n = 3 \)) or the polyphasic activity signature of their radio signals (\( n = 3 \), see supplemental material for exemplary diagnostic plots).

2.1.5 | (e) Statistical analyses

To test whether boldness type explained variation in risk taking, space use, survival, and reproductive success, we used linear or generalized linear mixed effects models (LMM or GLMM) run with the R package “lme4” (Bates et al., 2015). The underlying error distributions were specified as binomial for probabilities (survival, reproduction, visits of risky areas), as Poisson for count variables (number of visits of risky areas, number of offspring) and as Gaussian for continuous variables (home range size, core area size, duration of visits of risky areas). Given a biased distribution as based on visual inspection, we log-transformed the duration of visits. Models included our predictor boldness type (shy or bold) and sex (male or female) as fixed effects, and their interaction. Further, we included control variables as fixed effects into initial models: the starting month of the replicate (to control for seasonal variation as covariate) as a continuous covariate, and the experimental phase (with four levels, see Table 1 for details) for models on space use only. Experimental phase was specified in interaction with boldness type and sex because we expected space use to vary with the phases of the experiment and in particular with female reproductive activity in our artificially reproductively synchronized populations. As random effect, we included population replicate ID (specified as random intercept) to control for potential variation among replicates such as vegetation height, rain events, or predation pressure. Such external properties could potentially affect the behavior of the entire population. Furthermore, vegetation and weather may affect the quality of our tracking calibration, the recapture success. For exploration of the data, we experimented with different random structures (e.g., including the identity of the enclosure or adding the year as a fixed factor, but population ID as random factor captured this variation). In models of space use and risk taking, we had repeated measurements of individuals and therefore added individual ID as a second random effect to the mixed models.

We assessed model fit visually based on inspections of residual distribution (homogeneity of variances, normal distribution) and calculated conditional and marginal coefficients of determination (\( R^2 \)) using the R package MuMIn (Nakagawa & Schielzeth, 2013).
Predictors reflecting the experimental setup (boldness type and sex) were always kept in the final model. Based on log-likelihood ratio tests, control variables, covariates, and interaction of factors were removed if they did not increase the predictive value of the model. For behavioral variables from the field (risk taking and space use), we estimated repeatability over time using the R package rptR (Nakagawa & Schielzeth, 2013, Stoffel et al., 2017), using 1000 simulations to estimate confidence intervals and 1000 permutations to estimate p-values. Analyses were carried out with R, Version 3.0.2 (R Core Team, 2016).

3 | RESULTS

3.1 | Behavior in the field

Risky areas were visited 367 times by 44 of 113 individuals (39%), which visited at least one guided passage in the vegetation free zones of the enclosures. Females were less likely (yes or no) to visit the risky area (12 of 57 females) than males (32 of 44, $\chi^2 = 14.4, p < .001$, Figure 2a, Table 2) and visitation probability of an animal was independent of its boldness ($\chi^2 = 0.4, p = .524$). Among visiting animals males visited more often (7.4 ± 10 visits) than females (3.5 ± 6.2 visits, $\chi^2 = 14.1, p < .001$, Figure 2b), and bold individuals more often (8.3 ± 11.9) than shy individuals (4.7 ± 6.1 visits, $\chi^2 = 22.3, p < .001$), and the number of visits was higher in replicates later in the season (month; $\chi^2 = 4.8, p = .029$, Table 2). The duration of a visit was highly repeatable within individuals ($R = .632, CI 0.50–0.73$) and was depending on an interaction of experimental phase and sex ($\chi^2 = 17.53, p < .001$, Table 3, Figure 3a). Visited risky areas for shorter periods during the first pregnancy phase (Grav 1) compared with the exploration phase (Expl) and the second pregnancy (Grav 2). In males, we detected no effects of experimental phase or boldness type.

Daily home ranges ($n = 1004$ ranges, 58 animals) were highly repeatable within individuals (for $R_{adj} = .44, CI: 0.32–0.54$, $p < .001$). Range size was explained by an interaction of boldness type, sex, and experimental phase (three-way interaction: home range $\chi^2 = 22.9, df = 3, p < .001$, effect sizes Table 3, Figure 3). To disentangle the three-way interaction, we analyzed simple effects within different subsets. Among bold animals ($n = 30$ individuals of both sexes, $n = 553$ daily core areas), experimental phase affected day range size ($\chi^2 = 23.9, df = 3, p < .001$, Figure 3b) with smaller day ranges during the exploration phase (mean ± SD: 88% percent of respective population mean ± 45%), which differed from the first pregnancy phase (105% ± 55%) and the parturition and second mating phase (111% ± 55%) but not from the second pregnancy phase (98% ± 48%). Among shy animals ($n = 28$ individuals, $n = 451$ day ranges), sex differences varied between experimental phases (interaction $\chi^2 = 22.1, df = 3, p = .001$, Figure 3b). Within females, boldness type explained day range size depending on experimental phases (interaction: $\chi^2 = 12.2, df = 4, p = .006, n = 31$ females, $n = 542$ daily home ranges). Separate analyses of main effects within combinations of boldness types by sex revealed that shy females ($n = 15$ individuals, $n = 271$ daily home ranges) had larger day ranges during the exploration phase (106% ± 58%) and first pregnancy (108% ± 36%), and size decreased during later phases (mating: 92% ± 20% and second pregnancy: 79% ± 32%; experimental phase: $\chi^2 = 54.0, df = 3, p < .001$, Figure 3b). In contrast, bold females ($n = 16$ individuals, $n = 272$ daily home ranges) used smaller day ranges during exploration phase (86% ± 35%) compared to later phases (mean 95% 105%, experimental phase: $\chi^2 = 17.5, df = 3, p < .001$; Figure 3b). Across all phases, day ranges of males were larger (104% ± 40%) than those of females (94% ± 44%, $\chi^2 = 7.2, p = .007$, Table 4). Males’ ($n = 27$ individuals, $n = 461$ days) range sizes varied with experimental phase ($\chi^2 = 16.1, df = 3, p = .001$) but not with boldness types ($\chi^2 = 0.1, df = 1, p = .789$, interaction not significant). Males had the smallest day ranges during the exploration phase (87% ± 52%) compared to later phases (mean 103%–111%, Figure 4).

3.2 | Fitness of animals

In total, 73 of 113 (65%) released common voles were recaptured from the enclosures. Females tended to survive better (73%, Table 2, Figure 4a, mean ± SD: 2.9 ± 1.0 per population) than males (57%, 2.3 ± 1.3 per population, $\chi^2 = 2.8, p = .094$), and survival tended to decrease with month from 88% survival in June replicates to 35% in
### Table 2: Risk taking (visiting short vegetation areas of enclosures), survival, and reproductive success of common voles in experimental populations

| Dependent variable (model, error distribution) | Model estimates | Explained variance |
|-----------------------------------------------|-----------------|--------------------|
|                                              | Intercept       | Boldness type (shy) | Sex (male) | month | Boldness type*Sex | $R^2_{\text{marginal}}$ | $R^2_{\text{conditional}}$ |
| Visitation (probability, binomial) $n = 113$, 14 populations | $\beta \pm SE$ | $1.80 \pm 0.66$ | $0.26 \pm 0.41$ | $-1.60 \pm 0.42$ | ns | ns | 0.4 | 0.4 |
|                                              | $z$             | 2.7               | 0.64         | -3.80   | |
|                                              | $p$             | 0.008             | 0.52         | <0.001  | |
| No. of visits (Poisson) $n = 44$, 14 populations | $\beta \pm SE$ | $1.30 \pm 0.30$ | $-0.60 \pm 0.13$ | $0.79 \pm 0.21$ | $0.26 \pm 0.11$ | ns | 0.11 | 0.4 |
|                                              | $z$             | 5                 | -4.6         | 3.5     | 2.1 |
|                                              | $p$             | <0.001            | <0.001       | <0.001  | 0.029 |
| Survival (probability, binomial) $n = 113$, 14 populations | $\beta \pm SE$ | $4.04 \pm 2.18$ | $-0.52 \pm 0.44$ | $-0.74 \pm 0.44$ | $-0.47 \pm 0.25$ | ns | 0.25 | 0.57 |
|                                              | $z$             | 1.85              | 1.17         | 1.67    | 1.99 |
|                                              | $p$             | 0.64              | 0.224        | 0.94    | 0.06 |
| No. of offspring (Poisson) $n = 57$, 8 populations | $\beta \pm SE$ | $1.31 \pm 0.17$ | $0.22 \pm 0.18$ | $0.37 \pm 0.17$ | ns | $-0.61 \pm 0.25$ | 0.02 | 0.12 |
|                                              | $z$             | 7.44              | 1.25         | 2.12    | -2.40 |
|                                              | $p$             | <0.001            | 0.211        | 0.034   | 0.017 |

**Note:** Shown are model estimates of GLMMs based on different sample sizes of animals ($n$, given for each model) and different numbers of populations, which were included as a random factor. Covariate and interaction were removed if $p > .1$. The reference levels for categorical predictors are bold (for boldness type) and female (for sex). Shown are estimates ($\beta$) and their standard errors (SE), z-values and p-values as well as $R^2_{\text{marginal}}$ as the variance explained by fixed factors, and $R^2_{\text{conditional}}$ as the variance explained by fixed and random factors. Significant effects are marked with bold font.
October replicates in October, $\chi^2 = 3.5, p = .060$). Boldness type did not explain variation in survival ($\chi^2 = 1.4, p = .242$, Figure 4a, effect sizes in Table 2). During the time in the enclosures, animals gained on average 12.3 g body mass (mean ± SD body mass after the experiment: males 38.1 ± 6.5 g; females 32.7 ± 5.9 g, including gravid females).

Fifty-seven experimental animals were parental candidates, and offspring were assigned to 51 of them (89%). The number of offspring assigned to an individual ranged from 0–14 and was predicted by an interaction of sex and boldness type ($\chi^2 = 5.7, p = .017$, effect sizes Table 2, Figure 4b). Follow-up analysis within sexes (GLMM of offspring numbers, Poisson error distribution) showed that overall, bold males produced more offspring ($5.6 ± 4.6, n = 14$) than shy males ($3.9 ± 2.5$ offspring, $n = 13$), simple effects within males: $\chi^2 = 4.4, p = .036$, effect size $\beta = -0.38 ± 0.18$, Figure 4c shows distribution of offspring per male within the 8 populations analyzed). In females, boldness types did not predict reproductive output ($4.3 ± 3.1$ offspring, $\chi^2 = 1.4, p = .227$, $\beta = -0.22 ± 0.18, n = 15$ females per type, Figure 4b).

### 4 | Discussion

In experimentally created populations of known behavioral-type composition, we were able to show that individual differences in boldness behavior covaried with risk taking, space use, and fitness under near natural conditions. We created experimental populations combining the opposite ends of the distribution of behavioral phenotypes (bold and shy animals) and studied behavior in the field with automated tracking methods. We found that behaviors measured in the field were consistent within individuals over time, quantifying themselves for animal personality traits. We further found that bold animals of both sexes visited the risky edges of the enclosures more frequently than shy animals of the same sex. In females, effects of boldness type were detected during limited times only (Figure 4). While males ranged over larger areas than females (as shown earlier for this species, e.g., Briner et al., 2005), range sizes differed between shy and bold females immediately upon release. Shy females apparently explored larger areas initially and then settled in smaller ranges, while the opposite pattern was observed in bold females and males of both behavioral phenotypes. Bolder males took higher risks and fathered more offspring than shy males. Boldness did not explain survival probability in both sexes, however. Mortality of voles tended to increase in autumn (Figure 2a), probably due to colder weather and decreasing quality of forage, mirroring annual population dynamics of common voles (Eccard & Herde, 2013).

#### 4.1 Behavior and boldness types

Boldness as measured in many small mammals in laboratory settings using open field and exploration tasks may be a direct predictor of
risk taking (Dammhahn & Almeling, 2012). The main source of mortality for voles is predation, rendering them a key species in natural food chains with many ground and avian predators preying on them (Halle, 1988; Jędrzejewski & Jędrzejewska, 1993; Norrdahl & Korpimäki, 1995). Since boldness may be directly linked to mortality risk (Smith & Blumstein, 2008, but see Moiron et al., 2020), it should have a strong impact on spatial behavior, and the exposure to predators.

In the experimental populations in our study, bold males were taking higher risks by visiting the short vegetation edges of the enclosure more frequently than shy males. Boldness is often correlated with exploration at the phenotypic level, so bolder individuals were often reported to explore an area faster than shy individuals, which could result in a more superficial exploration and exploitation of resources (Mazza et al., 2018; Sih et al., 2004; Wolf et al., 2007a; Wolf et al., 2007b). In our study, we found some support for this pattern with bold males being registered in the risky area of the enclosure more often and for shorter periods than shy males, probably indicating a quicker and superficial exploration of these areas. The duration of single visits did not differ among males, but among females. Females visited the risky edge less frequently than males, and stayed very shortly at the passage counter during their first pregnancy, compared to the exploration phase and the second pregnancy (Figure 3). Since long stays indicated a slower and more careful passage, as indicated by our pilot experiment, we assume these phases are used for exploration of the novel area, or to find a new nest for giving birth to the second litter, while during the first pregnancy females passed the counters quickly and on paths known to them. Females never appeared at the passage counters during the second mating phase, probably because voles mate briefly during

FIGURE 3  Behavior of common voles in large grassland enclosures over 7 weeks, gray: boldness type “bold,” white: boldness type “shy”. Experiments were divided into four phases based on female reproductive biology: Expl: exploration of novel environment including social environment and mating (3 days); Grav1: first pregnancy (15 days), Mate: parturition of litters and mating after postpartum estrus (5 days), Grav2: second pregnancy (13 days). (a) Number of visits at low vegetation (risky) areas of the enclosures. (b) Duration of visits (n = 358 visits by 44 common voles in 14 populations) at low vegetation (risky) areas of the enclosures. Long visits indicate a careful and slow movement at the passage counter, short visits a quick passage. Missing observations: no visits. Width of bar indicates relative sample size. (c) Model predictions for relative size of 1004-day ranges (95% Kernel estimates) in relation to the respective population mean (= reference line: 100%) for 58 common voles (ID included as random effect) from nine enclosed populations (population included as random effects). Each day range was computed based on 90–120 location fixes per individual over 24 h. Asterisk refers to post-hoc differences at p < .05 (compare Table 3)
Directly after transfer to the novel environment (exploration phase), bold females and both types of males in our experimental populations used smaller daily ranges than at later stages of the experiment, indicating an initial reduction in mobility. Shy females used larger areas during the first three days, and settled in areas that later allowed them to maintain small home ranges (Figure S1). At first glance, this differs from established populations in different species where bold animals (of both sexes) had larger ranges than shy ones (rodents: Boon et al., 2008; Schirmer et al., 2020, birds: Minderman et al., 2010). Our finding is more in line with observations of shy animals being more thorough explorers in novel environments (Marchetti & Drent, 2000; Mazza et al., 2018; Mutzel et al., 2013; Veerbek et al., 1994), which might give them an advantage under changing and harsh environmental conditions.

4.2 | Fitness and boldness types

As predicted, bold males overall sired more offspring than shy males, although not in all populations. In line with our results, boldness scales positively with reproductive success in many species (Collins et al., 2019; Dingemanse & Réale, 2005; Godin & Dugatkin, 1996; Reaney & Backwell, 2007; Scherer et al., 2020; Smith & Blumstein, 2008). Several nonexclusive mechanisms might explain fitness benefits of being bold. (1) Bolder males might take higher risks in roaming in space to find receptive females and/or defend receptive females more successfully to monopolize paternity (Ophir et al., 2008; Smith & Blumstein, 2008; Wolf et al., 2007a; Wolf et al., 2007b); indeed, in our study bolder males were detected more often in risky areas of the enclosures. Appearance in such areas is sometimes used to infer dispersal tendencies (Hahne et al., 2011) and may also indicate wider roaming areas (Schirmer et al., 2020; Ward-Fear et al., 2018). (2) Females could have a preference for bolder males (Godin & Dugatkin, 1996). If boldness was selected for in males, we would expect males to generally be bolder than females (Schuett et al., 2010), which is not supported by our other studies on voles (M. arvalis: Eccard & Herde, 2013; Herde & Eccard, 2013; Myodes glareolus: Mazza et al., 2018; Schirmer et al., 2019). (3) Reproductive success of males may be primarily determined by dominance rank (Dewsbury, 1982; Ellis, 1995) rather than personality per se, but both traits can be highly entangled so that boldness may predict dominance. (4) Among-individual variation in behavior could be part of a larger pace-of-life syndrome (Dammhahn et al., 2018; Réale et al., 2010) and covariance between these traits might be maintained by density-dependent selection (Milles et al., 2022; Wright et al., 2019). Microtine voles, in most places, frequently and predictably undergo massive fluctuations in population density, which are accompanied by population-level differences in behavioral type (Eccard & Herde, 2013) and social environmental conditions. Further, for short-lived iteroparous animals in seasonally fluctuating environments, life history trajectories and social environmental conditions (e.g., density) are predictable (Eccard et al., 2017; Eccard & Herde, 2013). Selection may favor bolder behavioral types in high density and high

FIGURE 4 Fitness of common voles in experimental populations (a) survival of males and females (initial numbers: 4 each per population) in 14 populations over the season, year 1 solid lines, year 2 dashed lines, (b) number of assigned offspring per individual (8 populations, 57 parental candidates, 16–44 offspring per population assigned) at a significance level of p < .05 (asterisks), (c) number of assigned offspring per male, ∆ bold male, ○ shy male, gray lines = 8 populations sorted by months, year 1 white, year 2 gray symbols

a postpartum estrus and females may have spent this time close to their nests nursing new borne offspring.

Individual daily range sizes were highly repeatable over time, indicating intrinsic individual differences in space use. Similarly, home range and core area size, as well as microhabitat characteristics of bank voles (Myodes glareolus) and striped field mice (Apodemus agrarius)—automatically tracked under natural conditions—covaried with individual behavioral differences (Schirmer et al., 2020). Thus, overall among-individual differences in space use may contribute to individual niche specialization (Pearish et al., 2013; Spiegel et al., 2017), facilitating the coexistence of similar species (Schirmer et al., 2020). The distribution of individuals in space and time is an important determinant for key aspects of the social system, for example, the mating system (Heckel & von Helversen, 2002, 2003; Lukas & Clutton-Brock, 2013), and of foraging under risk. Hence, behavioral type-specific space use should have consequences for survival and reproductive success.
competition phases of the yearly fluctuation cycle, which ought to express a more risk-prone pace-of-life syndrome (Herde & Eccard, 2013). Shy behavioral types and risk-averse pace-of-life syndrome may potentially be favored by higher survival at low densities in winter or early in the breeding season, and subsequently have a higher contribution to the increasing population in spring than bold individuals. In our data, in the replicates run earlier in the breeding season shy males had the highest reproductive success (Figure 2c), while later in the season, when in wild, natural populations densities would be high, reproductive success was skewed in favor of single, bold males. This may indicate a density-dependent selection of different pace-of-life syndromes in fluctuating populations, triggered by seasonal cues. Future studies should test the relationships between social environment, predictable seasonal life history trajectories, animal personality, and fitness.

We did not detect an effect of boldness type on reproductive success in females. Thus, the fitness consequences of boldness might be sex-specific, similar to the findings in black banded albatrosses (Patrick & Weimerskirch, 2014). Access to food and safety are major determinants of reproductive success in female mammals (Crook & Gartlan, 1966; Emlen & Oring, 1977; Lukas & Clutton-Brock, 2013; Ophir et al., 2008; Terborgh & Janson, 1986); since common voles mainly eat grass and find shelter in underground burrows, our large grassland enclosures should not have provided a resource limited environment. Further, reproductive skew is generally lower in females than in males (Bateman, 1948) and once female voles reproduce, they usually produce entire litters. We expected bold females to occupy larger home ranges (as bank voles under natural conditions: Schirmer et al., 2019) and thus be able to provision their offspring better (e.g., as in blue tits: Mutzel et al., 2013) compared to shy females. However, differences in provisioning (lactation) would be difficult to detect among different types of female mammals in an outdoor study.

In contrast to our prediction but in line with results of a recent meta-analysis (Moiron et al., 2020), survival did not differ between boldness types. The finding may be caused by the rather benign setting of our experiment in a favorable season, a low population density, and reduced predation pressure since ground predators were excluded. Alternatively, limited space might be another explanation, since male voles might roam larger areas under natural conditions than offered in our enclosures. Overall, the survival rate of common voles in our study (35% over 7 weeks) seemed high compared to those reported elsewhere: 2 to 9% daily mortality of voles with radio transmitters (field voles, East European voles and bank voles; Norrдаhl & Korpimäki, 1995), or 50% mortality over four weeks in agricultural fields (common voles; Jacob, 2003). In our experiment, survival dropped toward the end of the season (Figure 2a) for animals of any boldness type, when in wild populations peak densities would crash (Eccard & Herde, 2013) and the adult animals captured during summer would reach the end of their life span. Meanwhile, if in voles mortality would follow a disruptive viability selection, such as found in Eastern chipmunks (Tamias striatus), and both high and low extremes of behavioral types would lead to intermediate types (Bergeron et al., 2013), we would not be able to detect this pattern since we selected extreme boldness types from the ends of a behavioral gradient.

5 | CONCLUSIONS

Overall, our results highlight that among-individual differences in behavior translate into variation in space use, risk taking, and reproductive success in near-natural populations. Reproduction was biased toward single bold males in late summer replicates. Since variation in boldness is maintained in natural populations, we assume that shy types may have fitness advantages in other seasons (Lonn et al., 2017) or at different population densities (Wright et al., 2019), which remains to be tested. With daily range sizes being highly repeatable within individuals, consistent individual space use patterns may facilitate individual niche specialization and thus affect among- and between-species ecological interactions. We show with this experiment, that behavioral phenotypes covary with risk-taking behavior in the field, and that behavioral differences are thus expressed in natural settings. We can further show that behavioral phenotypes are fitness relevant.

ACKNOWLEDGMENT

We thank all students and student helpers who trapped and observed common voles and helped in the enclosures. Thanks to Jennifer Schöpf who conducted the RFID-pilot tests in the laboratory. For technical support, we thank Elke Seydewitz, Angela Puschmann, Annika Schirmer, and Anne Gärtnner. Many thanks to Fritz Trillmich and Elke Hippiauf for help with the first steps of the paternity analysis at the paternity platform of the research group (FOR 1232) in Bielefeld. Financial support was provided by the German Science Foundation (DFG) with a grant to JAE (EC/361-6) in the research unit “Reduction in Phenotypic Plasticity in Behaviour by early Experience” (FOR 1232), and to MD during paper writing (DA 1377/4-1). Open Access funding enabled and organized by Projekt DEAL. This publication was funded by the Deutsche Forschungsgemeinschaft (DFG, German Research Foundation) - project number 491466077

CONFLICT OF INTEREST

Authors declare that there are no competing interests.

AUTHOR CONTRIBUTIONS

Jana A. Eccard: Conceptualization (lead); Formal analysis (equal); Funding acquisition (lead); Methodology (equal); Resources (lead); Supervision (lead); Visualization (equal); Writing – review & editing (equal). Antje Herde: Conceptualization (equal); Data curation (lead); Formal analysis (equal); Investigation (equal); Methodology (equal); Visualization (equal); Writing – original draft (lead). Andrea C. Schuster: Data curation (supporting); Investigation (supporting); Methodology (supporting). Thilo Liesenjohann: Investigation (equal). Tatjana Knopp: Investigation (equal). Gerald Heckel: Formal analysis (supporting); Methodology (supporting); Resources (supporting);
Data are accessible on dryad https://doi.org/10.5061/dryad.44j0z pcfs.

ORCID
Jana A. Eckett https://orcid.org/0000-0002-6151-2128
Gerald Heckel https://orcid.org/0000-0002-0162-323X
Melanie Dammhahn https://orcid.org/0000-0003-0557-740X

REFERENCES
Araya-Ajoy, Y. G., Kuhn, S., Mathot, K. J., Mouchet, A., Mutzel, A., Nicolaus, M., Wijmenga, J. J., Kempenaers, B., & Dingemanse, N. J. (2016). Sources of (co)variation in alternative siring routes available to male great tits (Parus major). Evolution, 70(10), 2308–2321. https://doi.org/10.1111/evo.13024
Arvidsson, L. K., Adriaensen, F., van Dongen, S., De Stobbeleere, N., & Matthysen, E. (2017). Exploration behaviour in a different light: Testing cross-context consistency of a common personality trait. Animal Behaviour, 123, 151–158. https://doi.org/10.1016/j.anbehav.2016.09.005
Bateman, A. J. (1948). Intra-sexual selection in Drosophila. Heredity, 2(3), 349–368. https://doi.org/10.1038/hdy.1948.21
Bates, D., Mächler, M., Bolker, B., & Walker, S. (2015). Fitting linear mixed-effects models using lme4. Journal of Statistical Software, 67(1), 1–48. https://doi.org/10.18637/jss.v067.i01
Bergeron, P., Montiglio, P.-O., Réale, D., Humphries, M. M., Gimenez, O., & Garant, D. (2013). Disruptive viability selection on adult exploratory behaviour in eastern chipmunks. Journal of Evolutionary Biology, 26(4), 766–774. https://doi.org/10.1111/jeb.12081
Bolnick, D. I., Amarasekare, P., Araújo, M. S., Bürger, R., Levine, J. M., Novak, M., Rudolf, H. W., Schreiber, S. J., Urban, M. C., & Vasseur, D. (2011). Why intraspecific trait variation matters in community ecology. Trends in Ecology & Evolution, 26(4), 183–192. https://doi.org/10.1016/j.tree.2011.01.009
Boon, A. K., Réale, D., & Boutin, S. (2008). Personality, habitat use, and their consequences for survival in North American red squirrels Tamiasciurus hudsonicus. Oikos, 117(9), 1321–1328. https://doi.org/10.1111/j.0300-1299.2008.16567.x
Borkowska, A., & Ratkiewicz, M. (2010). Promiscuity, male reproductive success and mate relatedness in a natural population of the common vole. Journal of Zoology, 280(2), 195–201. https://doi.org/10.1111/j.1469-7998.2009.00648.x
Borowski, Z., & Owadowska, E. (2010). Field vole (Microtus agrestis) seasonal spacing behavior: The effect of predation risk by mustelids. Naturwissenschaften, 97(5), 487–493. https://doi.org/10.1007/s00114-010-0663-1
Braaker, S., & Heckel, G. (2009). Transalpine colonisation and partial phylogeographic erosion by dispersal in the common vole (Microtus arvalis). Molecular Ecology, 18(11), 2518–2531.
Briner, T., Nentwig, W., & Airoldi, J.-P. (2005). Habitat quality of wildflower strips for common voles (Microtus arvalis) and its relevance for agriculture. Agriculture, Ecosystems & Environment, 105(1–2), 173–179. https://doi.org/10.1016/j.agee.2004.04.007
Careau, V., Thomas, D., Humphries, M. M., & Réale, D. (2008). Energy metabolism and animal personality. Oikos, 117(5), 641–653. https://doi.org/10.1111/j.0030-1299.2008.16513.x
Collins, S. M., Hatch, S. A., Elliott, K. H., & Jacobs, S. R. (2019). Boldness, mate choice and reproductive success in Rissa tridactyla.

Animal Behaviour, 154, 67–74. https://doi.org/10.1016/j.anbehav.2019.06.007
Cooper, E. B., Taylor, R. W., Kelley, A. D., Martining, A. R., Boutin, S., Humphries, M. M., Dantzer, B., Lane, J. E., & McAdam, A. G. (2017). Personality is correlated with natal dispersal in North American red squirrels (Tamiasciurus hudsonius). Behaviour, 154(9–10), 939–961. https://doi.org/10.1163/1568539X-00003450
Crook, J. H., & Garlan, J. S. (1966). Evolution of primate societies. Nature, 210(5042), 1200–1203.
Dammhahn, M., & Almeling, L. (2012). Is risk taking during foraging a personality trait? A field test for cross-context consistency in boldness. Animal Behaviour, 84(5), 1131–1139. https://doi.org/10.1016/j.anbehav.2012.08.014
Dammhahn, M., Dingemanse, N. J., Niemelä, P. T., & Réale, D. (2018). Pace-of-life syndromes: A framework for the adaptive integration of behaviour, physiology and life history. Behavioral Ecology and Sociobiology, 72, 62. https://doi.org/10.1007/s00265-018-2473-y
Dewsbury, D. A. (1982). Dominance rank, copulatory behavior, and differential reproduction. The Quarterly Review of Biology, 57(2), 135–159. https://doi.org/10.1086/412672
Dingemanse, N. J., Both, C., Drent, P. J., & Tinbergen, J. M. (2004). Fitness consequences of avian personalities in a fluctuating environment. Proceedings of the Royal Society of London. Series B: Biological Sciences, 271(1541), 847–852.
Dingemanse, N. J., & Réale, D. (2005). Natural selection and animal personality. Behaviour, 142(9–10), 1159–1184. https://doi.org/10.1163/15685390577459445
Eckett, J. A., Dammhahn, M., & Vlietin, H. (2017). The Bruce effect revisited: is pregnancy termination in female rodents an adaptation to ensure breeding success after male turnover in low densities? Oecologia, 185, 81–94. https://doi.org/10.1007/s00442-017-3904-6
Eckett, J. A., & Herde, A. (2013). Seasonal variation in the behaviour of a short-lived rodent. BMC Ecology, 13(1), 43. https://doi.org/10.1186/1472-6785-13-43
Eckett, J. A., & Klemme, I. (2013). Reducing mortality of shrews in rodent live trapping—A method increasing live-trap selectivity with shrew exits. Annales Zoologici Fennici, 50(6), 371–376. https://doi.org/10.5735/086.050.0602
Eckett, J. A., Liesenjohann, T., & Dammhahn, M. (2020). Among-individual differences in foraging modulate resource exploitation under perceived predation risk. Oecologia, 194, 621–634.
Ellis, L. (1995). Dominance and reproductive success among nonhuman animals: A cross-species comparison. Ethology and Sociobiology, 16(4), 257–333. https://doi.org/10.1016/0162-3095(95)00050-U
Emlen, S. T., & Oring, L. W. (1977). Ecology, sexual selection, and the evolution of mating systems. Science, 197(4300), 215–223.
Fink, S., Excoffier, L., & Heckel, G. (2006). Mammalian monogamy is not controlled by a single gene. Proceedings of the National Academy of Sciences of the United States of America, 103(29), 10956–10960. https://doi.org/10.1073/pnas.0602380103
Gliwicz, J. (1997). Space use in the root vole: Basic patterns and variability. Ecography, 20(4), 383–389. https://doi.org/10.1111/j.1600-0587.1997.tb00383.x
Godin, J. G., & Dugatkin, L. A. (1996). Female mating preference for bold males in the guppy, Poecilia reticulata. Proceedings of the National Academy of Sciences of the United States of America, 93(19), 10262–10267. https://doi.org/10.1073/pnas.93.19.10262
Graceeva, G., Herde, A., Groothuis, T. G., Koohaaas, J. M., Palme, R., & Eckett, J. A. (2014). Turning shy on a winter’s day: effects of season on personality and stress response in Microtus arvalis. Ethology, 120(8), 753–767.
Hahne, J., Jenkins, T., Halle, S., & Heckel, G. (2011). Establishment success and resulting fitness consequences for vole
Reaney, L. T., & Backwell, P. R. (2007). Risk-taking behavior predicts aggression and mating success in a fiddler crab. *Behavioral Ecology*, 18(3), 521–525. https://doi.org/10.1093/beheco/arm014

Roth, A. M., Dingemanse, N. J., Nakagawa, S., McDonald, G. C., Levieil, H., Robledo-Ruiz, D. A., & Pizzari, T. (2021). Sexual selection and personality: Individual and group-level effects on mating behaviour in red junglefowl. *Journal of Animal Ecology*, 90(5), 1288–1306. https://doi.org/10.1111/1365-2656.13454

Roth, A. M., Firth, J. A., Patrick, S. C., Cole, E. F., & Sheldon, B. C. (2019). Partner’s age, not social environment, predicts extrapair paternity in wild great tits (Parus major). *Behavioral Ecology*, 30(6), 1782-1793.

Saether, B.-E., Visser, M. E., Gruen, V., & Engen, S. (2016). Evidence for r- and K-selection in a wild bird population: A reciprocal link between ecology and evolution. *Proceedings of the Royal Society B: Biological Sciences*, 283(1829), 20152411.

Scherer, U., Godin, J.-G.-J., & Schuett, W. (2020). Do female rainbow kribbs choose males on the basis of their apparent aggression and boldness? A non-correlational mate choice study. *Behavioral Ecology and Sociobiology*, 74(3), 1–15. https://doi.org/10.1007/s00265-019-2794-5

Schirmer, A., Herde, A., Eccard, J. A., & Dammhahn, M. (2019). Individuals in space: Personality-dependent space use, movement and microhabitat use facilitate individual spatial niche specialization. *Oecologia*, https://doi.org/10.1007/s00442-019-04365-5

Schirmer, A., Hoffmann, J., Eccard, J. A., & Dammhahn, M. (2020). My niche: Individual spatial niche specialization affects within- and between-species interactions. *Proceedings of the Royal Society B: Biological Sciences*, 287(1918), 20192211. https://doi.org/10.1098/rspb.2019.2211

Schuett, W., Tregenza, T., & Dall, S. R. X. (2010). Sexual selection and animal personality. *Biological Reviews*, 85(2), 217–246. https://doi.org/10.1111/j.1469-185X.2009.00101.x

Sih, A., Bell, A., & Johnson, J. C. (2004). Behavioral syndromes: An ecological and evolutionary overview. *Trends in Ecology & Evolution*, 19(7), 372–378. https://doi.org/10.1016/j.tree.2004.04.009

Sih, A., Chang, A. T., & Wey, T. W. (2014). Effects of behavioural type, social skill and the social environment on male mating success in water striders. *Animal Behaviour*, 94, 9–17. https://doi.org/10.1016/j.anbehav.2014.05.010

Sih, A., & Del Giudice, M. (2012). Linking behavioural syndromes and cognition: A behavioural ecology perspective. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 367(1603), 2762–2772.

Smith, B. R., & Blumstein, D. T. (2008). Fitness consequences of personality: A meta-analysis. *Behavioral Ecology*, 19(2), 448–455. https://doi.org/10.1093/beheco/arm144

Solomon, N. G., & Jacquot, J. J. (2011). Characteristics of resident and wandering prairie voles, *Microtus ochrogaster*. *Canadian Journal of Zoology*, https://doi.org/10.1139/z10-202-053

Spiegel, O., Leu, S. T., Bull, C. M., & Sih, A. (2017). What’s your move? Movement as a link between personality and spatial dynamics in animal populations. *Ecology Letters*, 20(1), 3–18. https://doi.org/10.1111/ele.12708

Spritzer, M. D., Solomon, N. G., & Meikle, D. B. (2006). Social Dominance among male meadow voles is inversely related to reproductive success. *Ethology*, 112(10), 1027–1037. https://doi.org/10.1111/j.1439-0310.2006.01260.x

Stoffel, M. A., Nakagawa, S., & Schielzeth, H. (2017). rptR: repeatability estimation and variance decomposition by generalized linear mixed-effects models. *Methods in Ecology and Evolution*, 8, 1639–1644.

Terborgh, J., & Janson, C. H. (1986). The sociocology of primate groups. *Annual Review of Ecology and Systematics*, 17, 111–136. https://doi.org/10.1146/annurev.es.17.110186.000551

Tkadlec, E., & Krejčová, P. (2001). Age-specific effect of parity on litter size in the common vole (Microtus arvalis). *Journal of Mammalogy*, 82(2), 545–550.

Veerbek, M. E. M., Drent, P. J., & Wiekpema, P. R. (1994). Consistent individual differences in early exploratory behaviour of male great tit. *Animal Behaviour*, 48, 1113–1121.

Ward-Fear, G., Brown, G. P., Pearson, D. J., West, A., Rollins, L. A., & Shine, R. (2018). The ecological and life history correlates of boldness in free-ranging lizards. *Ecosphere*, 9(3), e02125. https://doi.org/10.1002/ecs2.2125

Wolf, M., van Doorn, G. S., Leimair, O., & Weissing, F. J. (2007a). Life-history trade-offs favour the evolution of animal personalities. *Nature*, 447(7144), 581–584. https://doi.org/10.1038/nature05835

Wolf, M., van Doorn, G. S., Leimair, O., & Weissing, F. J. (2007b). Evolution of animal personalities—Reply. *Nature*, 450(7167), E5–E6. https://doi.org/10.1038/nature06327

Wolf, M., & Weissing, F. J. (2010). An explanatory framework for adaptive personality differences. *Philosophical Transactions of the Royal Society B: Biological Sciences*, 365(1560), 3959–3968. https://doi.org/10.1098/rstb.2010.0215

Worton, B. J. (1987). A review of models of home range for animal movement. *Ecological Modelling*, 38(3), 277–298. https://doi.org/10.1016/0304-3800(87)90101-3

Worton, B. J. (1995). Using Monte Carlo simulation to evaluate Kernel-based home range estimators. *The Journal of Wildlife Management*, 59(4), 794–800. https://doi.org/10.2307/3801959

Wright, J., Bolstad, G. H., Araya-Ajoy, Y. G., & Dingemanse, N. J. (2019). Life-history evolution under fluctuating density-dependent selection and the adaptive alignment of pace-of-life syndromes. *Biological Reviews*, 94(1), 230–247. https://doi.org/10.1111/brv.12451

**SUPPORTING INFORMATION**

Additional supporting information may be found in the online version of the article at the publisher’s website.

**How to cite this article:** Eccard, J. A., Herde, A., Schuster, A. C., Liesenjohann, T., Knopp, T., Heckel, G., & Dammhahn, M. (2022). Fitness, risk taking, and spatial behavior covary with boldness in experimental vole populations. *Ecology and Evolution*, 12, e8521. https://doi.org/10.1002/ece3.8521