

1 The adaptive value of tandem communication in ants: insights
2 from an agent-based model

3 Natascha Goy¹, Simone M. Glaser¹ & Christoph Grüter^{1,2}

4 ¹Institute of Organismic and Molecular Evolutionary Biology, Johannes-Gutenberg University
5 of Mainz, Mainz, Germany.

6 ²Present address: School of Biological Sciences, University of Bristol, 24 Tyndall Avenue,
7 BS8 1TQ Bristol, UK

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9 **Correspondence:** goyna@students.uni-mainz.de

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20 **Abstract**

21 Social animals often share information about the location of resources, such as a food source or
22 a new nest-site. One well-studied communication strategy in ants is tandem running, whereby
23 a leader guides a recruit to a resource. Tandem running is considered an example of animal
24 teaching because a leader adjusts her behaviour and invests time to help another ant to learn the
25 location of a resource more efficiently. Tandem running also has costs, such as waiting inside
26 the nest for a leader and a reduced walking speed. Whether and when these costs outweigh the
27 benefits of tandem running is not well understood. We developed an agent-based simulation
28 model to investigate the conditions that favour communication by tandem running during
29 foraging. We predicted that the spatio-temporal distribution of food sources, colony size and
30 the ratio of scouts and recruits affect colony foraging success. Our results suggest that
31 communication is favoured when food sources are hard to find, of variable quality and long
32 lasting. These results mirror the findings of simulations of honeybee communication. Scouts
33 locate food sources faster than tandem followers in some environments, suggesting that tandem
34 running may fulfil the criteria of teaching only in some situations. Furthermore, tandem running
35 was only beneficial above a critical colony size threshold. Taken together, our model suggests
36 that there is a considerable parameter range that favours colonies that do not use communication,
37 which could explain why many social insects with small colony sizes forage solitarily.

38

39 **Introduction**

40 Finding food is critical for survival and reproduction, but also energy- and time-consuming.
41 Foraging for food can be done independently or by using information provided by other
42 organisms (Sumpter 2010; Hoppitt and Laland 2013). In social insects, such as ants, social bees
43 or social wasps, new food sources are usually discovered by scouts that explore the environment

44 on their own (Hölldobler and Wilson 1990; Seeley 1995). After finding a profitable food source,
45 they return to their nest and often communicate their discovery to nestmates. The communicated
46 information depends on the species, but often includes the location of the resource, e.g. by
47 means of laying a pheromone trail (Hölldobler and Wilson 1990; Jarau and Hrnčir 2009;
48 Czaczkes et al. 2015b). Recruitment communication allows colonies to exploit profitable
49 feeding sites fast, e.g. before competitors have discovered and consumed the food source. Once
50 the foragers have learned the location of the food source, they can use their route memory to
51 return to the feeding site (e.g. von Frisch 1967; Collett et al. 2013).

52 In social insects, foraging strategies should not only take into account short-term
53 individual success, but also how they affect colony foraging success. Thus, the value of
54 communication should ultimately be studied at the colony level. So far, most theoretical and
55 empirical studies that explored the value of communication for colony foraging success have
56 focused on honeybees (but see also e.g. Sumpter and Pratt 2003; Dechaume-Moncharmont et
57 al. 2005; Czaczkes et al. 2015a). These studies suggest that the value of communicating the
58 location of food sources by means of waggle dances depends on how food sources are
59 distributed (Sherman and Visscher 2002; Dornhaus and Chittka 2004; Dornhaus et al. 2006;
60 Beekman and Lew 2008; Donaldson-Matasci and Dornhaus 2012; Schürch and Grüter 2014;
61 I'Anson Price et al. 2019; reviewed in I'Anson Price and Grüter 2015). For example, Beekman
62 & Lew (2008) found that the value of the “dance language” (the spatial information provided
63 by the waggle dance) depends on the size and distance of the food patches. When patches were
64 large and close to the hive, colonies that did not use dance communication and instead followed
65 an individual foraging strategy were more successful. Dornhaus et al. (2006) concluded that
66 dance communication does not help colonies collect more energy if there are many food sources
67 that vary little in quality. Their models suggest that communication is beneficial if high-quality
68 food sources are available, but are hard to find and that dance communication could be

69 detrimental if food sources are easy to find (see also Dechaume-Moncharmont et al. 2005). In
70 the latter case, foragers should search for new food sources through scouting (independent
71 search) and return to known high-quality food sources using route memory (Schürch and Grüter
72 2014).

73 There is a well-known, but not yet fully understood link between colony size and the
74 method of recruitment in ants (Beckers et al. 1989; Planqué et al. 2010; Dornhaus et al. 2012).
75 While large colony size is associated with pheromone-based mass-recruitment, species with
76 smaller colony sizes often forage solitarily or they use a recruitment method called tandem
77 running (Beckers et al. 1989). In tandem running, an experienced ant (tandem leader) guides an
78 inexperienced nestmate (tandem follower) to a new nest-site or a rewarding food source
79 (Hingston 1929; Wilson 1959; Möglich et al. 1974; Franks and Richardson 2006; Pratt 2008;
80 Kaur et al. 2017; reviewed in Franklin 2014). It has been argued that tandem followers locate
81 resources quicker than scouts that search for resources by individual exploration and trial-and-
82 error learning (Franks and Richardson 2006). Additionally, ants that are recruited by a tandem
83 leader might find food sources of higher quality because foragers are more likely to perform
84 tandem runs after finding a better food source (Shaffer et al. 2013). On the other hand, tandem
85 running also has disadvantages. During a tandem run, both ants walk with reduced speed
86 (Franks and Richardson 2006; Kaur et al. 2017) and a substantial proportion of tandem runs fail
87 (e.g. Wilson 1959; Pratt 2008; Glaser and Grüter 2018; Grüter et al. 2018). Furthermore,
88 recruits experience time and opportunity costs as they wait inside their nest for a leader, rather
89 than search in the environment for food sources by themselves. These disadvantages could
90 explain why some ant species do not seem to use tandem communication when foraging, even
91 though tandem runs are used during colony migrations (Hölldobler 1984; Traniello and
92 Hölldobler 1984; Fresneau 1985; Maschwitz et al. 1986). More generally, a sizeable group of
93 ant species do not seem to use any form of communication during foraging (e.g. Beckers et al.

94 1989; Lanan 2014; Reeves and Moreau 2019). This raises the question whether, when and how
95 a communication method that is relatively slow and small-scale, like tandem running, improves
96 colony foraging success and whether the ecological circumstances that favour tandem running
97 match those that favour honeybee dance communication.

98 We developed an agent-based simulation model to investigate the importance of
99 recruitment communication in the form of tandem running for the foraging success of virtual
100 ant colonies. We compared colonies that could perform tandem runs with colonies that
101 consisted only of scouts, *i.e.* without tandem running in an environment that varied in the
102 number, quality, distance and longevity of food sources. Additionally, we tested whether colony
103 size affects the importance of communication for colony foraging success. Finally, we explored
104 the role of forager ratio (relative numbers of scouts and recruits) and tested if recruits indeed
105 locate food sources faster than scouts. Based on studies that simulated honeybee foraging, we
106 predicted that tandem running is beneficial when high quality food sources are hard to find
107 (Dornhaus et al. 2006; Beekman and Lew 2008), but is detrimental to colony success when food
108 sources are short-lived (Schürch and Grüter 2014). We also predicted that larger colonies
109 benefit more from tandem running.

110

111 **The agent-based simulation model**

112 An agent-based simulation model (ABM) was developed using the software Netlogo 6.1.1
113 (Wilensky 1999, Wilensky and Rand 2015) (the NetLogo file can be found in the online
114 material). The model simulates the foragers of an artificial ant colony in an environment
115 consisting of their nest and food sources. Some of the basic parameters, like the range of colony
116 sizes, walking speeds or energy collected by foragers were derived from the ant species
117 *Temnothorax nylanderi* (Glaser and Grüter 2018).

118

119 *Purpose*

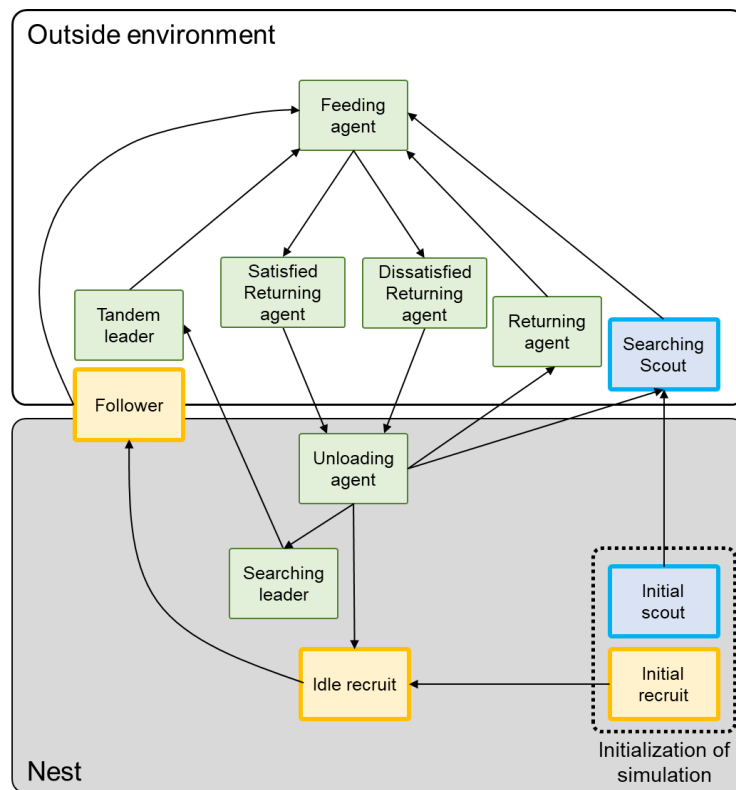
120 The aim of our model was to explore the adaptive value of tandem running in ants by measuring
121 the colony foraging success (as gained energy) and the time required by foragers to find a food
122 source. We compared colonies that could perform tandem runs with colonies that consisted only
123 of scouts, *i.e.* without tandem running. This latter situation is found in many ant species with
124 small colony sizes, such as *Diacamma* or *Neoponera* (Hölldobler 1984, Traniello and
125 Hölldobler 1984, Fresneau 1985, R. Kaur, pers. communication). In both situations, foragers
126 could also use route memory (or private information) to return to food sources they visited in
127 the past. We assessed the effects of communication depending on food source distribution
128 (number, distance), their quality and stability as well as colony size and the scout-recruit ratio.
129

130 *Entities, state variables and scale*

131 Netlogo operates with patches that can be used to measure distances and ticks for time steps. In
132 our model, 1 tick is equivalent to 1 second and 1 patch to 1 cm. The agents operate in a two-
133 dimensional square grid of 140×140 patches (arena) with a nest and either 2 or 10 food sources
134 (FS). This simulated environments with few or many food sources. The nest is located in the
135 center ($x=0, y=0$), with a radius of 10 patches. The food patches were at a distance of either 40
136 (default) or 20 patches from the outer edge of the nest, simulating natural conditions as *T.*
137 *nylanderi* mostly forages within 50 cm from their nest (Heinze et al. 1996). Each food source
138 had a size of 1 patch, which could represent a dead insect or a drop of honey dew, and could
139 either be of high or low quality (FS_{high} and FS_{low}), simulating a sugar solution of either 1 molar
140 or 0.1 molar concentration.

141 Since all agents are foragers, our default colony size of 100 would correspond to a
142 natural colony consisting of ~300-400 workers, assuming that foragers account for about 20-
143 30% of a *Temnothorax* colony (e.g. Shaffer et al. 2013). Simulated colonies consisted of varying

144 ratios of scouts that search for resources independently and recruits that waited in the nest until
 145 they are recruited to a food source. The default scout-recruit ratio was 1:4 (*i.e.* 20 scouts + 80
 146 recruits in the default situation), similar to what has been observed in honeybees where scouts
 147 represent about 5-35 % of the colony (von Frisch 1967; Seeley 1995). In colonies without
 148 tandem running, all foragers were scouts. In the default configuration, scouts and recruits can
 149 both assume any of the following seven states: (1) idle inside the nest, (2) feeding at food
 150 sources, (3) returning to the nest with food, (4) unloading food, (5) searching for a follower
 151 inside the nest, (6) leading a tandem run to the food source or (7) returning alone to the food
 152 source (*i.e.* use private information). Additionally, scouts search for food sources independently,
 153 while recruits wait inside the nest for a tandem leader. Recruits can then follow tandem runs to
 154 a food source (Fig. 1).



155
 156 **Fig. 1.** State diagram for the agent-based model for colonies with tandem runs. All foragers
 157 start inside the nest. Green boxes represent agent states that are possible for both scouts and

158 recruits, blue boxes are states that are only possible for scouts and orange boxes represent
 159 states that are only possible for recruits.

160

161 Colonies gain nest energy (NE) when agents return to the nest and “unload” the energy
 162 gained at food sources. We estimated the energy content of a crop load of a *T. nylanderii* forager
 163 the following way: we measured foragers (N = 21, from 3 colonies) and estimated that full
 164 foragers carry ~0.15 mg of sugar solution. Given the energetic value of sucrose, we calculated
 165 that a forager feeding at a 1 molar sucrose solution collects ~0.75 Joule per foraging trip,
 166 whereas a forager feeding at a 0.1 molar solution would obtain ~0.075 J.

167

168 **Table 1: Overview of the model parameters and the used values**

Variables	Description	Default values	Other values tested
FS distance	Distance from nest centre to food source	40	20
FS number	Number of food sources	2 or 10	
Colony size	Number of agents (foragers) in a colony	100	20-200
Scout-recruit ratio	The ratio of scouts and recruits in a colony	1:4 (r = 0.2) or (all scouts)	1:9 to 10:0 (r = 0.1 to 1.0)
FS _{High}	Energy gained from the high-quality food source	0.75 J	
FS _{Low}	Energy gained from the low-quality food source	0.075 J	
t_{max}	Duration of a simulation (1 tick ~ 1 second)	5400 ticks	21600 ticks
$v_{outside}$	Walking velocity of ants outside the nest	0.8 patch/ticks	
v_{nest}	Walking velocity of ants inside the nest	0.1 patch/ticks	
v_{tandem}	Walking velocity of Tandem leader and Tandem follower towards the respective food source	0.4 patch/ticks	
M_{cost}	Metabolic or energy cost of walking outside	$2.446 \times e^{-7}$ J/tick	$2.446 \times e^{-6}$, $2.446 \times e^{-8}$
t_{scouts}	Time a scout searches food before returning to nest	600-900 ticks	
$t_{nest-stay}$	Time a returned forager stays inside the nest	60 ticks	
$t_{tandemstarter}$	Time an active recruiter searches for a recruit inside the nest	120 ticks	
$t_{feeding}$	Feeding time of drinking agents	120 ticks	
$p_{break-up}$	Probability that tandems break up	0/tick	0.002/tick,

			0.005/tick
$p_{\text{recruitment}}$	Probability to recruit when “satisfied”	50%	

169

170

171 *Process overview*

172 The default simulation duration t_{max} was 5400 ticks (corresponding to 90 minutes), but we also
173 tested a duration of 21600 ticks (corresponding to 6 h). Time and distance in the model were
174 connected via the walking speeds (0.8 patches/tick outside the nest = v_{outside} , 0.4 patches/tick in
175 a tandem run = v_{tandem}), which were chosen to be similar to the walking velocity (in cm/sec) of
176 *T. nylander* ants (Glaser and Grüter 2018).

177 When the model was initialized ($t=0$), the nest and either 2 or 10 food sources and the
178 agents were created. In the situation without tandem running, only scouts were simulated. All
179 agents started in the centre of the nest. Scouts immediately started to perform a random walk to
180 search for food sources with the speed of v_{outside} , whereas recruits patrolled inside the nest with
181 speed v_{nest} (0.1 patches/tick) and waited to be recruited by another agent. All agents started with
182 an energy of zero. When leaving the nest, this energy decreases every tick by a metabolic cost
183 M_{cost} (see Table 1). M_{cost} was chosen so that the metabolic costs that accumulate during an
184 average foraging trip correspond to ~0.1% of the value of energy obtained during a typical
185 foraging trip (Fewell 1988). We estimated this by running several simulations and measuring
186 foraging trip duration of our agents. We also ran simulations with metabolic rates that were 10-
187 times higher or 10-times lower than our default value but found that this did not affect the
188 general patterns (Fig. S1).

189 When an agent finds a food source, it becomes a feeding agent and feeds for a duration
190 of 120 ticks. It gains either 0.75 J or 0.075 J, depending on whether the food source is of high
191 or low quality. If scouts do not find a food source within a certain time period (t_{scouts}), they
192 return to the nest. If they are at greater distances from the nest, unsuccessful scouts return

193 quicker (600 ticks). Unsuccessful scouts that are closer to the nest (35 patches from the center)
194 return if 900 ticks have passed. This was done to match observations that *T. nylanderi* scouts
195 often return to their nest if they had been searching unsuccessfully for several minutes (S.M.G.,
196 personal observation). After their return, unsuccessful scouts wait idle inside the nest for 60
197 ticks ($t_{\text{nest-stay}}$), before resuming to scout. At the end of the feeding time, agents return to the nest
198 either as “satisfied” or “unsatisfied” foragers. Foragers that found a high-quality food source
199 were always satisfied, whereas agents feeding at a low-quality food source had only a 10%
200 probability to become satisfied. After unloading for the duration of $t_{\text{nest-stay}}$, “satisfied” agents
201 become prospective tandem leaders with a 50% probability ($p_{\text{recruitment}}$), whereas unsatisfied
202 agents would not recruit. This leads to a recruitment probability of 5-50% per trip, which is
203 similar to what has been found in both *T. nylanderi* and *Pachycondyla harpax* (Glaser and
204 Grüter 2018; Grüter et al. 2018). Satisfied agents return to the same food source they had visited
205 before, either in a tandem run or alone. In other words, they use “route memory” to revisit a
206 high-quality food source, but were unlikely to return to a low-quality food source (10%
207 probability). Unsatisfied agents would not recruit and either wait inside the nest for a tandem
208 leader (recruits) or they search for a new food source (scouts). Fig. S2 is a screenshot of a
209 simulation showing the arrangement of the nest, food sources and some of the agent states.

210 Prospective tandem leaders stay inside the nest and search for a potential recruit for the
211 duration of 120 ticks ($t_{\text{tandemstarter}}$). A tandem run starts when a leader encounters a recruit on the
212 same patch. By default, tandem runs do not break up but we also tested situations with a break-
213 up probability of 0.002 and 0.005 per tick, which corresponds to tandem success rates of ~75 %
214 and ~50% for the default distance (calculated based on an average tandem run duration of 127
215 ticks for the default food source distance). Lost tandem followers first perform a random walk
216 for 180 ticks ($t_{\text{search-time}}$) and – if they do not find a food source – have an equal probability to
217 become either a scout or to return to the nest as an unsatisfied forager.

218 In the default settings, food sources were *ad libitum*, *i.e.* they did not disappear during
219 the simulations. Since this may not always be the case, we also simulated food sources that
220 disappeared after they were visited by 10 agents to create a more dynamic foraging environment.
221 If a food source disappears before ants return to it (either alone or in a tandem), agents reaching
222 the old food source location search randomly for 180 ticks ($t_{\text{search-time}}$), then they become
223 unsatisfied foragers and return to the nest. If the food source vanishes during feeding, the agent
224 becomes an unsatisfied forager. Food sources that have disappeared are replaced by an identical
225 food source at the same position after 600 ticks have passed, which means that it has to be
226 discovered again by scouts. For each simulation run, new inexperienced agents were created as
227 described above.

228

229 *Tested factors*

- 230 • Spatio-temporal distribution and variability of food sources: we tested the effects of food
231 source number (2 or 10), variability (only high-quality or variable quality. In the latter
232 situation, food sources alternated in quality, *i.e.* FS 1 high-quality, FS2 low-quality, FS3,
233 high-quality etc.), distance, foraging (simulation) duration and food source longevity (stable
234 or short-lived).
- 235 • Scout-recruit ratio: next to our default ratio of 1:4, we tested several other ratios, including
236 the extreme case with only scouts.
- 237 • Colony size: in addition to simulating a colony size of 100 agents, we tested a range of other
238 colony sizes (Table 1).
- 239 • Food discovery time of scouts and recruits: For each simulation run, we quantified the time
240 scouts needed to discover their first food source. In recruits, we measured both their waiting
241 time inside the nest and the duration of the tandem run. These durations were averaged per
242 forager type and per simulation run. Agents that did not discover a food source during an

243 entire simulation were given the maximum value of 5400 ticks. Only conditions with
244 successful tandem runs were considered.

245

246 The total nest energy NE (total J of all individual collection trips minus the total J of the
247 metabolic costs) were measured for each simulation run.

248

249 *Sensitivity of outcomes*

250 Due to the stochasticity of simulations we performed 30 simulation runs for each tested
251 combination of parameters. To assess how sensitive our model is to changes in the default
252 parameters, we varied many of them and explored their effect, as mentioned above (see Table
253 1). We also simulated environments with only low-quality food sources. A pure scouting
254 strategy was always better under these circumstances (Fig. S3). This is because tandem runs
255 are very rare when all food sources are of low quality and recruits spend most of their time
256 inside the nest.

257

258 *Statistical analyses*

259 All statistical analyses were performed using the software R 3.6.3 (www.r-project.org). Since
260 different treatments occasionally had unequal variance (heteroscedasticity) or contained zeros
261 and in order to provide a consistent statistical approach we used non-parametric statistical tests
262 throughout. It should be noted, however, that when we compared parametric and non-
263 parametric methods (Anova's), they yielded very similar results. We used Mann-Whitney U
264 tests to compare two independent samples and Wilcoxon signed-rank tests for paired data. In
265 addition to the p-values, the R software provides the test statistic value W, which is a linear
266 transformation of the usual rank sum statistic U. When three groups were compared, we used

267 Kruskal-Wallis tests and Dunn tests with sequential Bonferroni corrections for post-hoc pair-
268 wise comparisons (“FSA” package, Ogle et al. 2020) (Sokal and Rohlf 1995).

269

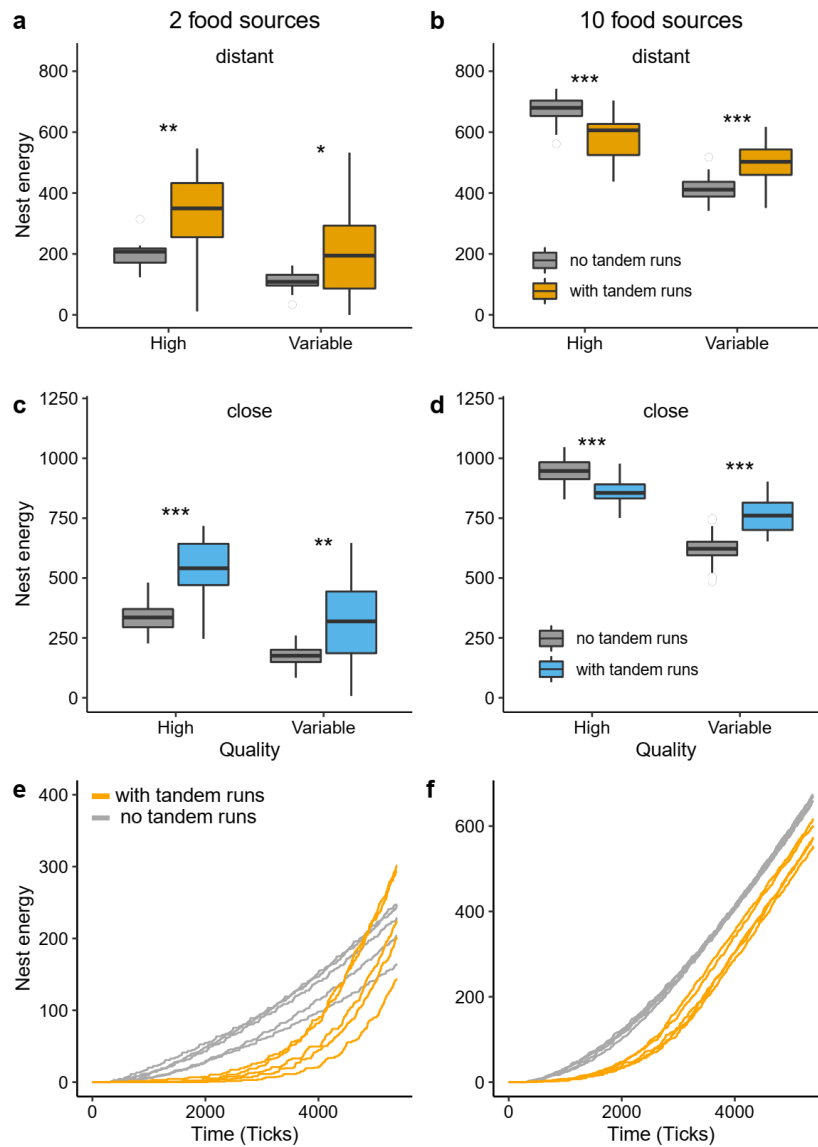
270 **Results**

271 *Distribution of food sources*

272 We first tested if the number of food sources and their distance from the nest affect the value
273 of communication. When colonies had access to few food sources, they were more successful
274 with tandem recruitment (scout-recruit ratio of 1:4) than colonies consisting only of scouts,
275 irrespective of whether food sources were of high-quality (Fig. 2) (Mann-Whitney U Test, W
276 = 215, $p = 0.0004$) or of variable quality ($W = 307$, $p = 0.034$). In a rich environment, with 10
277 food sources, colonies collected overall more energy (Fig. 2). Tandem communication was
278 beneficial when food source quality was variable ($W = 112$, $p < 0.0001$), whereas colonies
279 consisting only of scouts performed better when all 10 food sources were of high quality ($W =$
280 773 , $p < 0.0001$). This general pattern did not change when food sources were closer to the nest
281 (20 patches instead of 40 patches) (2 food sources, high-quality: $p < 0.0001$; variable-quality:
282 $W = 210$, $p = 0.0003$; 10 food sources, high-quality: $W = 827$, $p < 0.0001$; variable-quality: W
283 = 54, $p < 0.0001$), but colonies gained overall more energy when all food sources were close to
284 the nest (Fig. 2). Fig. 2e and 2f illustrate the temporal development of nest energy during
285 exemplary simulation runs that correspond to the conditions shown in Fig. 2a and 2b with high-
286 quality food sources.

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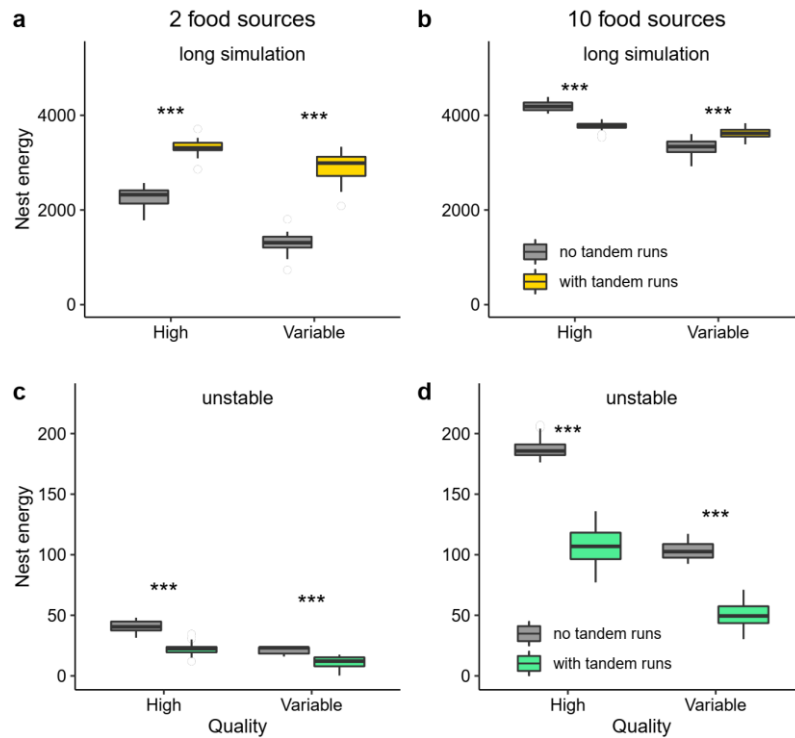
290 **Fig. 2.** Nest energy with 2 or 10 food sources. In (a) and (b) food sources were at a distance of 40
 291 patches, whereas in (c) and (d) food sources were at a distance of 20 patches. *p<0.05, **p<0.001,
 292 ***p<0.0001. In (e) and (f), nest energy is plotted over time for conditions as shown in (a) and (b) when
 293 all food sources were of high quality (5 simulation runs per treatment for visualisation of the trajectory).
 294

295

296 *Foraging duration and food source longevity*

297 When we increased the foraging duration (*i.e.* the simulation duration) from 5400 to 21600
 298 ticks, we found a similar pattern. Tandem running was highly beneficial when there were few
 299 food sources (high-quality: $W = 0$, $p < 0.0001$; variable-quality: $W = 0$, $p < 0.0001$). Tandem
 300 runs were also beneficial when there were many food sources of variable quality ($W = 56$, $p <$
 301 0.0001). In the case of many high-quality food sources, pure scout colonies performed better

302 (W = 900, $p < 0.0001$). It is noteworthy that colonies with tandem communication were almost
303 as successful in an environment with 2 food sources as in an environment with 10 food sources
304 (Fig. 3a,b).
305



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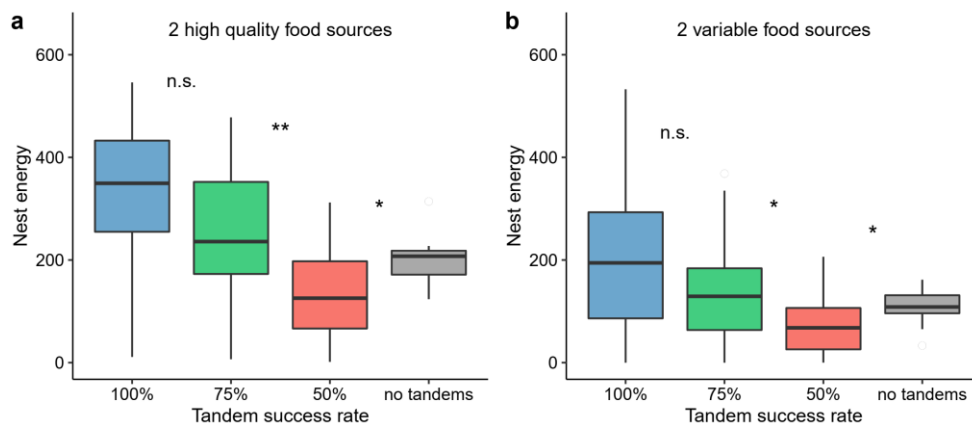
307 **Fig. 3.** Nest energy with 2 or 10 food sources. In (a) and (b), food sources simulations were 4-times
308 longer (21600 ticks instead of 5400). In (c) and (d), simulations lasted 5400 ticks and food sources
309 disappeared if they were visited by 10 ants. A new food source appeared after a delay.

310 So far, we assumed that food sources offered food during the entire simulation. Next,
311 we tested the effects of short-lived food sources. If food sources were unstable (sometimes
312 called non-renewable), a scouting strategy was more successful, irrespective of the number of
313 food sources and their variability (Fig. 3c,d) (2 food sources, high-quality: W = 897, p-value <
314 0.0001; variable-quality: W = 896, p-value < 0.0001; 10 food sources, high-quality: W = 900,
315 p-value < 0.0001; variable-quality: W = 900, p-value < 0.0001). Differences were particularly
316 pronounced when colonies were offered many food sources. Scouting remained the better
317 strategy when we increased the foraging duration to 21600 ticks (e.g. 2 food sources, high-
318 quality: W = 900, p-value < 0.0001; variable-quality: W = 900, p-value < 0.0001).

319

320 *Tandem success rate*

321 Tandems do occasionally break up and we tested how this affects the energy collected by
322 colonies. We compared colonies with 100% (default), ~75% and ~50% successful tandem runs
323 and colonies with only scouts in an environment with few food sources, *i.e.* under conditions
324 where tandem runs are beneficial (Fig. 2a). Our simulations show that a reduction in tandem
325 success rate has a negative impact on the energy intake that is collected by colonies (Fig. 4). If
326 only about 50% of the tandem runs are successful, colonies without any tandem running collect
327 more energy in an environment with few, stable food sources (Fig. 4) (high-quality, 50%
328 success rate vs. no tandems: $W = 246$, $p = 0.002$, variable-quality, 50% success rate vs. no
329 tandems: $W = 257$, $p = 0.004$).



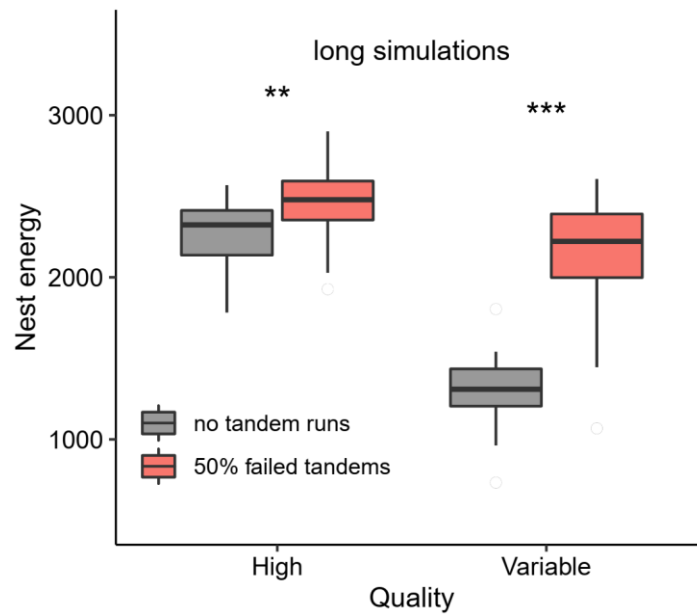
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331 **Fig. 4.** Nest energy with 2 food sources of high (a) and mixed (b) quality in relation to the tandem success
332 rate. Adjacent treatment groups were compared, as indicated by asterisks or “n.s.”. No tandems = only
333 scouts. Default settings were used for the other parameters.

334

335 However, tandem runs with a high rate of failure (50%) are not always a disadvantage
336 compared to having no communication. When colonies can forage for longer (simulations of
337 21600 ticks), colonies that perform tandem runs with a ~50% break-up rate are more successful
338 than colonies consisting of only scouts (Fig. 5) (high-quality, 50% success rate vs. no tandems:
339 $W = 866$ $p < 0.0001$, variable-quality, 50% success rate vs. no tandems: $W = 689$, $p = 0.0003$),
340 highlighting the benefits of imperfect communication over longer time periods.

341



342

343 **Fig. 5.** Nest energy with 2 food sources of high or variable quality and a long foraging duration. Colonies
 344 were scouting or could recruit with tandem runs that had a ~50% failure rate. Default settings were used
 345 for the other parameters.

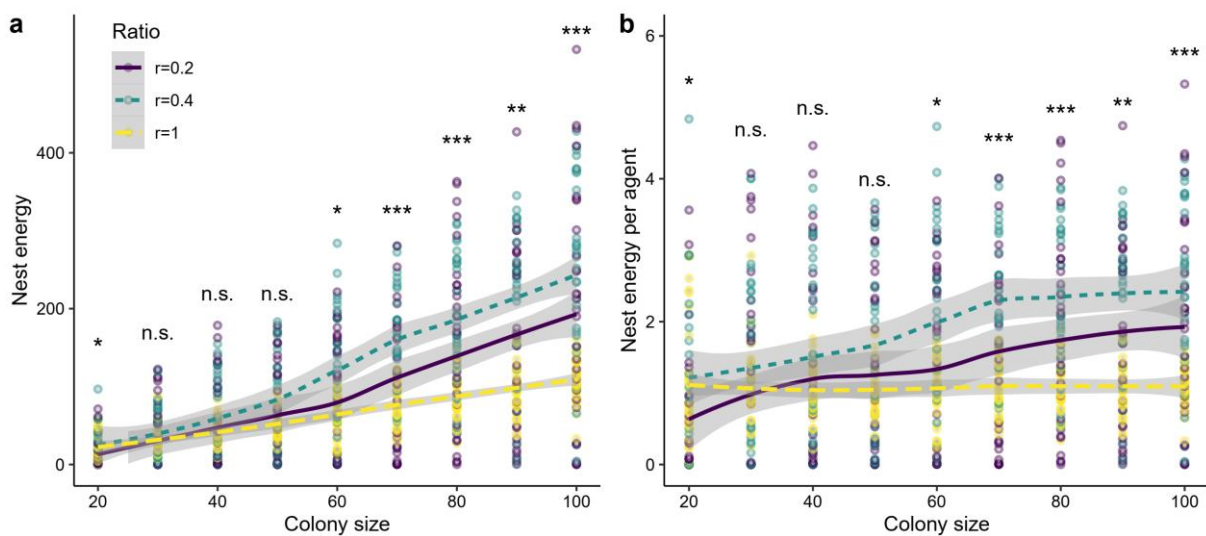
346

347 *Colony size and scout-recruit ratio*

348 We tested various colony sizes ranging from 20 to 200 agents in an environment with few,

349 variable food sources, *i.e.* an environment that favours tandem running under default conditions

350 (see Fig. 2a).



351

352 **Fig. 6.** The relationship between colony size and nest energy (a) and nest energy per agent (b) in
 353 colonies with and without tandem runs in environments with two food sources of variable quality. Three

354 scout-recruit ratios were simulated, $r = 0.2$ and $r = 0.4$ and colonies consisting only of scouts, $r = 1.0$.
 355 Grey area indicates confidence intervals. Significance tests refer to comparisons among ratios,
 356 separately for each colony size. P-values for total nest energy (a) or energy per agent (b) are identical.
 357 Default settings were used for the other parameters.

358

359 Colony size had a strong effect on the total collected energy that was collected (Fig. 6).

360 If colonies were very small (20 foragers), they were least successful if they performed tandem

361 runs and had a default scout-recruit ratio of 0.2 (Table 2). There was no difference in foraging

362 success when colony size ranged from 30 to 50 foragers. However, colonies with tandem

363 recruitment were more successful if they had at least 60 agents (Table 2). The most successful

364 colonies contained 40% scouts, suggesting that the scout-recruit ratio has a considerable impact

365 on colony success. Fig. 6b shows the nest energy collected per agent (nest energy/colony size).

366 In colonies with only scouts, individual agents collected a relatively constant amount of energy

367 irrespective of colony size (Spearman rank correlation: $\rho = 0.1$, $p = 0.09$). In colonies with

368 tandem running, on the other hand, individual agents collected more energy on average as

369 colony size increased from 20 to 100 agents ($r = 0.2$, $\rho = 0.34$, $p < 0.0001$; $r = 0.4$, $\rho = 0.35$,

370 $p < 0.0001$).

371

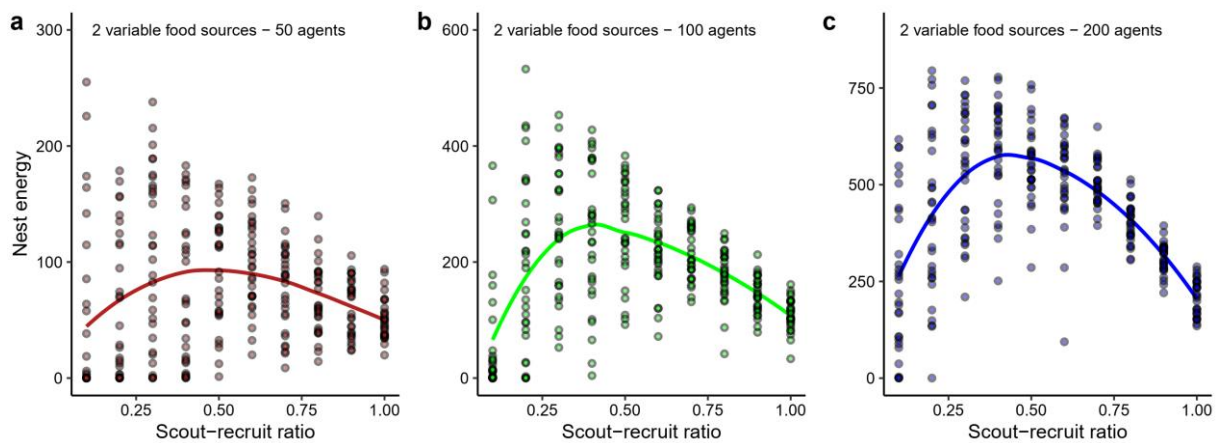
372 **Table 2:** Effect of colony size on nest energy. Three conditions were tested: in two conditions, colonies
 373 performed tandem runs and had a scout-recruit ratio of 0.2 or 0.4. In the third condition, colonies
 374 consisted only of scouts (1.0). Pair-wise comparisons were performed if the overall $p < 0.05$ and p-
 375 values were corrected using sequential Bonferroni.

376

Colony size	Kruskal-Wallis Test		p-value of pair-wise comparisons		
	χ^2	p-value	0.2 vs. 0.4	0.2 vs. 1.0	0.4 vs. 1.0
20	12.7	0.002	0.018	0.002	0.42
30	2.33	0.31	NA	NA	NA
40	3.92	0.14	NA	NA	NA
50	1.81	0.41	NA	NA	NA
60	8.98	0.01	0.077	0.4	0.01
70	22.85	<0.0001	0.0007	0.35	<0.0001
80	20.6	<0.0001	0.13	0.007	<0.0001
90	15.86	0.0004	0.09	0.048	0.0002
100	19.17	<0.0001	0.053	0.027	<0.0001
200	46.59	<0.0001	0.006	0.0001	<0.0001

377

378 To explore this further, we simulated different scout-recruit ratios and different colony
379 sizes to test how the balance between scouts and recruits affects colony foraging success.
380 Simulations suggest that the optimal proportion of scouts is ~40% for the simulated
381 environment, irrespective of colony size (Fig. 7). Interestingly, deviations from the optimal
382 ratio have a larger negative impact in larger colonies (see “pointiness” of curves in Fig. 7). For
383 example, there is no difference in success when colonies with 50 agents contain 40% or 80%
384 of scouts ($W = 119$; $p = 0.54$). When colony size is 200, however, colonies with 80% scouts
385 collect 31.5% less energy than colonies with 40% scouts ($W = 199$, $p = 0.0001$).
386



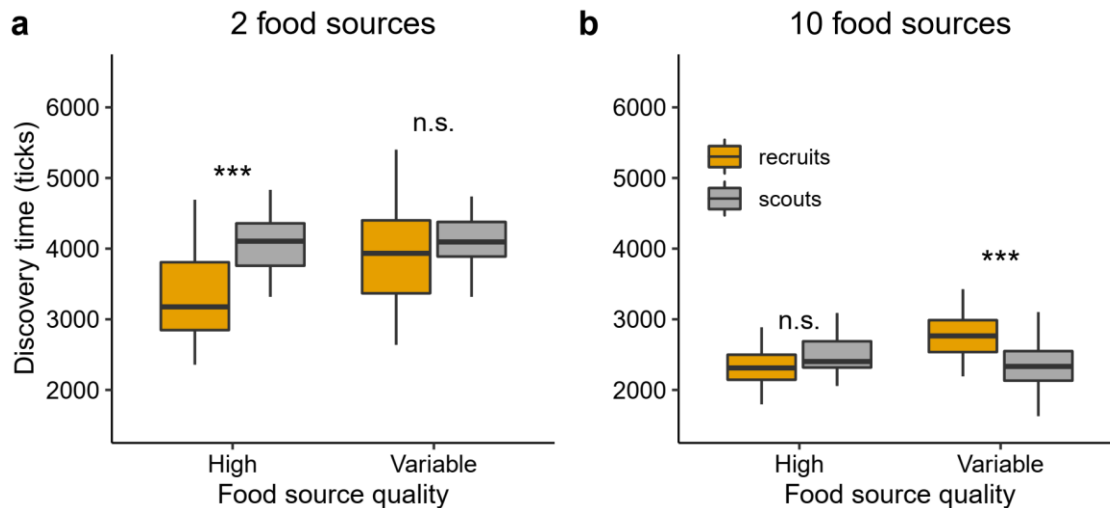
387
388 **Fig. 7.** The effect of the scout-recruit ratio with three different colony sizes. The line shows the best fit
389 line based on local polynomial regression using the LOESS method (locally estimated scatterplot
390 smoothing). The smallest ratio was 0.1. A ratio of 1.0 refers to colonies containing only scouts.
391
392

393 *Discovery times*

394 Unsurprisingly, foragers needed more time to find their first food source in an environment
395 with few food sources compared to when there were many food sources (Fig. 8). Recruits
396 needed less time in an environment with few, high-quality food sources compared to scouts
397 (Wilcoxon-signed rank test: $W = 143$, p -value < 0.0001), whereas there was no difference when
398 food sources were variable in quality (Fig. 8a) ($W = 348$, p -value = 0.13). However, in an
399 environment with many food sources, scouts did comparatively better and needed a similar

400 amount of time to locate their first food source when food sources were all high-quality ($W =$
401 327, p -value = 0.07). With many, variable food sources, scouts were significantly faster than
402 recruits (Fig. 8b) ($W = 720$, p -value < 0.0001).

403



404

405 **Fig. 8.** Time until agents located their first food source in environments with few (a) or many (b) food
406 sources of constant or variable quality. For recruits, the food discovery time consisted of the time waiting
407 inside the nest and the tandem run duration. Default settings were used for the other parameters.

408

409

410 Discussion

411 Our simulations show that the spatio-temporal distribution of food sources greatly affects
412 whether colonies with communication are more successful than colonies that employ a scouting
413 strategy. Tandem running was beneficial when colonies were in an environment with few food
414 sources (+ 57-83% nest energy) and when food sources were of variable quality (Fig. 2a,b).
415 Colonies without communication were more successful (~15%) in a rich environment that
416 offered only high-quality food sources. This is in line with studies that simulated honeybee
417 foraging and found that communicating food source locations by waggle dancing is most
418 beneficial if food sources are hard to find and of variable quality (Dornhaus et al. 2006;
419 Beekman and Lew 2008; Schürch and Grüter 2014; I'Anson Price et al. 2019). Under such

420 circumstances, the probability that scouts find high-quality food sources on their own is low
421 and communicating the location of a relatively small number of high-quality patches becomes
422 advantageous. As food source variability decreases and the number of high-quality food sources
423 increases, scouts become more successful. Even though colonies with communication also
424 collect more energy in such an environment, the benefits of communication no longer offset the
425 costs of recruits waiting for information inside the nest. This highlights that communication
426 often has considerable time and opportunity costs (Seeley 1983; Seeley and Visscher 1988;
427 Dechaume-Moncharmont et al. 2005; Schürch and Grüter 2014; I'Anson Price et al. 2019).

428 It has been hypothesised that recruitment communication is particularly beneficial in an
429 ephemeral environment (Sherman and Visscher 2002; Dornhaus and Chittka 2004; Grüter and
430 Ratnieks 2011), *i.e.* when food sources last only for short time-periods and, thus, need to be
431 exploited quickly. Counterintuitively, a simulation model of honeybee foraging has found that
432 communication was less beneficial if food sources were shorter-lived (Schürch and Grüter
433 2014). Our simulations support their findings by showing that tandem running was a very
434 successful strategy in a stable environment with relatively long foraging durations (*i.e.* with
435 longer simulations) and few, variable food sources (Fig. 3a). A long-lasting food source could
436 be a large insect (Lanan 2014), floral nectars or a group of honeydew secreting insects (Carroll
437 and Janzen 1973; Quinet and Pasteels 1996; Völkl et al. 1999; Mailleux et al. 2003; Lanan
438 2014). A very different pattern was observed when resources were shorter-lived: colonies
439 without communication were always more successful, irrespective of the foraging (simulation)
440 duration (Fig. 3c,d). The most likely explanation is that colonies with communication pay time
441 costs without being able to take advantage of the benefits of communication over longer time
442 periods (see also Schürch and Grüter 2014). Our model differs from theirs in that our food
443 sources only disappeared if they were exploited, rather than with a constant probability. A food
444 source that disappears after it has been exploited could be a droplet of honeydew that fell on

445 vegetation. Honeydew droplets on leaf surfaces represent an important food source for the
446 tandem recruiting *Temnothorax curvispinosus* (Lynch et al. 1988).

447 Tandem runs occasionally break-up and success rates of ~50% to 90% are not
448 uncommon (Wilson 1959; Pratt 2008; Kaur et al. 2017; Glaser and Grüter 2018; Grüter et al.
449 2018). We simulated different success rates and found that colonies with more successful
450 tandem runs collected more energy (Fig 4). If the success rate was about 50%, colonies
451 consisting only of scouts collected more energy in an environment with few food sources, *i.e.*
452 a virtual environment that normally favours tandem running. When foraging durations were
453 longer, on the other hand, colonies with tandem runs gained the upper hand over scouting
454 colonies even though half of all tandem runs failed (Fig. 5). Under these circumstances, even a
455 relatively low number of successful recruitment events can be very important because the
456 discovered high-quality food sources can be exploited for longer time periods by successful
457 recruits. Additionally, tandem recruitment can lead to an exponential increase of ants at a feeder
458 even if a leader recruits <1 follower per trip. With exponential growth, the impact of
459 communication will increase over time (Fig. 2e).

460 We found that colony size had a considerable effect on the value of tandem
461 communication (Fig. 6). This contrasts with models of honeybee communication, where colony
462 size did not greatly affect the benefits of communication (Dornhaus et al. 2006; Schürch and
463 Grüter 2014), but is consistent with an empirical study on honeybee colony foraging success
464 (Donaldson-Matasci et al. 2013) and a mathematical model of ant communication (Planqué et
465 al. 2010). If colonies contained 60 or more foragers, tandem communication was usually
466 beneficial. However, a pure scouting strategy was equally or more successful when colonies
467 had 20 to 50 foragers, even in environments with few and variable food sources, *i.e.* a virtual
468 environment that normally favours tandem running. This number of foragers could be expected
469 in ant colonies with ~80-250 workers (assuming that foragers make up 20-30% of the worker

470 population, e.g. Shaffer et al. 2013), which is also the typical colony size of many ant species
471 that use tandem running and species with solitary foraging (Beckers et al. 1989). Our simulation
472 results could explain why some species, e.g. in the genera *Diacamma* or *Neoponera*, do not
473 perform tandem runs during foraging even though they use this recruitment method during
474 migrations (Hölldobler 1984; Traniello and Hölldobler 1984; Maschwitz et al. 1986). Whether
475 colonies employ tandem running might depend on the food sources they collect (e.g. small or
476 large items) and whether they are risk-averse or risk-prone because tandem recruitment was
477 often associated with a more unpredictable outcome in our simulations (greater variation in nest
478 energy gain among simulations of a particular situation, see Fig. 2). A better understanding of
479 the natural history of these species and similar species that do perform tandem runs (e.g.
480 *Neoponera* vs. *Pachycondyla*) is needed to understand why some species use communication,
481 while others forage solitarily.

482 Colony foraging performance depended on the proportions of scouts and recruits (Fig.
483 7). In our simulations with few food sources, colonies were most successful if scouts
484 represented about 40% of the forager population, but this is likely to depend on the number and
485 variability of food sources (see Fig. 2). Interestingly, having the right scout-recruit ratio is more
486 important in larger colonies than in smaller ones, possibly because the foraging success of
487 smaller colonies depends more on chance events, such as the discovery of a high-quality food
488 source by a single scout. This suggests that larger colonies would benefit from having the ability
489 to assess their current environment and adjust their use of communication accordingly. Whether
490 this is indeed common is not well known, but it has recently been reported that honeybees are
491 able to assess the value of communication and reduce their reliance on waggle dances if dance
492 information is not beneficial in the current environment (I'Anson Price et al. 2019).

493 In the simulations, we measured the time recruits and scouts need to locate their first
494 food source in environments with many or few food sources. We found that the food discovery

495 time depends strongly on the environment. Recruits were faster in environments with few high-
496 quality food sources, whereas scouts found a food source sooner in an environment with many,
497 variable food sources. Our measurements also included the time that recruits wait inside the
498 nest to find a tandem leader. Franks and Richardson (2006) found that tandem followers found
499 a food source faster in their experiment with one food source, which, in combination with their
500 other findings, indicated that tandem running fulfils the criteria for animal teaching set out by
501 Caro and Hauser (1992; namely, a teacher [*i*] modifies its behaviour in the presence of a naive
502 observer, [*ii*] at some cost to the leader [*iii*] so that the observer can learn more quickly or
503 efficiently). Our simulations suggest that this is the case only in certain environments, namely
504 those with few, high-quality resources. In other situations, scouts are likely to learn food source
505 locations quicker and tandem running might no longer fulfil the criteria for animal teaching
506 (namely that a follower acquires knowledge or learns a skill more rapidly or efficiently than it
507 might otherwise do, or that it would not learn at all, see Caro and Hauser 1992).

508 Taken together, our simulations show that the value of tandem communication is highly
509 dependent on the environment and the size and composition of the colony. Future studies should
510 explore whether and how foragers can assess their foraging environment and modify their
511 communication behaviour (see also Grüter and Czaczkcs 2019). It would also be desirable to
512 test the conclusions from our simulations empirically, but so far it has been challenging to stop
513 ants from performing tandem runs without affecting their behaviour.

514

515 **Acknowledgments**

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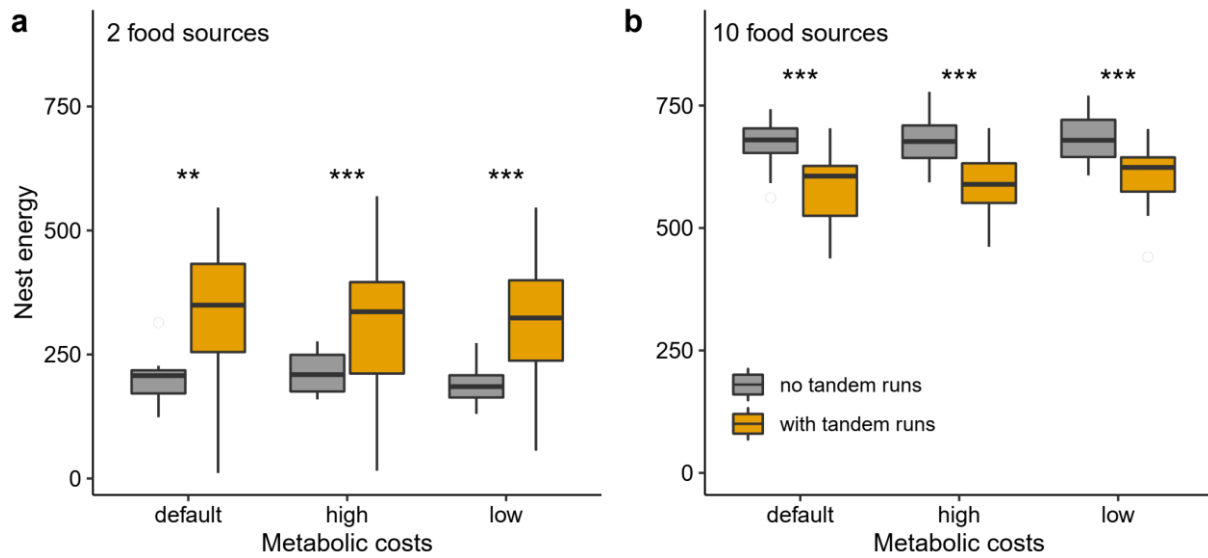
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631 **Fig. S1**



632

633 **Fig. S1.** The effects of 10-times higher and 10-times lower metabolic costs on nest energy
634 (see Table 1). Two (a) and ten (b) high-quality food sources were offered, default values were
635 used for all other parameters. The default conditions match those shown in Fig. 2a and 2b.
636 Mann-Whitney U tests, ** $p < 0.001$, *** $p < 0.0001$.

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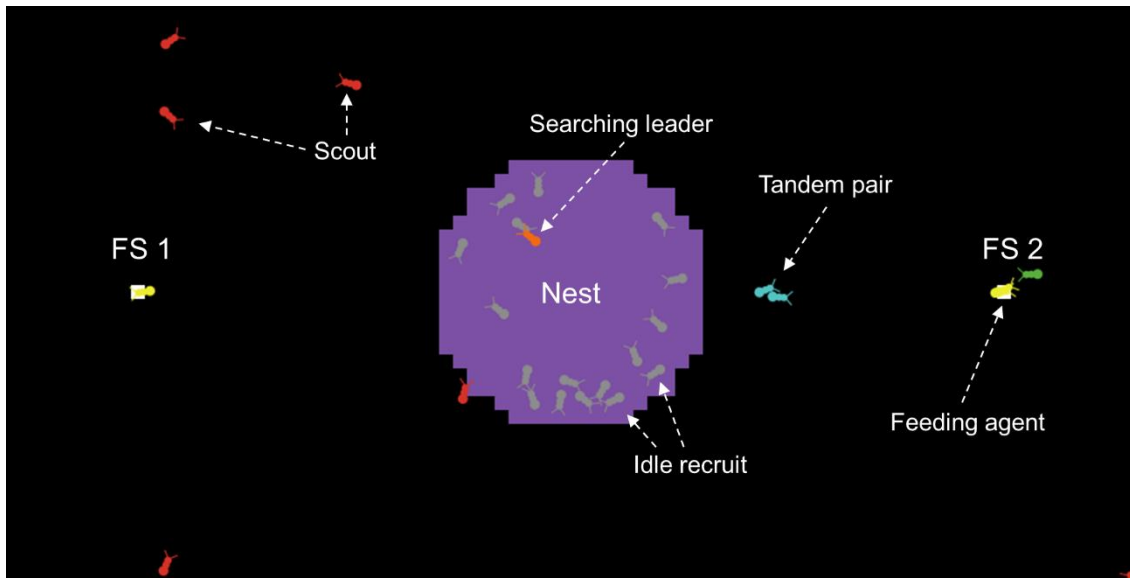
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647 **Fig. S2**



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649 **Fig. S2.** NetLogo interface showing some of the different agent types in different colours. In
650 this situation, two food sources (FS 1 and FS 2) were offered.

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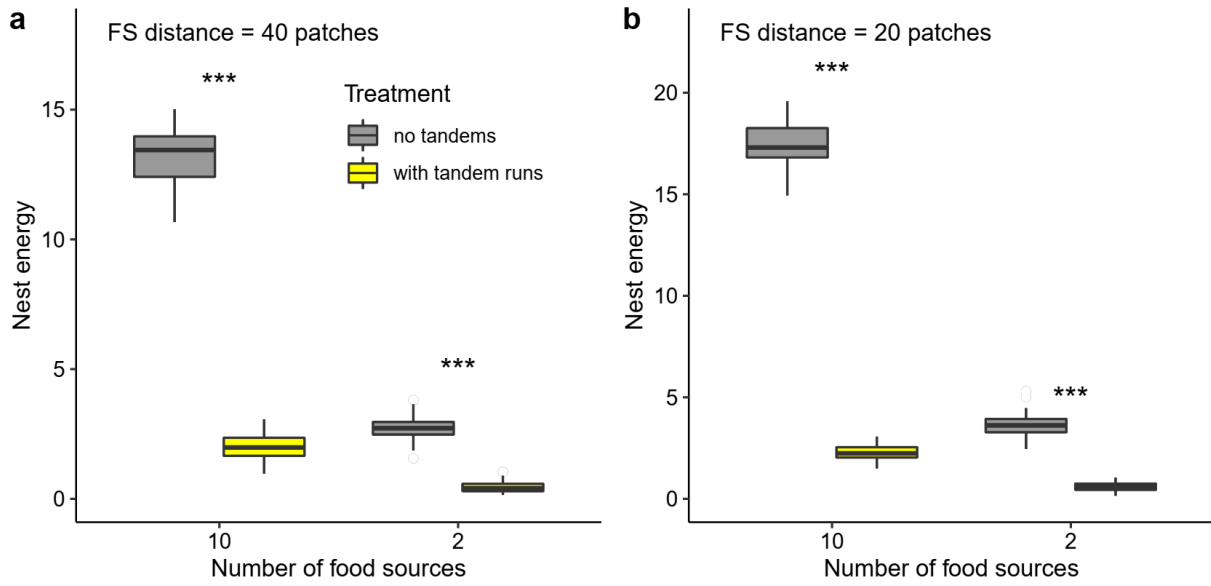
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662 **Fig. S3**



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664 **Fig. S3.** Nest energy of colonies with or without tandem runs when all food sources are of low
665 quality. Two food source distances were simulated, 40 patches (a) or 20 patches (b). Default
666 values were used for all other parameters. Mann-Whitney U tests, *** $p < 0.0001$.

667