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1 Inheritance of Behavioral Traditions underlying Nest Architecture in Stingless Bees

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20 Summary

21 The transmission of complex behavior and culture in humans has long been attributed to advanced forms of social learning^{1,2}, which play a crucial role in our technological 22 advancement³. While similar phenomena of behavioral traditions and cultural inheritance have 23 been observed in animals^{1,2,4-6}, including in primates⁷, whales⁸, birds⁹, and even insects¹⁰, the 24 25 underlying mechanisms enabling the persistence of such animal traditions, particularly in 26 insects, are less well understood. This study introduces pioneering evidence of enduring 27 architectural traditions in the stingless bee Scaptotrigona depilis, which are maintained without 28 any evidence for social learning. We demonstrate that S. depilis exhibits two distinct nest architectures, comprising either helicoidal or flat, stacked horizontal combs, which are 29 transmitted across generations through stigmergy $^{11-17}$ – an environmental feedback mechanism 30 whereby the presence of the existing comb structures guides subsequent construction 31 behaviors, thereby leading to a form of environmental inheritance¹⁸⁻²⁰. Cross-fostering 32 experiments further show that genetic factors or prior experience do not drive the observed 33 34 variation in nest architecture. Moreover, the experimental introduction of corkscrew 35 dislocations within the combs prompted helicoidal building, confirming the use of stigmergic

building rules. At a theoretical level, we establish that the long-term equilibrium of building in the helicoidal pattern fits with the expectations of a two-state Markov chain model. Overall, our findings provide compelling evidence for the persistence of behavioral traditions in an insect based on a simple mechanism of environmental inheritance and stigmergic interactions, without requiring any sophisticated learning mechanism, thereby expanding our understanding of how traditions can be maintained in non-human species.

42

43 Keywords: animal traditions; environmental inheritance; stigmergy; nest architecture;
44 stingless bees

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46 **Results and discussion**

47 The progress and prosperity of human civilizations have been significantly influenced by our ability to learn from others and accumulate knowledge, leading to technological progress and 48 innovation that is stably transmitted through cultural mechanisms²¹. While long perceived to 49 be exclusive to humans, behavioral traditions and social information transmission across 50 generations have also been observed in a variety of non-human species^{1,2,4-6,22}. In particular, 51 primates⁷, cetaceans⁸, birds⁹, and even insects¹⁰ have all demonstrated the capacity to maintain 52 and transmit complex behaviors or structures across generations. High cognitive skills and 53 54 advanced forms of social learning have long been thought to be required for the stable maintenance of behavioral traditions and culture^{7-9,23,24}. However, the persistence of distinct 55 traditions also in species with simpler cognitive capacities, like insects^{10,25}, challenges the 56 conventional understanding of cultural transmission, suggesting that alternative, less complex 57 mechanisms might sometimes be at $play^{10,25-28}$. 58

Recent insights into the architectural diversity of stingless bee colonies hint at the existence of 59 behavioral traditions that might not rely on direct social learning or high cognitive abilities²⁵. 60 61 In this case, it is likely that variation in nest architecture could be propagated based on the 62 comb structure itself, whereby the current building style guides the building by later generations of bees. Such a process could lead to environmental and behavioral inheritance¹⁸⁻ 63 ^{20,27,28}, even in the absence of any direct form of social learning. To test this theory, we here 64 present the first study of behavioral innovation and social transmission of alternative nest 65 66 architecture building patterns in a neotropical stingless bee, Scaptotrigona depilis. This species is one of 10 known stingless bee species that exhibit significant intraspecific variation in nest 67 architecture²⁹, building its brood combs in either a helicoidal shape built continuously without 68

69 the support of a central pillar, or as horizontally stacked parallel layers, supported by a central 70 pillar^{12,14} (Figure 1A,B). Given that stingless bees generate perennial colonies and reproduce 71 through swarming or managed splitting, we hypothesized that the observed architectural 72 variation might be maintained over periods of time extending beyond the lifespan of individual 73 worker cohorts.

74 To investigate the inheritance and prevalence of the two alternative nest architectural building patterns, we surveyed 413 S. depilis colonies in Jaguariúna, São Paulo State, Brazil, and 75 76 monitored the incidence and changes in nest architecture over one-month periods in 2022 and 77 2023 (see STAR Methods). We hypothesized that the prevailing comb configuration would guide the construction of successive worker cohorts²⁵, with the frequencies of helicoidal and 78 79 parallel architectures reflecting the stable equilibrium of a two-state continuous time Markov 80 chain model. Specifically, if a and b represent the monthly transition rates to helicoidal and parallel comb structures, each occurring at frequencies H and P, we predicted the dynamics to 81 82 adhere to the following system of ordinary differential equations:

$$\frac{dH}{dt} = aP - bH$$

$$\frac{dP}{dt} = bH - aP$$

At equilibrium (dH/dt = 0 and dP/dt = 0), and given that H + P = 1, the expected equilibrium 85 proportion building in the helicoidal configuration is $H^* = a/(a + b)$, while for the parallel 86 configuration it is $P^* = b/(a + b)$. Our empirical data show that parallel-building colonies (n 87 = 398) transitioned to a helicoidal architecture at a rate of a = 4.5% per month (binomial 95%) 88 CrI 2.8-6.9% per month), while helicoidal-building colonies (n = 15) switched to parallel at a 89 rate b = 87% per month (binomial 95% CrI 64-97% per month). These rates did not differ 90 significantly between the study years, supporting the Markov chain model's assumption of 91 constant transition rates (binomial GLM, posthoc tests, z ratio = -1.53, p = 0.13 and z ratio = 92 0.32, p = 0.75 for contrasts in switching rates a and b between years). 93

The calculated equilibrium frequency of the helicoidal architecture H^* of 5.0% (parametric bootstrapping 95% CI: 2.9-7.6%, n = 15) aligns with the observed average frequency over both years (binomial GLM, 4.3% with 95% CI 2.7-6.7%, n = 413, z ratio = -0.652, p = 0.51) and was also not significantly impacted by season or year (Jan.-Febr. 2022 vs. Aug.-Sept. 2023),

98 hive entrance orientation, or position within the bee yard (Figure S1 & Table S1). It has been suggested that a helicoidal nest architecture could be a thermoregulatory adaptation helping to 99 maintain the brood area at an optimum temperature^{30,31}. Nevertheless, our results indirectly 100 101 dismiss this hypothesis, since we would then have expected the helicoidal building type to be 102 spatially clustered or to be more common in the warm season (Jan.-Feb.), which was not the 103 case (Table S1). The inclination of the hives also did not impact the frequency of both types. 104 When 10 colonies were tilted, none of them switched to building helicoidal combs within 2 105 weeks, and on a per-month basis the rate of switching to the helicoidal type was not 106 significantly elevated compared to 2022 parallel-building control colonies (n = 299) (Firth's penalized logistic regression, z ratio = 0.66, p = 0.51). The lack of any significant 107 environmental effects argues against variation in nest architecture being driven by changes in 108 the external environment^{30,31}. Our results confirm the long-term persistence of both 109 architectural phenotypes, but with the helicoidal form continuing for a shorter average duration 110 $(T_H = 1/b = 1.2 \text{ months}, 1.0-1.5 95\% \text{ CI})$ than the parallel form $(T_P = 1/a = 23.4 \text{ months}, 14.7-1.5 \text{ months})$ 111 39.8 95% CI). Yet, the average duration of building in the helicoidal pattern still surpassed the 112 113 typical comb-building period of a single worker cohort, which transitions to other tasks after approximately two to three weeks²⁹. Consequently, our results imply that information on how 114 115 combs are constructed is transmitted across different worker cohorts and hence that some form of environmental and behavioral inheritance is going on^{18-20,27,28}. 116

To investigate the specific mechanism by which this transmission occurs, we carried out a full 117 factorial cross-fostering experiment in which we provided experienced workers sourced from 118 colonies building in either the helicoidal or parallel pattern with a comb lattice structure from 119 120 a genetically unrelated colony that was matching or was opposite to their own (see STAR 121 Methods, Figure 1, n = 6 replicates per treatment combination). According to our stigmergy-122 based environmental inheritance hypothesis, the expectation was that presenting an altered 123 comb structure in itself should cause the experienced workers to switch to a new building style, 124 merely by being guided by the comb structure itself. In that case, social information on how combs are built would be socially transmitted across worker cohorts, but it would be based on 125 126 environmental inheritance driven by stigmergy, whereby the change in the structure of the 127 environment – the change in the comb lattice structure – would act as a stimulus for continued construction in a manner consistent with the existing architecture¹¹⁻¹⁷. This form of social 128 transmission emphasizes the role of the artifacts themselves – in this case, the comb structure 129 130 - as repositories and conduits of social information, thereby bypassing the need for direct observation of behavior or instruction. By contrast, if the experienced workers were predisposed towards building in a particular style, based on their genetic makeup or prior experience, the expectation was that they should continue to do so even when provided with a comb template of the alternative type.

135 Our cross-fostering experiment results confirm the environmental inheritance hypothesis. 136 When exposed to alternative architectural styles, experienced workers did not adhere to their 137 original construction patterns, thereby implying that nest architectural variation was not 138 controlled by genes of large effect or by prior experience. Instead, they promptly adjusted their 139 behavior to align with the provided comb structure, as evidenced by the rapid change in 140 building patterns observed within the first week – a change that remained constant for several 141 weeks (Figure 2). These results are consistent with the hypothesis that stigmergy underlies the 142 observed environmental inheritance and the persistence of distinct nest architectural behavioral traditions. Nevertheless, over time, most colonies did eventually revert to building in the 143 144 parallel configuration, consistent with the dominance of this type in the wild. Given that the 145 observed changes in nest architecture were not affected by the original building style of the 146 experienced workers, our results clearly argue against genetic factors of large effects (e.g. 147 supergenes^{32,33}) or direct social learning playing a significant role. Instead, our results support 148 the concept that behavioral traditions can be maintained through environmental cues and 149 stigmergic responses, aligning with the broader definition of traditions as patterns perpetuated 150 over time by mechanisms that do not require direct social learning^{27,28}.

To test if stigmergic building rules underpin the observed variation in nest architecture, we carried out one final experiment in which we incised the top comb of parallel-building colonies to mimic "corkscrew dislocations", which are known theoretical precursors to helicoidal comb building¹⁴. In line with theoretical predictions¹⁴, we found that this manipulation of the comb substrate caused a marked increase in the transition to a helicoidal architecture compared to control colonies (Figure 3). This result corroborates that the physical structure of the comb itself can direct subsequent construction, in line with the concept of stigmergy^{11,13,16}.

Despite our clear conclusions on the mode of inheritance of alternative nest architectures in this species, several important questions remain. It is unclear, for example, what factors cause the actual transitions between building styles, and why helicoidal building tends to persist for a much shorter time than parallel building. In all likelihood, transitions to parallel building are initiated by the building of a central wax support pillar¹² (Figure 1A), while transitions to 163 helicoidal building are caused by occasional mistakes in the vertical alignment of newly built cells¹⁴. If correct, this suggests that the parallel configuration is the default building pattern and 164 165 that switching to the helicoidal pattern could be regarded as an error-correction mechanism to 166 absorb occasional building mistakes and avoid costly interruptions in cell building. Similar 167 error-correction mechanisms during comb construction have previously been described in other social insects³⁴⁻³⁷. Honeybees, for example, initiate comb building at multiple points within a 168 frame, and to be able to fuse those comb sections they occasionally switch to building 169 pentagonal or heptagonal cells instead of the standard hexagonal ones^{36,37}. This adaptability 170 reflects a sophisticated combination of stigmergy and behavioral flexibility, where bees 171 respond dynamically to the challenges presented by their construction environment^{36,37}. In any 172 173 case, our findings indicate no clear superiority of either architectural style, given that neither 174 significantly impacted cell construction rates (Figure S4). This absence of a distinct advantage 175 may explain the coexistence of both architectural styles in S. depilis, despite the exclusive adoption of the helicoidal style in some other stingless bee species^{12,14,29}. 176

177 Another important outstanding question is if the persistence of distinct nest architectures in our 178 system could also involve more complex forms of social learning, which some authors take as a defining characteristic of behavioral or cultural transmission^{6,38,39}. Indeed, Chittka & Rossi²⁵ 179 180 suggested that stingless bees might to some extent be able to copy how to build their comb 181 merely by observing and being exposed to the existing comb structure, which would be akin to an indirect form of social learning known as product learning⁴⁰. At present, we cannot exclude 182 that the stigmergy we describe also involves such a learning process and is not just based on 183 184 innate responses, but conclusive evidence to support this theory is currently lacking. 185 Personally, we believe it is more likely that the stigmergic building rules are largely innate, and 186 that the same set of building rules produce different nest architectures depending on the initial 187 comb substrate, or that particular innate building rules are conditionally expressed in function 188 of the existing architecture.

Testing whether or not the inheritance of alternative nest architecture is purely innate remains a task for future work and would require extensive and complex experiments. Establishing the involvement of social learning would entail showing that prior exposure to a specific comb structure affects the construction approach in subsequent generations, following swarming events. Suggestive evidence for this has been collected in the honeybee, where colonies maintain their original comb building orientation after swarming⁴¹. Nevertheless, as this study 195 did not use a cross-fostering setup, it could not preclude that the building style persisted merely as a result of varying genetic predispositions⁴¹. In addition, definite conclusions for our system 196 197 might be precluded by the fact that colonies presumably have a propensity to revert back to 198 parallel building when placed in an empty hive without any comb substrate.

199 In conclusion, our findings provide a compelling case for the role of stigmergy and 200 environmental feedback in the transmission of nest-building traditions in stingless bees. This 201 mechanism facilitates the persistence of distinct architectural styles within colonies, thereby 202 producing behavioral traditions that persist across the lifespan of individual bees, even in the 203 absence of any evidence for learning or advanced cognitive processing. Hence, our study 204 strengthens the case that animal traditions and the non-genetic transmission of behavior can be based on cognitively simple processes^{27,28}, such as via stigmergy-mediated environmental 205 feedback. Future research should aim to dissect the precise nature of these stigmergic 206 207 interactions further and explore whether beyond innate responses, there might also be a layer of learning that influences these behaviours²⁵. These investigations could shed light on the 208 continuum between innate building rules and the potential for learning in shaping the cultural 209 210 and behavioral landscapes of non-human societies. Our study not only advances our 211 understanding of stingless bee behavior but also contributes to the broader discussion on the 212 mechanisms of behavioral transmission and tradition maintenance in animal societies.

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223

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221 **Author contributions**

222 R.C.O. and T.W. conceived and supervised the study. V.D.P, C.M. and R.C.O.: experimental design. V.D.P, C.M., M.G.B.F. and D.J.P.: behavioral experiments. T.W.: theoretical

modelling. V.D.P., M.F., R.C.O., T.W. and H.M.F.: data analysis. V.D.P and H.M.F wrote the 224

- first draft of the manuscript. All authors contributed to revising and proofreading and approved
- 226 of the final version before submission.

227 Declaration of interests

228 The authors declare no competing interests.



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Figure 1. Cross-fostering experiment to study how intraspecific variation in nest 230 231 architecture in the stingless bee Scaptotrigona depilis is transmitted. Strikingly, workers in this species construct their brood combs either in a parallel arrangement (A) with a central 232 233 support pillar from which the first cell of the comb is built (*) or in a helicoidal pattern (**B**), 234 without any central pillar (+). To study the mechanism by which this variation in nest 235 architecture is transmitted, we carried out a full factorial cross-fostering experiment (panels C-**F**), where we provided experienced workers (EW), deriving from colonies with either parallel 236 237 (blue) or helicoidal building patterns (red), with a comb lattice structure that was either of the 238 same type or the alternative type. The combs used as a template in the experiment contained 239 late-stage genetically unrelated brood that was ready to emerge, to allow for interactions and 240 potential social information exchange between experienced workers and naïve young workers 241 (EW and YW).





244 Figure 2. Stigmergy, rather than genetics or prior experience, drives the inheritance of 245 alternative nest architectures. Estimated changes in nest architecture over 7 weeks in our 246 cross-fostering experiment (n = 6 replicates/treatment). When experienced workers (EW), with 247 a history of building in either parallel or helicoidal styles, encountered the opposing comb structure, they immediately switched to building in the presented style (panels C and D). This 248 249 swift change was likely driven by stigmergy, where the existing comb structure guided 250 subsequent building. These results also clearly show that nest architecture was not determined 251 by prior experience or genetic makeup. As the weeks progressed, most colonies eventually 252 switched towards the parallel style (panels A-D), mirroring its dominance in the wild and the 253 observed preferential switching to this building style. The presence of EWs from parallel-254 building colonies did not accelerate this shift (panels **B** and **D**), thereby implying that direct 255 social learning did not significantly influence building patterns. The shaded areas denote 95% credible intervals from a Bayesian Gaussian process logistic regression model (see STAR 256 257 Methods). For detailed results see Figures S2 & S3.

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261 Figure 3. Stigmergy underlies the transition to helicoidal comb building. To investigate the role of stigmergy in the social transmission of helicoidal comb construction, we modified 262 263 the top comb of colonies that were previously building in a parallel pattern by incising and tilting one side to simulate "screw dislocations", a theoretically identified initiator of helicoidal 264 comb building¹⁴ (A,B). With only the comb's lattice structure changed and all workers 265 inexperienced in helicoidal building, a subsequent shift to this construction style would 266 267 implicate stigmergy. Results confirmed this hypothesis: 43% of the manipulated colonies 268 (9/21) transitioned to helicoidal building within one week (c, red line indicates initial incision, 269 obscured by newly built helicoidal comb), which amounted to a monthly transition rate to 270 helicoidal building of 75.0% [55.8-87.7] 95% CI (*n* = 21) compared to a mere 3.1% [1.5-6.3] 95% CI in the parallel-building 2022 controls (n = 299) (binomial GLM analysis, odds ratio to 271 272 switch over one month: 95.1 [30.3-299.2] 95% CI, *z* ratio = 7.79, *p* < 1E-14).

- 273
- 274
- 275 STAR Methods

276 **RESOURCE AVAILABILITY**

277 Lead contact

Further information and requests for resources should be directed to and will be fulfilled by the
lead contact, Viviana Di Pietro (viviana.dipietro@kuleuven.be).

280 *Materials availability*

281 This study did not generate new unique reagents.

282 Data and code availability

All datasets used for analysis in this study have been deposited at Mendeley Data 283 284 (10.17632/bhjp3m2sbp.1) and will be publicly available as of the date of publication. Accession numbers are listed in the key resources table. For review the data can be downloaded 285 286 through this link provided by Mendeley Data: 287 https://data.mendeley.com/preview/bhjp3m2sbp?a=cd4b4d79-ba16-4248-8754-1af2f5eda915

- All original code has been deposited at Mendeley Data and is publicly available as of the dateof publication. DOI is listed in the key resources table.
- Any additional information required to reanalyze the data reported in this paper is availablefrom the lead contact upon request.

292 EXPERIMENTAL MODEL AND STUDY PARTICIPANT DETAILS

293 Stingless bees, a group of more than 500 highly eusocial bees in the tribe of the Meliponini, display an astounding diversity in terms of diet, behaviour and nest architecture, not only 294 between but also within species^{29,42,43}. Unlike honeybees, which construct vertical parallel wax 295 combs and use them both for food storage and brood rearing⁴⁴, the nest building of stingless 296 bees is more complex and diverse⁴². They use a range of building materials, including soil 297 particles, resin, plant fibres, and even excrement^{29,42,45,46} with food pots and brood combs being 298 spatially separated⁴². Nest architecture can vary both across and within species, with brood 299 300 cells being arranged either in horizontal combs, vertical combs, clustered cells or some intermediates between these forms²⁹. Our focus in this study is on *Scaptotrigona depilis*, a 301 Neotropical stingless bees and one of 10 species that are known to show large bimodal 302 intraspecific variation in nest architecture²⁹. The species constructs nests with combs consisting 303 304 either of flat, stacked parallel combs, supported by a central pillar, or builds combs arranged in 305 a helicoidal configuration, in which case there is no central support pillar. Both forms of comb can coexist within the same colony simultaneosly⁴⁷, suggesting that minor changes in building 306 behaviour can cause a switch between both configurations^{14,48} and the switch from one 307 308 configuration to the other can occur at any stage during building. The precise trigger for such 309 transitions remains unknown. The colonies used for this study were kept at Embrapa Meio

Ambiente in Jaguariúna, São Paulo, Brazil. These colonies were housed in identical wooden
 boxes (19 x 19 x 16 cm) and the bees were allowed to forage freely in their natural environment.

312

313 METHOD DETAILS

314 Observational data collection

To determine the frequency of helicoidal and parallel comb building in the stingless bee Scaptotrigona depilis and determine the rate of switching between both types within the same colony, we recorded the nest architecture of 413 colonies at one month intervals, in January-February 2022 (n = 241), and August-September 2023 (n = 172). Hive entrance orientation and position in the bee yard (placed individually on a stand or placed on the upper or lower shelf under a shelter) were noted to evaluate environmental influences on nest architecture.

321 Cross-fostering experiment

322 Our factorial cross-fostering experiment involved four groups with six colonies each: two 323 control groups with congruent comb structures and worker backgrounds, and two treatment 324 groups with mismatched comb structures and worker backgrounds (Figure 1). Standardized 325 hives (19 x 19 x 16 cm) were populated with three to four layers of brood combs from which 326 bees where ready to emerge, the original mother queen and approximately 600 genetically 327 unrelated bees of varying ages, from the youngest to foragers, of a given nest architectural 328 background. After a one-week acclimation period during which colonies were kept closed and 329 where they were provided with wax, pollen and honey, we tracked nest architecture weekly for 330 seven weeks, in a blinded manner, between February and March 2022.

331 Stigmergy experiment and hive inclination effect

To investigate the role of stigmergy in the social transmission of helicoidal comb building, we experimentally manipulated small newly built brood comb layers (diameter ca. 5 cm) of 21 parallel-building colonies by making an incision and tilting up one side to mimic "screw dislocations", which theoretically have been posited to initiate helicoidal comb building¹⁴ (Figure 3). After seven days, we recorded the nest architecture to check if helicoidal comb building had stably persisted.

In a similar fashion, we examined whether inclination had any impact on nest configuration by

selecting 10 colonies that were building in parallel and tilting them at an angle of 20 degrees.

340 Following a two-week interval, we examined these colonies to determine if any of them had

341 transitioned to helicoidal architecture.

342 *Colony performance analysis*

343 To test if nest architecture affected colony performance, six free-foraging observation hives 344 (three per nest architecture) of similar colony size were set up and videotaped to allow us to 345 measure and compare cell building rates. The brood combs were exposed by removing the 346 involucrum, and colony temperature was maintained with halogen lights. The number of newly built cells over 5 subsequent days were measured from the video footage using ImageJ's "Cell 347 Counter" plugin⁴⁹. Colony size was controlled for by using that same plugin to count the 348 349 average number of workers visible in 20 video frames collected at 30-second intervals for a 350 period of 10 minutes.

351

352 QUANTIFICATION AND STATISTICAL ANALYSIS

353 *Observational data collection*

Bayesian binomial credible intervals for the frequency of each nest architecture and the 354 monthly transition rates were estimated under a Jeffrey's prior using the *propCI* function from 355 the prevalence package⁵⁰ in R. We employed binomial generalized linear models (GLM) with 356 357 a logit link function to test for annual and environmental variations in architecture switching rates, using the *glm* function in R and carried out two-sided Sidak posthoc comparisons with 358 the emmeans package⁵¹ (Figure S1). To derive confidence intervals for the equilibrium 359 frequency of helicoidal types and the expected duration of architectural forms, we applied 360 361 parametric bootstrapping with one million replicates. All statistical analyses were performed in R v. 4.2.2⁵². 362

363 *Cross-fostering experiment*

We employed a Bayesian Gaussian process logistic regression model⁵³, fit using the *brms* 364 package⁵⁴, to analyses the change in nest architecture over time. This approach allowed us to 365 model time series data flexibly, accounting for potential autocorrelations and inter-colony 366 367 variability by including a random intercept for each colony. We assumed an exponentiated 368 quadratic covariance function to describe the covariance between two time points as a function 369 of the number of weeks that separates them. We selected weakly informative zero-centered 370 normal priors for intercepts and scales (with a SD of 3), and a moderately informative inverse 371 gamma prior (with a shape and scale of 5) for length scales. We ran four MCMC chains of 372 2,000 iterations each, of which the first 1,000 iterations were discarded as warmup. Convergence diagnostics included inspection of traceplots and calculation of the Potential 373 Scale Reduction Factor (R-hat), which for all parameters was smaller than 1.01⁵⁵. To visualize 374

the effects of time and treatment on the outcome type, we plotted the conditional temporal trajectory of each treatment along with 95% equal-tailed credible intervals (Figure 2). We also performed a prior sensitivity analysis to ascertain that our findings were robust against alternative prior specifications (Figure S2). In addition, we visualized the posterior distributions of the estimated fraction of parallel comb architecture for each treatment group at each week using the *ggdist* package⁵⁶ (Figure S3). Posterior estimates were compared pairwise among treatments and weeks.

382 Stigmergy experiment and hive inclination effect

A logit link binomial GLM was used to compare the rate of switching to helicoidal comb building in manipulated colonies (n = 21) vs. in unmanipulated parallel-building control colonies observed over the same time period (February 2022, n = 299). Differences in the period over which switching was observed, i.e. one week versus one month, were adjusted by including log(period) as a model offset. To facilitate comparison, switching rates were expressed on a fixed per month basis.

A binomial GLM with logit link function was employed to compare the rate of transitioning to helicoidal comb construction in tilted colonies (n = 10) with that in unmanipulated control colonies that naturally built parallel combs during the same observation period (February 2022, n = 299). Given that there was separation, this model was fitted using the *brglm2* package⁵⁷ using a Jeffrey's prior. To account for differences in the observation period in the manipulated vs. in the control colonies (two weeks versus one month), we made adjustments by including log(period) as a model offset.

396 *Colony performance analysis*

The data were then analyzed using a Poisson GLMM with a log link function fitted using the *lme4* package⁵⁸. The model included time, nest architecture, and their interaction as fixed effects, log(colony size) as a model offset and colony ID as a random intercept. Overdispersion was addressed by incorporating an observational-level random effect. Two-sided Sidak posthoc pairwise comparisons and estimation of marginal means were performed with the *emmeans* package⁵¹.

403 Supplemental Information figures and tables



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Figure S1. The frequency of the helicoidal nest architecture was consistent across time 405 406 and space and not significantly affected by environmental factors. The frequency of the 407 helicoidal nest architecture in Scaptotrigona depilis was not significantly affected by season & 408 year (January-February 2022, n = 241 colonies, vs. August-September 2023, n = 172 colonies), hive entrance orientation (facing north, east, south or west) or position in the apiary (placed 409 410 individually on a stand, or on the bottom or top shelf in the apiary) (main effects logistic regression, anova Type III likelihood-ratio tests, effect for year/season: $\chi_1^2 = 0.095$, p = 0.76, 411 effect for hive entrance orientation: $\chi_3^2 = 2.65$, p = 0.45, effect for position in the apiary: $\chi_2^2 =$ 412 0.23, p = 0.89). Plots show the expected marginal means and Sidak adjusted 95% confidence 413 intervals, calculated using the emmeans package. As each colony was observed twice per year 414 415 with a one month interval, we used prior observation weights of 0.5 in our logistic regression 416 to avoid pseudoreplication.





418 Figure S2. Prior sensitivity analysis of nest architecture change over time in the cross-419 fostering experiment. Plots show the estimated percentage of colonies building parallel combs 420 through time for the four treatment groups (n = 6 replicates per treatment combination, Figures 421 1 & 2) under five different prior specifications for a Gaussian process logistic regression fitted using *brms*, in which colony was included as a random intercept (n = 6 colony replicates per 422 423 treatment combination, data were collected weekly). Full lines indicate posterior mean 424 trajectories, while dashed lines and the shaded zones show 95% credible intervals. The regular 425 prior specification (a Normal (0, 3) prior on the model intercept, an Inv-Gamma (5, 5) prior on the Gaussian process' length scale, a Normal (0, 3) prior on the Gaussian process' marginal 426 427 scale and a Normal (0, 3) prior on the random intercepts' scale), used for the main analysis 428 (Figure 2), is shown in blue. Results of an alternative model in which the Gaussian process' 429 length scale is replaced by an Inv-Gamma (18.5, 10) prior, favoring slower temporal changes, 430 is shown in red. Results of a model in which the Gaussian process' length scale is replaced by 431 an Inv-Gamma (18.5, 5) prior, favoring faster temporal changes, is shown in green. A model 432 in which wider Normal (0, 9) priors on the model intercept, the Gaussian process' marginal 433 scale and the random intercepts' scale are used, is shown in orange. Finally, a model in which tighter Normal (0, 1) priors on the model intercept, the Gaussian process' marginal scale and 434 the random intercepts' scale are used, is shown in purple. The prior sensitivity analysis reveals 435 436 that alternative prior specifications do not affect the general patterns and qualitative 437 conclusions presented in the main analysis, and only lead to minor changes in the width of the credible intervals. 438



Treatment (comb template - old worker origin)

440 Figure S3. Weekly estimates of nest architecture variation in transplant experiment. The 441 figure presents the model's predicted probabilities of colonies building in the parallel pattern across different treatments plus their uncertainty at 1 to 7 weeks after the start of the experiment 442 443 (points and line ranges show median posteriors plus 95% credible intervals, n = 6 replicates per treatment combination). Treatments show the combinations of the provided comb lattice 444 445 template (parallel or helicoidal) and the origin of the experienced old workers (from colonies 446 that previously built in the parallel or helicoidal arrangement), i.e. "par-par", "hel-par", "par-447 hel" or "hel-hel" (cf. Figures 1 & 2). The estimates are derived from a Bayesian Gaussian process logistic regression model, which incorporated random intercepts for individual bee 448 449 colonies. The colors distinguish between treatments, with blue and red indicating the 450 architecture of the provided comb lattice structure (parallel or helicoidal).

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Figure S4. Mean cell building rates in colonies with parallel and helicoidal nest 453 454 architectures. This figure illustrates the average number of new cells constructed per day over 455 a period of 5 days in colonies kept in observation hives that were constructing their combs in 456 either a parallel or a helicoidal arrangement (n = 6 colony replicates for each). The plotted estimates show marginal means and 95% confidence intervals derived from a Poisson 457 generalized linear mixed model (GLMM), calculated using the *emmeans* package (for details 458 see Table S1). No significant differences were found at any observation day, which shows that 459 460 nest architecture did not significantly affect colony performance.

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462

463 Table S1. Analysis of cell building rates in colonies with parallel and helicoidal nest 464 architectures. (A) Model coefficients of a Poisson generalized linear mixed model (GLMM) 465 in which we compared the number of new cells built per day, measured at 24-hour intervals for a period of five days in function of nest architecture (n = 6 colony replicates per type). The 466 467 model accounts for fixed effects time (day of observation) and nest architecture plus their 468 interaction, and included the natural logarithm of colony size as a model offset (average count 469 of workers visible in 20 video frames collected at 30-second intervals for a period of 10 470 minutes). Colony ID was included as a random intercept and an observation-level random 471 effect was included to take into account overdispersion. (B) Pairwise contrasts in daily cell 472 building rates between the two nest architectures show no significant differences in building 473 efficiency (ratio in the number of new cells built per day for colonies of average size) at any of 474 the observation days. The table shows coefficients, standard errors (SE), lower and upper 95% confidence intervals (*LCL* and *UCL*), z values and p values (NS = not significant, i.e. p > 0.05). 475

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(A) Model coefficients	term	coefficient	SE	z value	p value	
fixed effect	(Intercept)	-0.15	0.19	-0.80	0.42	
fixed effect	scale(time)	0.03	0.05	0.57	0.57	NS
fixed effect	nest_architecture1	-0.04	0.19	-0.23	0.82	NS
fixed effect	scale(time):nest_architecture1	-0.08	0.05	-1.54	0.12	NS
random intercept: observation	SD(Intercept)	0.19				
random intercept: colony ID	SD(Intercept)	0.46				
(B) Contrast (day)	ratio	LCL	UCL	z ratio	<i>p</i> value	
helicoidal / parallel (1)	0.95	0.43	2.09	-0.12	0.90	NS
helicoidal / parallel (2)	1.04	0.49	2.24	0.11	0.91	NS
helicoidal / parallel (3)	1.14	0.54	2.43	0.35	0.73	NS
helicoidal / parallel (4)	1.26	0.58	2.69	0.58	0.56	NS
helicoidal / parallel (5)	1.38	0.62	3.04	0.79	0.43	NS

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