# Individual recognition in a jumping spider (*Phidippus regius*)

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#### <sup>13</sup> Summary

Individual recognition is conceptually complex and computationally intense, leading to 14 the general assumption that this social knowledge is solely present in vertebrates with 15 larger brains, while miniature-brained animals in differentiating societies eschew the 16 evolutionary pressure for individual recognition by evolving computationally less demanding 17 class-level recognition, such as kin, social rank, or mate recognition. Arguably, this social 18 knowledge is restricted to species with a degree of sociality (sensu [1], for a review [2]). 19 Here we show the exception to this rule in an asocial arthropod species, the jumping spider 20 (*Phidippus regius*). Using a habituation - dishabituation paradigm, we visually confronted 21 pairs of spatially separated spiders with each other and measured the 'interest' of one 22 spider towards the other. The spiders exhibited high interest upon initial encounter 23 of an individual, reflected in mutual approach behaviour, but adapted towards that 24 individual when it reoccurred in the subsequent trial, indicated by their preference of 25 staying farther apart. In contrast, spiders exhibited a rebound from habituation, reflected 26 in mutual approach behaviour, when a different individual occurred in the subsequent 27 trial, indicating the ability to tell apart spiders' identities. These results suggest that P. 28 regius is capable of individual recognition based on long-term social memory. 29

## **30 Keywords**

Recognition memory, long-term social memory, individual recognition, flexible learning,
 arthropod, invertebrate, salticid, jumping spider, *Phidippus regius*

#### <sup>33</sup> Main text

Recognising individuals is a complex cognitive process requiring flexible learning and 34 recognition memory. Arthropod species possessing the social ability of individual recognition 35 would, thus, stand in stark contrast to the commonly accepted notion that animals 36 with smaller brains are cognitively less advanced due to reduced computational power 37 of nervous systems with smaller and fewer neurons [3]. And yet, there is evidence 38 for an arthropod species displaying face learning [4] and long-term social memory [5]. 39 That is, a social wasp species (*Polistes fuscatus*) showed mammal-like face learning 40 [4, 6], arguably providing social benefits by reducing aggression and stabilizing social 41 interactions. With this, being one of the few reported cases of individual recognition in 42 arthropods (also see [7]), it is considered unlikely that asocial arthropod species would 43 evolve such complex cognitive processes. The reasons being high energy consumption, 44 long processing times and, thus, increased predation risk that would never be outweighed 45 by the few social encounters between individuals and the additional survival benefit 46 [2, 8]. The general consensus, thus, is that a certain degree of sociality sensu Wilson 47 [1] is required for the emergence of individual recognition [8]. Here, we challenge this 48 consensus: In a naturalistic experimental procedure, we put to the test the ability of 49 individual recognition in a notoriously associal and miniature-brained arthropod species, 50 a member of the *Salticidae* family, the jumping spider (*Phidippus regius*). 51

In a first step, we assess the ability of *P. regius* to individually recognise other members 52 of its species, commonly referred to as individual recognition [9] or individuation of 53 conspecifics [10]. For this purpose, we used a habituation - dishabituation procedure, 54 where, in general terms, one individual habituates to the presence of another individual 55 in its close proximity and dishabituates when, after a short phase of visual separation, 56 another individual is present in close proximity, assuming that the one individual is 57 capable of discriminating the identities of the two individuals it was confronted with 58 [11, 12]. In other terms, with this habituation - dishabituation paradigm we expect to 59 see that the rebound in 'interest' following changes in a spider's identity is greater than 60 the rebound in 'interest' following a repetition of identity. To experimentally control 61 the animal pairs, we placed the individuals in separate containers with one side and the 62 top panel being transparent. We then pairwise confronted the individuals by placing 63 the containers such that the transparent sides faced each other in the following fashion: 64

Individual A and individual B were exposed to each other for 7 minutes, triggering an 65 initial 'interest' in each other, and were then visually separated by means of an opaque 66 slider for 3 minutes. Subsequently, they were either exposed to the same individual again 67 (A vs B, habituation trial) for 7 minutes, or to a different individual (A vs C, or B vs 68 D, dishabituation trial) for 7 minutes, followed by a 3-minute period of visual separation. 69 The relative interest is quantified by approximating spatial distances between the spider 70 pairs in the xy-plane, where high interest is reflected in smaller values (i.e. spiders 71 go close) and low interest in larger values (spiders stay apart). Therefore, under the 72 assumption that the spiders are capable of individuating each other, we predict that 73 in the *habituation* condition, involving the same individuals, the relative interest in each 74 other decreases and, hence, spiders' distances increase (Figure 1a-c; 'Habituation', dashed 75 line: towards maximal distance; solid line: medium distance), while in the dishabituation 76 condition, involving a different individual, the relative interest in each other increases and 77 spiders approach each other, hence, distances decrease (Figure 1a-c; 'Dishabituation'). 78 We divided a total of 20 individuals into five groups of four individuals each. Each 79 individual of each group was exposed to the three group members in both habituation and 80 dishabituation trials, resulting in six trials per session, equivalent to one hour of recording. 81 We repeated this procedure twice, resulting in 18 trials across three sessions and an exact 82 repetition of a given trial (and pairing of individuals) in 1-hour intervals (for a detailed 83 description of the procedure see Materials and Methods and Tables 1-2). We found that 84 habituation and dishabituation trials (i.e. predictor variable condition) were significantly 85 dissociated as a function of inter-individual distances (i.e. predictor variable *distance*), 86 leading to a significant improvement of model fitting the interaction of the predictors 87 distance and condition (LRT:  $\chi^2_{\Delta 3}$  = 63.66, p < 0.001; Figure 2a, Supplementary Table 88 1): dishabituation trials (blue discs) showed a greater proportion of close-distance values 89 than *habituation* trials (red discs), whereas *habituation* trials showed a greater proportion 90 of far-distance values. The interaction between the predictors distance and condition 91 further significantly interacted with the predictor *session*, modulating the level of the 92 dissociative effect of *condition* over the progression of the testing period, showing the 93 strongest modulation in session 1 and the weakest modulation in session 3 (LRT:  $\chi^2_{\Delta 6}$  = 94 34.14, p < 0.001; Figure 2a, Supplementary Table 1, exemplar trial: Figure 1 d-f). The 95 systematic dissociation of distance values between *habituation* and *dishabituation* trials 96

<sup>97</sup> suggests that *P. regius* possesses the ability to individuate conspecifics.

The question arises whether *P. regius*'s decreasing interest over session repetitions is 98 caused by a general fatigue effect due the prolonged testing procedure or whether, in later 99 testing sessions, *P. requis* actually recognises the current individual after having seen it 100 before at least once (when encountering it again in session 2) or twice (when encountering 101 it again in session 3) and, thus, would not dishabituate any longer. Such recognition 102 capability would further emphasise the role of long-term memory representations in the 103 individuation of conspecifics, due to the prolonged retention interval beyond the minute 104 range into the hours. In a second step, we therefore assessed the extent to which a 105 presentation of an individual novel and unseen across the three experimental sessions 106 would trigger a rebound in interest at the end of session 3, henceforth referred to as 107 dishabituation [long-term] trials, as opposed to the dishabituation trials of sessions 1-3, 108 henceforth referred to as *dishabituation* [short-term] trials (see Table 3). If such rebound 109 occurs, we conclude that the habituation across sessions is the result of recognition of 110 repeatedly presented individuals and not the result of a fatigue effect due to prolonged 111 testing procedures. In other words, such rebound would suggest a 'cognitive' fatigue 112 towards seeing the same 'old' individuals, subserved by long-term memory formation, 113 rather than a 'physical' fatigue effect. To this end, we re-ran the experiment in an 114 additional 16 spiders, arranged to four groups and added a memory dishabituation [long-term] 115 trial at the end of session 3. The memory dishabituation [long-term] trials were generated 116 by cross-combining individuals from two groups (group 1: A, B, C, D; group 2: E, F, 117 G, H; Table 3), which were run in parallel, at the end of session 3, resulting in novel 118 pairings (A - E, B - G, C - F, D - H). We replicated our previous results and found a 119 dissociation of the factors distance and condition (LRT:  $\chi^2_{\Delta 3} = 29.52$ , p < 0.001; Figure 120 2b, Supplementary Table 2), showing a greater proportion of close-distance values for 121 dishabituation (blue discs) than for habituation trials (red discs) and a greater proportion 122 of far-distance values for habituation than for dishabituation trials. Critically, we also 123 found that the dishabituation [long-term] trials at the end of session 3 elicited a rebound 124 in interest that exceeded the rebound in the *dishabituation* [short-term] trials of session 125 3 by far, reflected in the interaction of *condition* (i.e. *dishabituation* [short-term] vs 126 dishabituation [long-term]) and distance (F(3,127) = 3.91, sum sq. = 0.92, mean sq.127 = 0.31, p < 0.01, Figure 2b (right subfigure, white diamonds), exemplar trials: Figure 128

1g-i, j-l). Thus, the habituation across sessions reflects a decrease in interest for the same
repeatedly presented individuals on the basis of long-term memory capabilities.

Our findings show, firstly, that *P. regius* recognised individuals to which it was exposed 131 to for a short period of 7 minutes and that reoccurred after a visual separation period 132 of 3 minutes. Secondly, *P. regius* habituated in the long-term, i.e. 1 hour and 2 hours 133 after initial presentation of a given individual. Thirdly, despite long-term habituation, P. 134 regius showed an unprecedented rebound in interest towards an entirely novel individual, 135 ruling out a physical fatigue effect in favour of a cognitive fatigue on the basis of long-term 136 social memory capabilities. For these reasons, our results are the first to suggest that 137 P. regius, an asocial arthropod species, possesses long-term memory, which allows it to 138 individuate conspecifics and recognise novel individuals. 139

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Recognising members of one's own species is crucial for survival and a requirement for 141 various social behaviours. Individual recognition allows the receiver animal to distinguish 142 between friend and foe, to identify a mating partner, its offspring or a kin member. 143 Individual recognition is achieved via the production of individually-distinct features (e.g. 144 visual) or signals (e.g. acoustic) by the sender and extracting those features and signals 145 by the receiver [13]. Individual recognition bears particular significance for *social* animal 146 species mainly in three contexts: territoriality, aggressive competition and parental care 147 [9]. However, jumping spiders, as many other spider species, are solitary and aggressive 148 towards conspecifics, raising the question about the biological relevance of individual 149 recognition in *P. regius*: One of the few social instances in the life of a jumping spider 150 occurs during reproductive communication, encompassing a complex visual courtship 151 display of coordinated movement patterns of the body and bodily features. It is believed 152 that the typical colouration of the appendages (*Chelicerae*) and the colouration and 153 facial hair characteristics serve as important features for species and sex classification 154 in jumping spiders and as a general indicator about the quality of an individual as 155 a mating partner [14, 15]. Hence, colouration (sender) and the ability to distinguish 156 certain colours (receiver) seem to be sufficiently beneficial to sexual selection [16, 14]. 157 Similarly, in aggressive interactions, often due to territorial disputes, fighting abilities 158 are largely associated with the size and colour of the *Chelicerae* [16, 17], rendering 159 territoriality and aggressive competition needless as an ultimate explanation [18] for 160

individual recognition in *P. regius*. Some jumping spiders exhibited parental care [19], 161 protecting the nest through the spiderlings' first molt. One particular jumping spider 162 (Toxeus magnus) has been documented to provide a nutritious milk-like substance to 163 the spiderlings, which compares functionally and behaviourally to lactation in mammals 164 [20]. Whether individual recognition solves a survival problem in this context, is however 165 questionable. The remaining solitary behaviours of *P. regius* give little additional reason 166 to predict that individual recognition is a requirement for survival, setting the minimal 167 recognition needs (Minimum needs hypothesis [21, 2]) at the basic-level of classification 168 [22], e.g. colour-based distinction, size assessment of *Chelicerae*. Moreover, the neural 169 implementation of a subordinate-level classification [22] system that operates at the level 170 of abstraction required for a more detailed classification, such as individual recognition, 171 involves specialised processing in dedicated neural correlates [23]. For all these reasons it 172 is therefore more likely to assume that the ability of recognising conspecifics is related to 173 *P. regius*'s general learning capabilities via pleiotropy, also referred to as the 'generalised 174 learning hypothesis' [21]. For example, salticids' rather complex foraging and navigation 175 strategies [24, 25, 26], requiring high degrees of learning and adaptability, may translate 176 into the flexible learning ability required for recognising conspecifics, at a level of abstraction 177 more fine-grained than their minimum recognition needs [13]. In other words, while for 178 social animal species, including social arthropod species [6, 4, 7], there is an ultimate 179 explanation [18] addressing the function (or adaptation) of individual recognition, we 180 cannot conclusively infer the survival benefits gained by individual recognition in P. 181 regius. Instead we put forward the idea that individual recognition in P. regius is a 182 byproduct of fairly sophisticated cognitive processing capabilities. Critically, individual 183 recognition relies on recognition memory, a form of long-term memory, where a previously 184 encountered event or entity, here an individual, stored as memory representation is 185 neuronally activated upon re-experiencing that event or entity [27]. Such memory representation 186 might well serve as guidance to *Portia fimbriata*, allegedly the most intelligent jumping 187 spider, when, after scanning the access route to a prey target, it follows the path to 188 the prey under lack of visual control [28]. In our study, we demonstrated retrieval of 189 information from memory representations in various ways: First, P. regius's dissocative 190 behavioural responses upon perceiving an individual for a second time in succession as 191 opposed to perceiving a different individual suggests recognition of individually distinct 192

characteristics or cues, which manifest in solid memory representations. Moreover, P. 193 regius systematically reduced the overall interest over a series of repeated exposure to the 194 same individuals at a 1-hour presentation interval to the point where it became indifferent 195 to the presented individual, suggesting that *P. regius* successfully retrieves memory-stored 196 information at least one hour after memory consolidation. Thirdly, P. regius's interest in 197 novelty was restored at the end of session 3 upon perception of individuals that had not 198 been encountered before, highlighting that the loss of interest in the long-term was not 199 due to a general physical fatigue, but a 'cognitive' fatigue, i.e. to literally perceive the 200 same individuals over and over again. A novel individual, consequently, did not activate 201 memory representations of individuals, and led, as a response, to dishabituation, a fortiori 202 amplifying the notion of memory representation in *P. regius*. 203

Together, our study challenges the notion of spiders being stimulus-response driven 204 automata, by not only contributing to an increasing body of evidence that spiders and 205 saliticids in particular produce a wide spectrum of intelligent behaviour [29], but by 206 pinpointing the presence of two fundamentally important mechanisms for any higher 207 cognitive processing: flexible learning and recognition memory. The key building blocks 208 of these mechanisms are representations, mental images of external entities, that are 209 not present to the sense organs, allowing more elaborate information processing, such 210 as in complex decision making and goal-directed behaviour. The existence of which in 211 arthropods in general and spiders in particular triggers rethinking of miniature brain 212 cognition [29]. 213

#### <sup>214</sup> Materials and methods

#### 215 Subjects

Our subjects were 36 jumping spiders (*Phidippus regius*), kept individually in enclosures (7 x 7 x 12 cm) at room temperature (21 - 25°C) and supplied with a moist water-pad, exchanged every other day, and two small-sized cockroaches (*Shelfordella lateralis*) per week. All spiders were adult laboratory-bred and had no direct encounters with conspecifics during adulthood. Behavioural enrichment [30] was provided by means of climbing and nesting structures (i.e., natural wood branch) and by interaction with human caretakers and experimenters during handling and maintenance procedures. In Experiment 1,

spiders were assigned to five experimental groups, three of which contained females, two of which males; in Experiment 2, spiders were assigned to four experimental groups, i.e. two groups per sex.

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#### 227 Apparatus

In the following we describe how pairs of spiders were brought into direct visual contact 228 under controlled conditions and in a manner that allows to reassign individuals easily and 229 without interruption to form novel pairings. To this end, we built a cubical experimental 230 arena of 60 by 46 bx 65cm dimensions (L x W x H), consisting of white polypropylene 231 plastic panels, mounted in a frame of T-slotted aluminium profiles (20 Series; Misumi 232 Group Inc., Bunkyo City, Tokyo, Japan). Two LED light sources (Mettle<sup>®</sup> SL400, 233 45W, 2100lm, 350 x 250mm surface area, Mettle Photographic Equipment Corporation, 234 Changzhou, China) were placed outside the cubicle at 25cm distance from the side panels 235 of the cubicle, illuminating the inside of the cubicle uniformly. We also mounted two 236 FLIR® 1.3MP, Mono Blackfly USB3 cameras with a 1/2" CMOS sensor (BFS-U3-13Y3M-C, 237 FLIR<sup>®</sup> Integrated Imaging Solutions, Inc, 12051 Riverside Way, Richmond, BC, Canada) 238 equipped with 8mm UC Series lenses from Edmund Optics<sup>®</sup> (Stock #33-307, Edmund 239 Optics<sup>®</sup>, Barrington, New Jersey, USA) on T-slotted aluminium profiles, facing downwards 240 onto the arena surface at a distance of 60cm. For each spider we 3D-printed a white 241 container with outer dimensions (L x W x H) of 7 x 7 x 5cm and inner dimensions of 6.3 242  $x 6.3 \times 4.5$  cm. The upper side of the container and one of the four side walls were made 243 of a transparent .5mm thick acrylic sheet. While the acrylic sheet on the upper side of 244 the container was screwed onto the side walls of the container, the acrylic sheet on one of 245 the sides of the container can be lifted up to open the container, allowing easier transfer 246 of the spider from the home enclosure. 247

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#### 249 Procedure

In Experiment 1, each group involved four same-sex spiders, with each spider being placed inside a container prior to the experiment. During the experiment, the spiders remained in their own container. We allowed the spiders sufficient time (10-15min) to get used to the new environment. We then placed the containers of the four spiders such that the transparent side walls of two containers were facing each other, resulting in two pairs

of spiders with direct visual contact to each other. During the process of arranging the 255 containers and prior to the initiation of every new trial, visual contact was prevented 256 by an occluder slid between the transparent side walls of the containers. Each trial was 257 initiated by removing this occluder, allowing visual contact. For simplicity, let the four 258 individuals be symbolised by the letters 'A', 'B', 'C', and 'D': An arrangement of trials 259 where each individual is opposed to each other individual is described in Table 1. To tease 260 apart, whether or not *P. requis* was capable of visually discriminating other individuals 261 two types of trials were required: (a) a *habituation* trial, where the same individual was 262 presented in the trial preceding the current trial, and (b) a *dishabituation* trial, where a 263 different individual was presented in the trial preceding the current trial. Therefore, every 264 dishabituation trial followed a habituation trial, forming a habituation or dishabituation 265 phase, respectively, as shown in Table 2. A trial, e.g. A - B (and in parallel C - D), 266 lasted for 7 minutes allowing the spiders to visually inspect each other, before isolating 267 the spiders visually for 3 minutes with a non-transparent white occluder, fully covering 268 the transparent side wall. After the occluder phase, another exposure phase of 7 minutes 269 was initiated, which consisted of either the same individual (habituation trial) or another 270 individual (*dishabituation* trial) than the individual in the preceding trial. During each 271 trial, the individuals distance to each other was quantified at 10Hz temporal resolution, 272 and taken as a measure reflecting the 'interest' in each other: Short distances between 273 individuals signal greater 'interest' in each other, while large distances signal reduced 274 'interest' in each other. We predict a dissociation of distances between *habituation* and 275 *dishabituation* trials. With the outlined procedure (Table 2), we can form sequences of 276 exposure phases, where each first of two exposure phases is a habituation phase, and 277 every second of those exposure phases is a dishabituation phase and at the same time 278 a habituation phase for the subsequent exposure phase. In this manner, we created a 279 trial list, containing 12 trials in total, six of which result in *habituation* phases and six of 280 which result in *dishabituation* phases (Table 2). This session of trials was repeated twice, 281 resulting in a total of 36 trials per experiment. Each experiment lasted 180 min, where 282 each trial contained 7 min exposure and 3 min visual separation. Each group of spiders 283 was subjected to this protocol. Two amendments were introduced in Experiment 2: (a) 284 We ran two groups of four individuals in parallel, and (b) additional *cross-group* trials 285 were introduced at the end of session 3. This resulted in a modified procedure described 286

<sup>287</sup> in Table 3.

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#### 289 Data logging and analysis

Camera control and image acquisition were done using Matlab (Mathworks<sup>®</sup>, Natick, 290 Massachusetts, USA) and the image acquisition and processing toolboxes. The frame 291 rate was set to 10Hz. Cameras were placed perpendicular to the xy-plane at a distance of 292 about 60cm from the ground. The lens aperture was set to f/4, allowing a sufficient depth 293 of field. Analysis was done with Matlab (Mathworks<sup>®</sup>, Natick, Massachusetts, USA). We 294 pre-processed the video recordings by segmenting the spider body from the background 295 in each frame using functions for image intensity adjustment, image enhancement, image 296 binarization and image properties measurement to extract the largest available 'region', 297 the spider body, and its centroid. For each trial we approximated the distance between 298 the individuals in the xy-plane as a function of time, using the Euclidean distance 299 weight function based on the centroid coordinates of the two individuals. We then 300 pooled the distance values of each trial into 4 equally-sized and non-overlapping bins 301 (bin centers [mm]: [20, 60, 100, 140]; bin size 40mm; maximal distance  $\approx$  160mm) and 302 calculated the proportion of time spent at a given distance. Each bin was normalised 303 by the total number of events. Differences between proportions were then calculated 304 for every trial comparison according to Tables 2, 3: For instance, the proportions of 305 time spent at a given distance for individual A in trial 1 was subtracted from the 306 proportions of time spent at a given distance for individual A in trial 2, resulting in 307 an assessment for the relative rebound of interest following a repetition of exposure to 308 the same spider B (*habituation*). Subsequently, the proportions of time spent at a given 309 distance for individual A in trial 2 was subtracted from the proportions of time spent 310 at a given distance for individual A in trial 3, resulting in an assessment for the relative 311 rebound of interest following changes in spider's identity (dishabituation). We used 312 linear mixed-effects models, where the differences in proportions served as the dependent 313 variable. We fitted two separate models for each experiment (Full model 1 and 2), and 314 followed a commonly accepted model fitting procedure [31]: To fully account for the 315 dependent variable, we fitted three predictor variables: (1) The bin number ([1 to 4]), 316 reflecting a discretised distance measure and henceforth referred to as factor distance ([1 317 to 4]), (2) the session of comparisons ([1, 2, 3]), as outlined in the Procedure above (Table 318

2, 3), and (3) the *condition*, referring to whether the given comparison was a *habituation* 319 or *dishabituation* comparison. We also fitted all two-way interactions between the three 320 main predictors: distance: session, distance: condition, and session: condition, as well as the 321 three-way interaction *distance:session:condition*. Of particular interest are the two-way 322 interaction between the factors *distance* and *condition*, since we predict a modulation 323 of *distance* values by *condition* as a function of *distance*, and the three-way interaction 324 between the factors *distance*, *condition* and *session*, since we predict a modulation of 325 *condition* as a function of *distance* which becomes weaker over time and repetitions, i.e. 326 session. We further defined sex of the subjects and subject as random factors in all 327 models. We fitted a linear mixed-effects model (fitlme function in Matlab) with normal 328 error structure and identity link function to our data set. We then created a null model 329 for each corresponding full model, which consisted of the similar structure as the full 330 model, however leaving only *distance* as fixed effect, while preserving all random effects. 331 Using likelihood ratio test (LRT), we compared the null models with the corresponding 332 full models. Assuming a significant improvement for the full model over the null model, 333 the non-significant interaction terms were removed from the full model, reaching a model 334 containing only significant interaction terms and both significant and non-significant main 335 effects [32, 33], henceforth referred to as the final model. Evaluation of fixed effect were 336 on the basis of the final models and are referred to as the *Final model 1* (Experiment 337 1), and *Final model 2* (Experiment 2). This procedure resulted in the following models 338 (Wilkinson notation): 339

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341 Final model 1 and 2:
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<sup>342</sup> 'Response \sim 1 + \text{Distance} + \text{Session} + \text{Condition} + \text{Distance:Condition} + \dots
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<sup>343</sup> Distance:Session:Condition + (1|Sex) + (1|Subject)'
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An additional analysis of variance was performed comparing the *dishabituation [long-term]* trials at the end of session 3 with the *dishabituation [short-term]* trials from session 3 (Table 3) as a function of *distance*. No statistical methods were used to predetermine sample size. The experiments were not randomized. The investigators were not blinded to allocation during experiments and outcome assessment.

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## 430 Author contributions

431 CDD: study design, data collection, analysis and interpretation, writing article, provision
432 of necessary tools; YC: data collection, writing article, provision of resources.

#### 433 Competing interest

The authors declare that they have no competing interest. The authors have no affiliations
with or involvement in any organization or entity with any financial interest, or non-financial
interest in the subject matter or materials discussed in this manuscript.

#### 437 Additional information

All authors have seen and approved the manuscript. The manuscript has not been
accepted or published elsewhere. Supplementary Information is available for this paper.
Correspondence and requests for materials should be addressed to Christoph D. Dahl.
Codes and materials are available (https://osf.io/gpnct/).

## 442 Ethical approval

According to Taiwan's Animal Protection Act, issued by the Council of Agriculture
(Executive Yuan), experiments on invertebrates are allowed to be conducted without
any special permission in Taiwan.

## $_{446}$ Tables

Table 1: Pairwise comparisons.

Trial	Pair 1	Pair 2		
1	А - В	C - D		
2	A - C	B - D		
3	В - С	A - D		

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Table 2: Procedure of Experiment 1.

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Trial	Pair 1	Pair 2		
1 2 3 4 5 6 	A - B A - B A - C A - C B - C B - C	C - D C - D B - D B - D A - D A - D	Habituation Habituation Habituation	<pre>Dishabituation Dishabituation Dishabituation</pre>
3 sessions				

448

Table 3: Pairwise comparisons as habituation and dishabituation trials.

Group 1			Group 2			
Trial	Pair 1	Pair 2	Pair 1	Pair 2		
1	A - B	C - D	E - F	G - H	Habituation	
2	А - В	C - D	E - F	G - H	filabituation	Dishabituation
3	A - C	B - D	E - G	F - H	Habituation	$\int [\text{short-term}]$
4	A - C	B - D	E - G	F - H	Habituation	Dishabituation
5	В - С	A - D	F - G	Е-Н	Habituation	[short-term]
6	В - С	A - D	F - G	Е-Н	/ nabituation	)
 3 sessions						Dishabituation [long-term]
7	A - E	B - G	C - F	D - H	-	J

449

## 450 Figure

#### 451 Figure 1



<sup>452</sup> Figure caption 1: Theoretical assumptions and exemplar trials. a-c Predicted spider

distances for *baseline* (red dots), *habituation* (green dots) and *dishabituation* comparisons 453 (blue dots). Habituation can manifest either in equal inter-spider distances (solid line) 454 as in the *baseline* comparison or in an increase of distances (dashed line). What is 455 referred to as *baseline* in this context is the *dishabituation* trial of the previous comparison 456 (see Table 2). Distance samples are predicted to fall into distributions as shown in b. 457 Contrasts between *baseline*, *habituation* and *dishabituation* comparisons would result in 458 distributions as shown in c. d-f An exemplar trial consisting of *baseline*, *habituation* 459 and *dishabituation* comparisons from the first session of trials is shown. The *short-term* 460 dishabituation comparison shows a decrease of inter-spider distances, indicating increasing 461 interest in a different individual than the previously perceived one (*habituation* comparison). 462 g-i; j-l Two exemplar trials from the third session of Experiment 2 are shown, where a 463 presentation of an individual novel and unseen across the three experimental sessions 464 triggered a great rebound in interest (i,l, 'Dishabituation - habituation'). 465

#### <sup>466</sup> Figure 2



Figure caption 2: The relative change in distance between pairs of individuals, upon 467 being confronted with the same individual as in the preceding trial (habituation trial; red 468 discs) or a different individual from the individual in the preceding trial (dishabituation 469 trial; blue discs). Each panel refers to an experiment (panel a. for Experiment 1; panel 470 b. for Experiment 2), consisting of three sessions of trials. The dependent data is shown 471 as the proportion of time spent at a given distance binned into 4 equally spaced bins. 472 The x-axis labels refer to the proportional distances from the transparent acrylic sheet, 473 ranging from 'proximal' to 'distal'; the y-axis refers to the proportion of time spent at a 474 given distance, i.e. the relative number of samples that fall into a given bin. Discs show 475 the mean proportion across all individuals (i.e. 20 for Experiment 1; 16 for Experiment 476 2). The whiskers indicate the standard errors of the mean. White diamonds in the lower 477 right subfigure b show the long-term dishabituation trials. Light blue bars indicate the 478 side of the transparent acrylic sheet (proximal); grey bars indicate the back wall of the 479

480 container (distal).



#### Exemplar trial (short-term)



Exemplar trial (long-term)



x=0.25

x=0.38 x=0.22 x=0.19

