Match or mismatch: Tokay geckos adjust their behaviour to familiar and unfamiliar handlers but according to the context

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1. **Summary**

Animals need to distinguish among different conspecific and heterospecific individuals to be able to adjust behaviour appropriately. Behavioural responses towards familiar individuals might vary based on context in which they are encountered. However, such context dependent responses, while beneficial in the wild, can impact experimental results, increase error, decrease reproducibility and threaten scientific advancement. Consequently, it is essential to understand how and when experimenters influence animal behaviour. Here, we tested captive Tokay geckos (*Gecko gecko*) ability to discriminate familiar and unfamiliar handlers across two situations (novel and routine). In the novel situation, we induced tonic immobility through physical constraint, a protocol the animals had never experienced before. In the routine situation, we fed lizards live prey with tweezers (as during regular husbandry). Geckos behaved differently towards familiar and unfamiliar handlers in a routine situation but not in a novel situation. Nevertheless, we found high individual repeatability in tonic immobility (R = 0.41-0.56). Our study, therefore, has implications for animal behaviour, cognition and welfare, while revealing important insights into context specific responses in relation to handler identity, factors that are rarely considered in experimental animal studies but that can significantly impact results.

*Keywords*: death feigning, feeding, personality, reptile, Squamata, tonic immobility
2. Introduction

To be able to behave appropriately during interactions with members of the same or different species, animals need to discriminate among different individuals (e.g. familiar versus unfamiliar, kin versus non-kin, or single individuals) [1]. Importantly, behaviour towards familiar individuals might be specific to the context in which they are encountered. For example, male rhesus monkeys' (Macaca mulatta) support in agonistic interactions depends both on the identity and relative dominance status of the receiver and the aggressor [2]. Ants (Formica xerophila and F. integroides) can behave differently towards heterospecific neighbours and strangers based on resource value. They show more aggression towards strangers within their general territory, but similar amounts of aggression towards both when near their nest [3]. Even though context dependent responses towards different individuals can be crucial in the wild, similar context specificity might however be detrimental for experimental outcomes.

Research worldwide is currently facing a reproducibility crisis, in which the findings of previous scientific studies are challenging or impossible to replicate [4]. Given that reliable, high quality results are critically important for scientific advancement, there is an urgent need to identify the root causes of this lack of reproducibility to reduce potential sources of variation. Recently, it has been shown that the subjectivity involved in data analysis can lead to vastly different results [5]. However, even if statistical analyses become more standardized, underlying issues might persist, potentially arising at any stage of a project. In studies with animals, the sampling and study design, such as where and how individuals are collected, the acclimation period to the procedures or laboratory, or the level of environmental enrichment, can impact the behaviour of animals during experiments and thus produce altered experimental results [6-8]. Importantly, researcher identity might also create behavioural differences that are not promoted by or linked to the experiment itself [9-10]. For example, unfamiliarity with the experimenter increases anxiety scores in laboratory rats [10]. Given that many animals across taxa can distinguish between human individuals [11], it is surprising that this aspect is often overlooked in experimental settings, and its impact on results should not be neglected.
Some animal species can recognize and discriminate specific human faces or human individuals. Captive fishes can recognize many different human faces displayed on a virtual screen (in archerfish, *Toxotes chatareus*) [12], and differentiate between two human caretakers that perform different husbandry tasks (in zebrafish, *Danio rerio*) [13]. Similarly, corn snakes (*Pantherophis guttata*) can distinguish between a familiar handler and a stranger, when living in enriched environments [8]. Research has also shown that some animals adjust their behaviour according to the perceived threat level associated with different individuals. For example, some bird species known for their cognitive abilities, such as jackdaws (*Corvus monedula*) [14], Antarctic brown skuas (*Stercorarius antarcticus*) [15-16], black-billed magpies (*Pica pica*) [10], Northern Mockingbirds (*Mimus polyglottos*) [17] and wild American crows, (*Corvus brachyrhynchos*) [18] can discriminate between threatening and non-threatening humans, and adjust their mobbing behaviour to directly target threatening individuals. Thus, it is likely that most captive animals can at least distinguish their caretakers from strangers [11] and that this might exert an impact during experiments [9-10]. In order to account for potential variation caused by differences in behaviour towards handlers, it is crucial to assess whether animals adjust their behaviour based on the familiarity with handlers and under which circumstances such a differentiation might occur.

In this study, we aimed to understand if captive Tokay geckos (*Gecko gecko*) would behave differently towards familiar and unfamiliar handlers depending on the context: in a novel and a routine situation. Tokay geckos are a facultative social lizard species that forms temporary family groups, showing pair-bonding and parental care [19], which requires them to be able to discriminate at least their mate and offspring among conspecifics. Indeed, they can discriminate familiar from unfamiliar mates [20] and their own odour from that of an unfamiliar same-sex conspecific [21]. Therefore, we expect them to have the sensory capacity to discriminate at least categories (familiar versus unfamiliar) of human handlers. To simulate a novel situation, we induced tonic immobility, a procedure that individuals never experienced before. Tonic immobility is induced by constraining an animal on its back and applying pressure to the spine [22], which triggers the animal to enter a state where it appears to be
dead for a period of time, after which it returns to its normal activity [23]. This anti-predator behaviour aims to distract a predator so it loses interest in the prey aiding its escape [24]. When employing tonic immobility, lizards can evaluate the threat level of the situation and adjust this strategy accordingly [25-27]. To simulate a routine situation, we presented live prey in forceps as during geckos’ usual husbandry procedure.

We hypothesised that 1) if geckos cannot discriminate between handlers, they would behave similarly towards unfamiliar and familiar researchers across situations. 2) If they can discriminate handlers and base their behaviour on previous knowledge with the handlers but ignore their experience with the situation (novel or routine), they would behave differently towards unfamiliar and familiar researchers in both situations. 3) If they can discriminate handlers and also base their behaviour on previous experience with each situation (novel or routine), they would show context-dependent behaviour and behave similarly towards unfamiliar and familiar researchers in the novel situation (mismatch between handler and context familiarity), but behave differently in the routine situation (match between handler and context familiarity).

3. Materials and Methods

Animals

In the novel situation (tonic immobility), we tested 14 adult captive bred Tokay geckos (7 males: Snout-to-vent length (SVL) range = 14.45 – 15.99 cm, 7 females: SVL range = 12.97 – 14.61 cm) [19], and in the routine situation (feeding from forceps) we tested 39 captive bred geckos (16 males: SVL range = 12.25 – 15.99 cm, 23 females: SVL range = 11.76 – 14.91 cm) including the 14 adults used in the tonic immobility test. 22 individuals were purchased from different breeders, while 17 were bred from these adult individuals in our facility. Geckos were between 2 to 7 years of age at the time of the study. Sex of individuals was determined based on the presence (male) or absence (female) of femoral glands [19].
Captive conditions

All gecko enclosures are equipped with a compressed cork wall screwed to the back, and enriched with live plants. We provide cork refuges (cork branches cut in half, hung on the back wall with hooks) as well as branches for climbing. Enclosures are set-up bioactive. They contain a drainage layer of expanded clay on the bottom, covered with mosquito mesh (to prevent mixing of the expanded clay and the soil) and topped with organic rainforest soil (Dragon BIO-Ground). Additionally, we spread autoclaved red oak leaves and sphagnum moss on top of the soil to provide shelter and food for the isopods and earth worms that break down the faecal matter produced by the geckos. Enclosures are made of rigid foam slabs with a mesh top and glass front doors.

We keep enclosures across three rooms on shelves with small enclosures on the top and large enclosures on the bottom (we tested all 11 individuals from one room and three from the second room in the novel situation and all individuals in the routine situation). The environment in the rooms is fully controlled by an automatic system that aims to mimic natural conditions. Geckos are kept under a reversed 12h:12h photo period (light: 6pm to 6am, dark: 6am to 6pm). A red light (PHILIPS TL-D 36W/15 RED) not visible to geckos [28] ensures that researchers are able to work with the geckos during the “night” when they are active. The system simulates sunrise and sunset. The day/night changes are accompanied by a change in room temperature from approximately 25 °C during the night to about 31 °C during the day. During the day, we also provide UVB (Exo Terra Reptile UVB 100, 25 W) light from directly above the enclosures. A heat mat (TropicShop) fixed to the right outside wall of each enclosure increases the temperature locally by 4-5 °C and allows lizards to thermoregulation to their optimal body temperature at any time. Base room humidity is kept at 50% but 30s of daily rainfall with reverse osmotic water approximately every 12h (at 5pm and 4am) increases the humidity within enclosures to 100% for a short period of time.

During the first three trials of the novel situation, three female geckos were kept singly in plastic terraria of the size 45 L x 45 B x 70 H cm, one male was kept singly in a terrarium of
the size 90 L x 45 B x 100 H cm and the other eight individuals were kept in pairs in terraria of the size 90 L x 45 B x 100 H cm. During the last trial, all except two individuals (G011 and G020) were housed singly (females: 45 L x 45 B x 70 H cm; males: 90 L x 45 B x 100 H cm). During the routine situation, 30 individuals were kept in pairs in terraria of the size 90 L x 45 B x 100 H cm, one male and one females were kept singly in a terrarium of the size 90 L x 45 B x 100 H cm and the remaining 7 females were kept in terraria of the size 45 L x 45 B x 70 H cm.

**Husbandry**

We feed geckos with either 3-5 adult house crickets (*Acheta domesticus*), mealworms (*Tenebrio molitor*) and/or cockroaches (*Nauphoeta cinerea*), three times per week on Monday, Wednesday and Friday individually, using 25 cm long forceps. Prior to feeding, insects are fed with cricket mix (various brands), high protein dry cat food (various brands), fresh carrots and apples to ensure that they provided optimal nutrition (Vitamin D and calcium). In gecko enclosures, water is provided *ad libitum* in a water bowl. To keep track of lizards’ health, we weigh (±1g) them once a month and measure their snout vent length (±0.5cm) approximately every three-four months.

**Tonic immobility (novel situation)**

Experimental set-up

Tonic immobility was induced inside an empty glass testing tank (45 L x 45 B x 60 H cm) with a mesh top (Exo Terra Glass tank). The testing tank was placed inside the gecko rooms on a table ensuring the same basic climatic conditions during testing as provided under normal housing. All sides, except for the doors and mesh top, were wrapped in black plastic to make them opaque. Lizards were tested under red light and a piece of cardboard was placed on the floor of the testing tank to prevent lizards from losing body heat. The testing tank was placed so that the transparent doors were facing away from the room door. Trials were recorded from
above using a Samsung S20 smartphone (108 Megapixel, 8K-FUHD) or a GoPro Hero 8 Black (linear mode, 1080 resolution, 24 FPS) placed on the mesh top of the testing tank. We ran four trials per individual for a total of 56 trials.

Procedure

First, a lizard was captured by hand from within its home enclosure by one researcher (trial 1: BS, trial 2: LB, trial 3: LB, trial 4: LB) and then handed to a second researcher who would induce tonic immobility (trial 1: IDM - unfamiliar, trial 2: BS - familiar, trial 3: ER - unfamiliar, trial 4: BS – familiar; Figure 1). All researchers involved in the study were female, experienced in the capture of geckos and with prior training on inducing tonic immobility in Tokay geckos. Next, the lizard was turned on its back (head facing to the left) within the testing tank on top of the piece of cardboard and the video recording was started. For the next 45 seconds the lizard was held on its back, left hand flat over its head and front legs, while the hind legs (thighs, Figure 1, position 4 and 5) were gently held down with two fingers of the researchers right hand. Thereafter, the experimenter changed the position of their left hand putting the pinkie finger on the lizards chin (Figure 1, position 1), and the thumb and index finger on the lizards shoulders (Figure 1, position 1 and 2). All other fingers were stretched out so as to prevent the gecko from holding on with their pads. The lizard was gently held down in this position for the remaining 75 seconds (until a total of 2 minutes had elapsed). At this point, the experimenter removed their hands, closed the testing tank doors, locked them and moved away always to the right in the direction of the lizards tail (see supplementary video S1 for the whole procedure). If the lizard did not stay on its back, the experimenter resumed induction as described above until tonic immobility was induced. Individuals were given a trial of 15 minutes to upright themselves. At the end of the trial, lizards were captured by hand and released back into their home enclosure. If a lizard had not righted itself at the end of a trial its right hind leg was gently touched to induce righting, before being transported back into its enclosure.
The researcher who induced tonic immobility washed their hands thoroughly with soap between lizards and the cardboard was either flipped or replaced each trial to avoid odour cues from other individuals influencing tonic immobility. Lizards were tested between 07:30h and 14:00h in a random order between each trial (inter-trial interval of approximately one month). We made sure not to test two lizards from the same enclosure consecutively. Half of the geckos in the study (4 males and 3 females) performed another behavioural experiment between trials 2 – 3 and 3 – 4 (scan sampling of spatial behaviour [29]; chemical mate recognition [20]). All trials were conducted between December 2022 and March 2023.

Data collection

Videos were scored using the behavioural coding software BORIS [30]. We scored the latency to induce tonic immobility in seconds, from the moment an individual was first held down using all five locations on its body until the trial start (closing of the testing tank doors). We also scored if uprighting occurred (yes = 1, no = 0) and the time taken (seconds) from trial start (closing of the testing tank doors) until an individual uprighted (duration of immobility). All latencies were scored to an accuracy of 1 second. Additionally, we scored if a tail movement occurred (yes/ no; movement of the tail in a curling manner performed as an antipredator display [31]) and which side the individual used to upright itself (left or right, side closest to the ground when turning). We used the moment the lizard had half turned around as the endpoint of the trial. If lizards did not upright within 15 minutes, they received a truncated duration of immobility of 900 seconds, occurrence of 0 and side to upright as NA. In addition, for each trial, we recorded room temperature (measured within 5 minutes of trial start), and lizards’ weight (closest measure in time to the date of the trial) and snout vent length (average across the experimental period).
Inter-observer reliability

We were unable to score videos blind to animal identity. Therefore, 50% of videos were scored by two independent observers (one trial = 25% of videos each). Scores across observers were highly consistent (Trial 1: Spearman rank correlation, $R_{\text{turning latency}} = 1$, $p_{\text{turning latency}} = 2.2\times10^{-16}$; $R_{\text{latency to induce}} = 0.96$, $p_{\text{latency to induce}} = 5.3\times10^{-8}$; Cohen’s Kappa, $k_{\text{occurance}} = 1$, $N_{\text{occurance}} = 14$; $k_{\text{side}} = 1$, $N_{\text{side}} = 10$; $k_{\text{tail}} = 1$, $N_{\text{tail}} = 14$; Trial 4: Spearman rank correlation, $R_{\text{turning latency}} = 0.99$, $p_{\text{turning latency}} = 2.2\times10^{-16}$; $R_{\text{latency to induce}} = 0.99$, $p_{\text{latency to induce}} = 4\times10^{-13}$; Cohens Kappa, $k_{\text{occurance}} = 1$, $N_{\text{occurance}} = 14$; $k_{\text{side}} = 1$, $N_{\text{side}} = 8$; $k_{\text{tail}} = 0.87$, $N_{\text{tail}} = 8$).

Feeding from forceps (routine situation)

Experimental set-up

Lizards were tested within their home enclosure on two feeding days (21st and 23rd of January 2024, between 9:00 and 11:00 am). Beforehand, we randomly split lizards into two groups, one was first tested by the familiar handler, while the other half was first tested by the unfamiliar handler. In this experiment the unfamiliar handler was male, while the familiar handler was female. The order of testing was reversed on the following test day. Furthermore, within a day, lizards were tested in a random order. The unfamiliar handler received prior training (one day) on how to feed and perform video recordings of gecko behaviour.

Procedure

At the start of the test, a dim white light (LED, SPYLUX® LEDVANCE 3000 K, 0.3 W, 17 lm), that lizards were accustomed to (used during regular husbandry), was placed on top of the tank. Next, a focal lizard was located within its enclosure. If necessary, cork shelters were gently removed to be able to take video recordings of their behaviour. Once the focal individual was visible, a video recording was started using a Samsung S20 smartphone (108 Megapixel, 8K-FUHD). Then, a live cockroach was presented to the individual within 4-5 cm in front of its snout using 25 cm long forceps (Figure 1). The behaviour of the lizard was recorded either
until an attack occurred, it walked away or did not respond for 30s (this time was deemed appropriate as lizards usually attack prey immediately). Each handler was alone in the room while performing the experiment.

Data collection

Videos were scored using the behavioural coding software BORIS [30]. We measured the time from when the lizard first noticed a food item until the first attack regardless of if the food was captured or not (latency) as well as its' occurrence (yes = 1 and no = 0). We assumed that a food item was first noticed when a lizard moved its' head or eyes to focus on the prey [32]. Lizards that did not attack the prey within 30s were given a latency of 18.5 sec (longest latency + 1 s) instead of excluding them from the analysis. The handler was not visible in the videos and previously, inter-observer reliability was high when measuring latency to attack prey (r > 0.9 [32], unpublished data). Therefore, we did not perform an inter-observer reliability test for this experiment.

Statistical analyses

Tonic immobility (novel situation)

First, we investigated if the probability of uprighting (Bernoulli variable, turn = 1, no turn = 0) was influenced by the fixed effects of sex (male, female), the latency to induce tonic immobility, if tail movement occurred (yes = 1, no = 0), handler familiarity (familiar - BS, unfamiliar - IDM & ER), room temperature (degree celcius) and the body condition of the lizard (scaled mass index [33]). Originally, we also included the interaction between handler familiarity and the latency to induce tonic immobility in the model but as this was not significant the interaction was removed to ensure better model performance. We used a Bayesian generalised linear mixed model (GLMM) with a Bernoulli distribution from the package brms [34-36] with random effects of animal identify (intercept) and trial (1-4, slope).
Second, we investigated if the duration of immobility (log-normal variable) was influenced by the fixed effects handler familiarity (familiar - BS, unfamiliar - IDM & ER), sex (male, female), the latency to induce tonic immobility, if tail movement occurred (yes = 1, no = 0), room temperature and the body condition of the lizard. In this model, we also removed the interaction between handler familiarity and the latency to induce tonic immobility to ensure better model performance. Because the duration of immobility measures were censored (cut off at 900 seconds), we used a censored Bayesian GLMM with a log-normal distribution and random effects of animal identify (intercept) and trial (slope).

Third, we investigated if the probability of uprighting to the right (Bernoulli variable, right turn = 1, left turn = 0) was influenced by the fixed effects of handler familiarity (familiar, unfamiliar), sex (male, female), room temperature and the body condition of the lizard. Again, we used a Bayesian GLMM with a Bernoulli distribution and random effects of animal identify (intercept) and trial (slope).

Finally, we investigated agreement repeatability in the duration of immobility using the package rptR [37]. We log-transformed the duration of immobility to fit a normal distribution. We calculated individual repeatability from the whole dataset and also after removal of trials in which a lizard did not upright (with a censored latency of 900 seconds) as we wanted to know if the truncated trials would bias repeatability. Due to the small sample size we did not calculate individual repeatability in the probability of uprighting and the side to upright.

Feeding from forceps (routine situation)

First, we investigated if the probability of attacking prey (Bernoulli variable, eaten = 1, not eaten = 0) was influenced by the fixed effects of sex (male, female), handler familiarity (familiar - BS, unfamiliar - PG), repetition (Wednesday = 1, Friday = 2), room temperature and the body condition of the lizard. We included the interaction between handler familiarity and sex, which was further analysed using post hoc least squares means tests (LSM, package emmeans [38]). We used a Bayesian GLMM with a Bernoulli distribution and a random effect of animal identify (intercept).
Second, we investigated if the latency to attack (log-normal variable) was influenced by the fixed effects handler familiarity (familiar, unfamiliar), sex (male, female), room temperature and the body condition of the lizard. Again, we included the interaction between handler familiarity and sex which was further analysed using post hoc least squares means tests. We used a Bayesian GLMM with a log-normal distribution and a random effect of animal identify (intercept).

All analyses were run in R version 4.2.2 [39]. For all Bayesian models, we ensured that Rhat was 1, that the ESS was above 2000 and checked the density plots and correlation plots to ensure that the models had sampled appropriately. We used a diffuse normal prior with a mean of 0 and a standard deviation of 1. We used a test for practical equivalence to determine whether to accept or reject a "null hypothesis", formulated as "no difference" or "no relationship", for each fixed effect in a model using the equivalence_test function from the package bayestestR [40]. We report results in which the null hypothesis was accepted (100% within the Region of Practical Equivalence – ROPE) or was undecided as no evidence and results in which the null hypothesis was rejected (0% within the ROPE) as evidence. Additionally, we provide Bayes factors (BF) to further evaluate the results by determining Bayes Factors from marginal likelihoods using the package brms [34-36] or Bayes Factor pairwise comparisons from the package pairwiseComparisons [41] where appropriate. Bayes factors below 1 indicate more support for no difference while above 1 more support for a difference [42]. We report cases in which the equivalence test produced “undecided” results but Bayes factors were above 1 as evidence.

4. Results

Tonic immobility (novel situation)

We were able to induce tonic immobility to all geckos, across all 56 trials. We found no evidence for the probability of uprighting to differ between familiar and unfamiliar handlers.
(GLMM, estimate_{unfamiliar} = 0.796, 95% CI_{low} = -0.790, 95% CI_{up} = 2.431, 12.19% inside ROPE, BF = 0.701). Moreover, we found no evidence that the probability of uprighting was associated with temperature (GLMM, estimate = -0.469, 95% CI_{low} = -1.654, 95% CI_{up} = 0.645, 19.05% inside ROPE, BF = 0.825), body condition (GLMM, estimate = -0.007, 95% CI_{low} = -0.184, 95% CI_{up} = 0.167, 99.91% inside ROPE, BF = 0.090), sex (GLMM, estimate_{male} = -0.076, 95% CI_{low} = -1.854, 95% CI_{up} = 1.701, 16.27% inside ROPE, BF = 0.251), if tail movement occurred (GLMM, estimate_{yes} = -0.422, 95% CI_{low} = -1.990, 95% CI_{up} = 1.207, 16.06% inside ROPE, BF = 0.926) or with the latency to induce tonic immobility (GLMM, estimate = -0.023, 95% CI_{low} = -0.055, 95% CI_{up} = 0.003, 100% inside ROPE, BF = 0.061).

Similarly, we found no evidence that the duration of immobility differed between familiar and unfamiliar handlers (GLMM, estimate_{unfamiliar} = -0.627, 95% CI_{low} = -1.655, 95% CI_{up} = 0.376, 0.89% inside ROPE, BF = 0.349; Figure 2A). Furthermore, we found no evidence that the duration of immobility was associated with temperature (GLMM, estimate = -0.198, 95% CI_{low} = -0.916, 95% CI_{up} = 0.418, 2.51% inside ROPE, BF = 0.320), body condition (GLMM, estimate = 0.028, 95% CI_{low} = -0.045, 95% CI_{up} = 0.105, 18.03% inside ROPE, BF = 0.050), sex (GLMM, estimate_{male} = 0.522, 95% CI_{low} = -0.945, 95% CI_{up} = 1.931, 0.96% inside ROPE, BF = 0.232), or the latency to induce tonic immobility (GLMM, estimate = 0.008, 95% CI_{low} = -0.001, 95% CI_{up} = 0.018, 67.24% inside ROPE, BF = 0.020). However, we found evidence that the probability that tail movement occurred was higher when individuals took longer to uprighten themselves (GLMM, estimate_{yes} = 0.675, 95% CI_{low} = -0.192, 95% CI_{up} = 1.533, 0.58% inside ROPE, BF = 1.429).

We found evidence for individual agreement repeatability of the duration of immobility of R = 0.414 (CI_{low} = 0.15, 95% CI_{up} = 0.74; Figure 2B). Similarly, after removal of trials in which lizards did not uprighten, we still found evidence for individual agreement repeatability in the duration of immobility of R = 0.555 (CI_{low} = 0.086, 95% CI_{up} = 0.815).

We found no evidence that the probability of uprighting to the right side was associated with temperature (GLMM, estimate = 0.043, 95% CI_{low} = -0.860, 95% CI_{up} = 0.945, 32.65% inside ROPE, BF = 0.461), or body condition (GLMM, estimate = 0.031, 95% CI_{low} = -0.061,
95% CI up = 0.136, 100% inside ROPE, BF = 0.056), nor did it differ between males and females (GLMM, estimate<sup>male</sup> = -0.408, 95% CI<sub>low</sub> = -1.791, 95% CI<sub>up</sub> = 0.986, 18.04% inside ROPE, BF = 0.437) or familiar and unfamiliar handlers (GLMM, estimate<sup>unfamiliar</sup> = -0.586, 95% CI<sub>low</sub> = -2.061, 95% CI<sub>up</sub> = 0.925, 14.12% inside ROPE, BF = 0.312). Some of the lizards showed a side bias when uprighting (Table 1).

Feeding from forceps (routine situation)

We found an almost 13 times stronger support for a difference in the probability to attack across males and females (GLMM, estimate<sup>male</sup> = 1.410, 95% CI<sub>low</sub> = 0.070, 95% CI<sub>up</sub> = 2.789, 1.08% inside ROPE, BF = 12.958) as well as a 20 times stronger support for a difference in the probability to attack prey presented by a familiar versus an unfamiliar handler (GLMM, estimate<sup>unfamiliar</sup> = -1.576, 95% CI<sub>low</sub> = -2.719, 95% CI<sub>up</sub> = -0.475; 0% inside ROPE, BF = 20.304). Post hoc tests revealed, that females but not males were less likely to attack a prey when presented by an unfamiliar handler (LSM, females: estimate<sub>familiar-unfamiliar</sub> = 1.560, 95% CI<sub>low</sub> = 0.45, 95% CI<sub>up</sub> = 2.740, 0% inside ROPE; males: estimate<sub>familiar-unfamiliar</sub> = 1.160, 95% CI<sub>low</sub> = -0.399, 95% CI<sub>up</sub> = 2.820, 3.86% inside ROPE; Figure 3a). Furthermore, males were more likely to attack prey than females, when the handler was unfamiliar (LSM, unfamiliar handler: estimate<sub>familiar-male</sub> = -1.420, 95% CI<sub>low</sub> = -2.760, 95% CI<sub>up</sub> = -0.060 , 0.26% inside ROPE; unfamiliar handler: estimate<sub>familiar-male</sub> = -1.820, 95% CI<sub>low</sub> = -3.320, 95% CI<sub>up</sub> = -0.308, 0% inside ROPE; Figure 3a). We found no evidence that the order of testing (familiar or unfamiliar handler first; GLMM, estimate = -0.109, 95% CI<sub>low</sub> = -1.205, 95% CI<sub>up</sub> = 0.956, 26.64% inside ROPE, BF = 0.541) or body condition (GLMM, estimate = 0.040, 95% CI<sub>low</sub> = -0.012, 95% CI<sub>up</sub> = 0.096, 100% inside ROPE, BF = 0.075) were related to the probability to attack prey. We found evidence that geckos were more likely to attack prey at lower enclosure temperature (GLMM, estimate = -0.862, 95% CI<sub>low</sub> = -2.062, 95% CI<sub>up</sub> = 0.302, 9.09% inside ROPE, BF = 1.689).

Similarly, we found almost nine times as much support for a difference in the latency to attack when prey was presented by a familiar or unfamiliar handler (GLMM, estimate<sup>unfamiliar</sup>
= 0.852, 95% CI_{low} = 0.435, 95% CI_{up} = 1.263, 0% inside ROPE, BF = 8.993) and over six
times as much support for a difference across the sexes (GLMM, estimate_{male} = -0.894, 95% CI_{low} = -1.591, 95% CI_{up} = -0.193, 0% inside ROPE, BF = 6.370). Both males and females took
longer to attack prey presented by an unfamiliar handler compared to a familiar handler (LSM, females: estimate_{familiar-unfamiliar} = -0.854, 95% CI_{low} = -1.280, 95% CI_{up} = -0.454, 0% inside ROPE, BF = 5.479; males: estimate_{familiar-unfamiliar} = -0.736, 95% CI_{low} = -1.210, 95% CI_{up} = -0.228, 0% inside ROPE, BF = 1.461; Figure 3b). Furthermore, regardless of handler familiarity, males attacked prey faster than females, but this difference was larger when prey was presented by the unfamiliar handler (LSM, familiar handler: estimate_{female-male} = 0.886, 95% CI_{low} = -0.200, 95% CI_{up} = 1.570, 0% inside ROPE, BF = 2.185; unfamiliar handler: estimate_{female-male} = 1.001, 95% CI_{low} = 0.268, 95% CI_{up} = 1.700, 0% inside ROPE, BF = 7.086; Figure 3b). We found no evidence that the order of testing (familiar or unfamiliar handler first; GLMM, estimate =0.039, 95% CI_{low} = -0.330, 95% CI_{up} = 0.415, 4.60% inside ROPE, BF = 0.188), temperature (GLMM, estimate = 0.215, 95% CI_{low} = -0.350, 95% CI_{up} = 0.792, 2.35% inside ROPE, BF = 0.380) or body condition (GLMM, estimate = -0.015, 95% CI_{low} = -0.039, 95% CI_{up} = 0.009, 32.26% inside ROPE, BF = 0.026) were related to the latency to attack prey.

5. Discussion

We found that Tokay geckos can discriminate between familiar and unfamiliar human individuals but show context-dependent behavioural responses. In the tonic immobility experiment, during which geckos experienced a novel, stressful situation, they did not exhibit behavioural differences when tested by a familiar or an unfamiliar handler. Instead, individuals behaved consistently in their duration of immobility across four trials with an inter-trial interval of one month. Contrary, in the feeding experiment, a routine familiar situation that did not involve direct handling, their behaviour differed when tested by a familiar compared to an unfamiliar experimenter, exhibiting overall more caution with the unfamiliar handler.
Our results support our third hypothesis showing that lizards can discriminate between human handlers but take the context into account when deciding how to respond. Geckos performed similarly in the novel situation, but adjusted their behaviour to familiar and unfamiliar handlers in the routine situation. Similar to the results from a study modelling decision making based on risk [43], our results show that geckos rely more strongly on past experiences (i.e. the familiarity with the handler) when the information regarding the risk level was more predictable (in the routine feeding situation). Alternatively, it is possible that when the threat level is high, as in the novel situation, geckos still discriminate between handlers, but even familiar humans may be perceived as threatening when the outcome is uncertain. A number of studies focusing on domesticated animals show that the sole presence of humans can act as a social buffer in stressful situations, modulating the animals’ stress levels (e.g. in dogs [44-47] and goats [47]). Yet, even though our geckos are captive bred and have extensive experience with humans, geckos behaved more similarly to wild than domesticated animals.

Remarkably, this is one of the very few studies demonstrating that reptiles can discriminate individuals of a different species and adjust their behaviour according to context (alongside with [8] in corn snakes). While this finding is exciting in itself and enhances our understanding of reptilian behaviour and cognitive abilities (related to context dependent decision-making), it also raises implications for data quality and research reproducibility. Our results show that (1) the identity of the researcher does introduce error into the data which needs to be accounted for, and (2) that the effect might vary from protocol to protocol. It is also possible, that animals do not just discriminate familiar from unfamiliar humans but might show more nuanced discrimination with certain researchers introducing even more error into the data leading to increased bias complexity. To better understand the discrimination ability of these animals, future studies could also manipulate experimenter roles (bad vs good [19]). In any case, it highlights that a first step to mitigate the reproducibility crisis in experimental studies could be to consider the effect of handler identity in animal experiments, as this might impact the animals’ behaviour in complex ways [9-10].
As of yet, we have no information regarding which cues lizards use to make the discrimination between human handlers. Geckos rely heavily on chemicals for social communication [47-48], but they also have a well-developed visual system [49]. Therefore, any of these cues or combinations might be used. It is also possible that, the more information across different modalities is available at a given moment, the better their ability to discriminate and this should be tested in the future. Moreover, even though our results point towards a context specific use of past experiences, the difference in gecko behaviour across experiments could be, at least in part, related to handler sex. For example, a study in mice demonstrated that individuals avoided the scent of a male experimenter and showed increased stress susceptibility after the administration of an antidepressant [50]. However, Tokay geckos habituate to new human handlers across time regardless of their sex (unpublished data). Therefore, it seems unlikely that handler sex rather than context led to our result.

It is also worth noting that we found high intra-individual consistency in the duration of immobility across time regardless of who performed the protocol. Consistent tonic immobility behaviour across trials was found in birds (Yellow-crowned bishop, *Euplectes afer*; Tree sparrow, *Passer montanus* [51]), amphibians (smooth newt, *Lissotriton vulgaris* [52]), and insects (yellow mealworm beetle, *Tenebrio molitor* [53]), and here we add evidence in a gecko. Moreover, we found a repeatability of 0.41 (and 0.55 after removing trials where lizards did not upright) for the duration of immobility, which is higher than average in studies on animal behaviour (average = 0.37) [54]. This is quite remarkable, as these lizards never lived in the wild (and thus never encountered a natural predator), were habituated to humans, and underwent repeated trials with intervals of one month. This individual repeatability can be interpreted as a personality trait [55] potentially measuring boldness or antipredator behaviour [51], and due to its consistency in the current experiment, might have a genetic basis rather than being based on experience [56]. Finally, we found that room temperature was related to behavioural responses. Lizards are ectotherms and temperature has an impact on behaviour, especially feeding behaviour as temperature is directly related to digestion [58-60]. This
highlights temperature should always be either controlled or accounted for (as a covariate similarly to our analysis) in behavioural research in ectotherms.

6. Conclusion

Our findings demonstrate context dependent behavioural responses in Tokay geckos in which individuals behave according to a match or mismatch between handler and context familiarity. When the context was a novel, unfamiliar test, geckos behaved similarly when handled by familiar and unfamiliar researchers; when the context was a routine, familiar test, geckos behaved differently when tested by familiar or unfamiliar handlers, showing more cautious behaviour with the latter. Hence, geckos are able to discriminate among heterospecifics such as different human individuals, but they act upon it depending on the context. Accounting for the effect of handler identity in experiments can thus be crucial for refining study design and mitigating potential sources of measurement error, which can have implications for data quality and contribute to the global reproducibility crisis in research. Overall, our study bears implications for experimental practices, while further contributing to our understanding of Tokay gecko behaviour and cognition.

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Ethical statement

Our tests followed the guidelines provided by the Association for the Study of Animal Behaviour/ Animal Behaviour Society for the treatment of animals in behavioural research and Teaching (ASAB Ethical Committee and ABS Animal Care Committee, 2022) and the
Guidelines for the ethical use of animals in applied animal behaviour research by the International Society for Applied Ethology (Sherwin et al, 2003). We also complied with the ARRIVE guidelines. Experiments were approved by the Suisse Federal Food Safety and Veterinary Office (National No. 33232, Cantonal No. BE144/2020). Captive conditions were approved by the Suisse Federal Food Safety and Veterinary Office (Laboratory animal husbandry license: No. BE4/2022). All lizards were part of our permanent captive stock and were retained in our facility after the experiment.

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Data Accessibility

Our code and dataset can be accessed at OSF under the link: https://osf.io/zeshv/?view_only=33667cfa1a774ea9b5391a7301f58536).

Declaration of AI use

AI-assisted technologies were used to aid in the creating of the abstract.

Competing Interests

We have no competing interests.

Authors' Contributions

IDM, BS - Conceptualization; IDM, BS - Data curation; BS - Formal analysis; BS - Funding acquisition; IDM, ER, LB, BS - Investigation; IDM, BS - Methodology; BS - Project
administration; ER, BS - Resources; BS - Validation; IDM, BS - Visualization; IDM, ER, LB, BS - Roles/Writing - original draft; IDM, ER, LB, BS - Writing - review & editing.
References


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### Tables

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### Figures

**Figure 1**
RUNNING HEAD: Context dependent responses to handlers in geckos

Figure 2

![Graph showing context-dependent responses to handlers in geckos.](image)

**Average duration of immobility (sec)**

- **Handler familiarity**: familiar vs. unfamiliar
- Box plots for each condition

![Duration of immobility (sec)](image)

**Duration of immobility (sec)**

- **Individual (ordered by mean latency value)**
- Scatter plot with confidence intervals

- **R = 0.41 [0.15-0.74]**

---

Figure 3

![Graph showing context-dependent responses to handlers in geckos.](image)

**Percentage of individuals that ate the prey**

- **Percentage**
- **Unfamiliar vs. familiar**

- **Latency to attack prey (sec)**
- **Unfamiliar vs. familiar**
Figure and table captions

Table 1. Laterality index (LI) calculated as \( \frac{N_{\text{right}}}{N_{\text{left}} + N_{\text{right}}} \) for each individual. The table presents individuals’ identity (ID), sex, the number of trials the geckos uprighted to the right \( (N_{\text{right}}) \) or to the left side \( (N_{\text{left}}) \), the total number of trials in geckos uprighted \( (N_{\text{total}}) \), and the laterality index to upright to the right side \( (LI_{\text{right}}) \). A "-" indicates no uprighting occurred.

Figure 1. Experimental procedures. Tonic immobility (novel situation): White circles on the ventral area of the individual indicate the locations at which the lizard was held by the handler during the induction of tonic immobility in all trials. Each gecko was tested four times, one month apart, by unfamiliar (trial 1 and 3, by two different researchers) and familiar handlers (trial 2 and 4, same researcher). Each lizard was allowed 15 minutes to upright. Feeding from forceps (routine situation): Geckos were fed with forceps by an unfamiliar and a familiar handler (twice by each in different days), and given 30 seconds to complete the trial.

Figure 2. Tonic immobility (novel situation) - Duration of immobility after handling by familiar and unfamiliar researchers, and individual gecko consistency in this behaviour. a) Boxplots of the duration of immobility (grey points represent the individual average per treatment) between trials in which a familiar (BS) and an unfamiliar handler (IDM and ER) induced tonic immobility. The bold line shows the median, the upper and lower edge of the boxes shows the upper and lower quartile, respectively, and the top and bottom edge of the whisker shows the maximum and minimum, respectively. b) Individual behavioural consistency over the four trials (ordered by mean latency for visual purposes). Open circles represent raw data from each trial, closed circles represent individual mean, and black vertical lines show individual variation. For both a) and b) we tested 7 females and 7 males.
**Figure 3.** Feeding from forceps (routine situation) - Feeding behaviour towards prey presented by unfamiliar and familiar handlers. a) Percentage of individuals that attacked the prey presented by the unfamiliar (PG) and familiar handler (BS). Individuals that attacked and ate the prey are represented in solid lines, and those that did not attack are represented in dashed lines. b) Boxplots of the latency to attack the prey (grey points represent the individual data points) between trials in which a familiar (BS) and an unfamiliar handler (PG) presented a prey. Females are represented in light grey and males in dark grey. The bold line shows the median, the upper and lower edge of the boxes shows the upper and lower quartile, respectively, and the top and bottom edge of the whisker shows the maximum and minimum, respectively. The asterisk represents significant difference (confidence interval not crossing 0). For both a) and b) we tested 23 females and 16 males.