Error in the Honeybee Waggle Dance Improves Foraging Flexibility

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The honeybee waggle dance communicates the location of profitable food sources, usually with a certain degree of error in the directional information ranging from 10°–15° at the lower margin. We simulated one-day colonial foraging to address the biological significance of information error in the waggle dance. When the error was 30° or larger, the waggle dance was not beneficial. If the error was 15°, the waggle dance was beneficial when the food sources were scarce. When the error was 10° or smaller, the waggle dance was beneficial under all the conditions tested. Our simulation also showed that precise information (0°–5° error) yielded great success in finding feeders, but also caused failures at finding new feeders, i.e., a high-risk high-return strategy. The observation that actual bees perform the waggle dance with an error of 10°–15° might reflect, at least in part, the maintenance of a successful yet risky foraging trade-off.

Results

Measurement of parameters and modeling. We categorized the foraging-related behavior of an individual bee into eight behavioral states to create the iMoAD-f (Fig. 1c). Transitions between the behavioral states were
determined by the probabilities discussed in the literature\(^{25}\) (Table 1) and by state-specific rules (see Supplementary Methods). Importantly, probabilities related to dancing and following depended on the concentration of sugar solution in the feeder (Table 2), based on previous observations\(^{20,26,27}\).

For parameterization of the simulations, we first videotaped the bee behavior, and then extracted and measured the parameters (Supplementary Fig. S1 and Supplementary Table S1). We found that dancing was performed in a restricted area, the so-called dance floor (Supplementary Fig. S1a). About 85% of followers (476 follows from 414 followers) turned away from their dancer after one or two waggle runs (Supplementary Fig. S1b). Walking bees also tended to be found on the dance floor (Supplementary Fig. S1c). Walkers moved forward but not far, i.e., at most 2.0 cm/s (Supplementary Fig. S1d,e,f), and changed their body axis between \(-40^\circ\) and \(40^\circ\) (about 80% of 1933 1-sec walks) (Supplementary Fig. S1g). The body weights of bees with a crop full of nectar and bees with an empty crop (mean \(\pm SE\)) were 119 \(\pm\) 2.75 mg (\(n = 10\)) and 78.5 \(\pm\) 1.62 mg (\(n = 14\)), respectively. The bees with full crops were likely to be successful foragers, and those with empty crops to be failed foragers. About 85% (304 out of 358 runs) of directional information variation ranged within \(15^\circ\), as we previously observed\(^{9,10}\). In addition to our own present and previous results, results from the literature were used to run the simulation (Supplementary Table S1). Most importantly, only measurable parameters were used, and thus our simulation results might be evaluated in future biological experiments.

**Evaluation of the model.** We evaluated iMoAD-f by comparison with a well-known experiment by Seeley et al.\(^{20}\), which is often used as a framework for modeling honeybee foraging\(^{28-32}\). In

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**Table 1 | Transition probabilities between two behavioral states**

| next behavioral state | Res | Wan | Sco | For | Suc | Fai | Dan | Fol |
|-----------------------|-----|-----|-----|-----|-----|-----|-----|-----|
| Res                   | 0.79| 0.21|     |     |     |     |     |     |
| Wan                   | 0.44| 0.5597| 0.0003\(^{a}\) |     | 0.0003\(^{a}\) |     |     |     |
| Sco                   | 0.0003\(^{a}\) |     | 0.9975| 1.0\(^{b}\) | 0.0025, 1.0\(^{b}\) |     |     |     |
| For                   |     |     |     | 1.0\(^{b}\) | 1.0\(^{b}\) |     |     |     |
| Suc                   |     | variable |     | variable | 1.0\(^{b}\) |     |     | variable |
| Fai                   |     |     |     |     |     |     |     | 1.0\(^{b}\) |
| Dan                   |     | 0.5 |     | 0.5 |     |     |     | 1.0\(^{b}\) |
| Fol                   |     |     |     |     |     |     |     | 0.6 |

For details, see Supplemental Methods.

\(^{a}\) The probability was dependent on the bee’s experience.

\(^{b}\) The probability was 1.0 if the specific condition was met (for details, see Supplementary Methods), and otherwise was 0. Variable: See Table 2. Res, resting; Wan, wandering; Sco, scouting; For, foraging; Suc, success; Fai, failure; Dan, dancing; Fol, following.

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**Figure 1 | The waggle dance and our model.** (a), Variation of directional information encoded by waggle runs. Each run had a certain degree of variation from the mean waggle-run direction. (b), Sequences of information variation in consecutive waggle runs of two dancers (Bee 11 and Bee 12). The mean direction is represented as 0\(^\circ\). (c), iMoAD-f. Eight behavioral states are indicated and transitioned by a specific probability and state-specific rules. For visualization, scouting (SCO) and foraging (FOR) are represented in the same box, because these two states were essentially the same, except that a bee could be flying in search of a feeder (i.e., lacking directional information about a feeder), or it could be flying to a known feeder (i.e., with directional information about the feeder). The numbers beside the solid lines indicate the transition probabilities (see Tables 1 and 2). Dotted lines indicate that the states automatically transitioned, i.e., the transition probability was 1, under state-specific rules (see Supplementary Methods). The probability of scouting from success (Vd(c)), of dancing from success (Vd(c)), and of foraging from following (f(c)) depended on the concentration of feeders (see Table 2). DAN, dancing; FAI, failure; FOL, following; FOR, foraging; RES, resting; SCO, scouting; SUC, success; WAN, wandering.
Seeley’s experiment, two feeders were placed 400 m away from the hive to the south and the north, respectively. The concentrations of sugar water in the south and north feeders were 2.5 M and 1.0 M, respectively. After 4 hours, the concentrations were changed to 0.75 M and 2.5 M, respectively. They found that the colony altered its feeder visitation habits in response to the change in the concentration of sugar water. We reproduced Seeley’s experiment, and 20 simulations yielded similar results (Supplementary Fig. S2). Thus, we concluded that iMoAD-f reliably expressed the behavior of an actual bee colony.

**Availability of the waggle dance.** Do waggle dances really confer an advantage for food collection? To examine the first function, we compared the foraging results among four virtual colonies characterized by their foraging strategies: a random-search colony, a non-communication colony, a random-information dance colony, and a normal waggle dance colony (see Methods). Twenty simulations of one-day foraging by a 1000-bee colony under experimental conditions mimicking those of Seeley et al. revealed a large difference in the total number of visits to all feeders for the entire foraging time among colonies (Fig. 2a). For the random-search colony, no-communication, random-information dance and waggle dance colonies, the mean total numbers of visits, with standard deviations, were 409.5 ± 14.1, 1348.1 ± 75.5, 1978.0 ± 292.3, and 2756.4 ± 237.0, respectively. The analysis of variance (ANOVA) showed a significant effect of colony type \( (P < 0.001) \). Among the colonies tested, the waggle dance colony made a significantly greater number of successful visits to food sources than the other three colonies (Fig. 2a, \( P < 0.001 \) for all combinations, Tukey-Kramer test).

For evaluating foraging success, the energy-balance results are more general and useful than the number of visits, because a gain in energy will enhance the survivability of the colony. We calculated the energy balance by subtracting the energy loss from the energy gain for one-day foraging using the values in Figure 2a and found that the waggle dance colony gave the best foraging performance, as expected (Fig. 2b). Interestingly, there were no longer significant differences in energy balance between the no-communication and random-information dance colonies under the present conditions (Fig. 2b). Another 100 simulations under the same conditions as used in Figure 2a, i.e., 2 feeders at 400 m, demonstrated that the rates of success expressed as a percentage of total visits of the two colonies were almost identical (38.6% for the no-communication colony and 38.5% for the random-information dance colony), i.e., the efficiency was the same (Supplementary Fig. S3).

When the number of feeders was increased from 2 to 10, the waggle dance colony showed the greatest energy gain, but this result did not differ significantly from that for the random-information dance colony (Fig. 2c), indicating that both the informative and the non-informative dances performed best under this feeder condition. These findings suggest that the best foraging strategy is dependent on the number of feeders.

**Optimal conditions for dance execution.** We then examined the relationship between the spatial distribution of feeders and the efficiency of the waggle dance as assessed by energy balance, taken together with variations in directional information. iMoAD-f revealed that either informative or non-informative dancing behavior was beneficial in 139 (82.7%) of 168 tested conditions (Fig. 2d), suggesting that interactions between individuals per se are important for collective foraging, irrespective of the information exchanged. In addition, random searching may not be strategic, as no best-cases were found in this group (Fig. 2d).

As shown in Figure 2c, the informative dance (waggle dance colony) was not always beneficial (Fig. 2d). When the variation was 5° or less, only the waggle dance colony was the best colony (colonies with a performance not significantly different from the best colony were also classified as best colonies; see Methods) under all conditions tested (Fig. 2d). When the variation was 10°, the waggle dance colony remained the best performer, and there were cases in which the random-information dance colony and/or the no-communication colony performed equally well (Fig. 2d). Under this variation (10°), only the waggle dance colony was the best colony if there were a small number of feeders, which suggested that the feeders were hard for the bees to locate. In contrast, when the variation was large (30° or 60°), the waggle dance colony did not perform the best. Interestingly, when the variation was 15°, the results were highly mosaic (Fig. 2d). Basically, when the feeders were not easy to find due to the small number of feeders, the waggle dance colony performed the best, and the random-information dance colony performed the best when the feeders were easy to find. The no-communication colony performed the best when the feeders were far away (2000 m). If the bees persisted with informative dancing when the feeders were at this latter distance, the variation had to be 10° or less (Fig. 2d). Otherwise the colony could not expect an advantage from dancing. Consistently, our video analysis of the actual dance behavior yielded the same conclusions as the simulation analysis (Fig. 2e), i.e., the variation in directional information was reduced as the distance from the hive to the feeder increased. In summary, iMoAD-f suggested that a variation of 15° is the upper limit for gaining a foraging advantage from informative dancing.

**Switching feeders.** Although Figure 2d indicates that a colony always gained an advantage from dancing when the variation was below 5°, it does not explain why real bees dance with an error of 15° (just below the upper limit) or larger. In other words, does a variation of 15° have any biological significance? To answer this question, we next examined the other function, i.e., the adaptability of a colony to change in an environment, by focusing on the time course of the number of visits to each feeder. The time course showed a clear switch in the feeders visited by the waggle dance colony, but not by any of the other colonies (Fig. 2f). The random-search colony, no-communication colony, and random-information dance colony never ceased visiting the low concentration feeders.

### Table 2 | Concentration-dependent transition probabilities of dancing and following

| Concentration (M) | Successful Flight | Following |
|-------------------|-------------------|-----------|
| 0.75              | 0.15t             | 0.05t     |
| 1.0               | 0.26t             | 0.05t     |
| 2.5               | 1.0t              | 0.1t      |
|                   |                   |           |
| 0.75              | 0.13t             | 0.35t     |
| 1.0               | 0.14t             | 0.35t     |
| 2.5               | 0.6t              | 0.3t      |
|                   |                   |           |
| 0.75              | 0.72t             | 0.6t      |
| 1.0               | 0.6t              | 0.6t      |
| 2.5               |                   |           |

\( t \): Seeley et al. Although Seeley et al. did not show the probabilities for wandering and foraging for 2.5 M sucrose solution, we can easily obtain these probabilities based on the 1.0 probability of dancing.

\( t \): This study. From Supplementary Figure S1b, we obtained 0.6 by simple probability calculation.

\( t \): We assumed that the probability for foraging could be higher, i.e., that a colony could tend to collect more food, if the concentration were higher. This is because a higher nectar concentration was associated with a reduction in begging. The reduction of begging is associated with collecting food.

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We calculated the rate of visits (see Methods), and found that in phase 1 (at 4 hours), the waggle dance colony had clearly visited the currently more profitable feeder (Fig. 3a, b), when the two feeders were available. Other colonies did not show such distinct behavior, although there were slightly higher visit rates for the more profitable feeder. In the case of the larger variations (30° and 60°), the visit rate for the more profitable feeder by the waggle dance colony was lower than that of the other variations at each of the distances (Fig. 3a, b). When five or more feeders were made available, the visit rates for the most profitable feeder in phase 1 were more intensive than those for any other feeders (Fig. 3b). In phase 2 (at 8 hours), clearly higher visit rates for the most profitable feeder (feeder 2) by the waggle dance colony were seen, but no distinct differences were noted with the other colonies. When the number of feeders increased, the correct responses to the change in feeder concentration were observed only in the waggle dance colony (Figs. 2f and 3b). Detailed observations, however, revealed cases in which the waggle dance colony failed to switch (Fig. 3c and d). Specifically, increases in both the distance to the feeder and variation were associated with a higher number of failures (Fig. 3c).

Quantitative analysis of switching. To quantify the rate of success in switching the visited feeder, further analysis was carried out using only the waggle dance colony with a variation of 0–15°, because iMoAD-f suggested that only the waggle dance colony could perform the switch and because iMoAD-f revealed that the waggle dances were not advantageous when the variation was 30° or 60°. We calculated the switching index (see Methods) for all runs tested. The bees exhibited good switching, i.e., a high switching index, when the feeders were located 400 m or 750 m (Fig. 4) away from the hive, regardless of the number of feeders. When the feeders were 1000–1250 m away from the hive, negative switching indices were found in more cases than under the shorter-distance conditions (Fig. 4). Surprisingly, negative switching indices were seen even when the variation was 0°, although positive switching indices were also observed under these conditions. When the feeders were 1500 m or farther away from the hive, the bees failed to switch more often than when the feeders were closer. Strikingly, the negative switching indices were always near −100, while the range of positive switching indices was broader. Thus, the occurrence histogram of the switching indices showed two peaks—namely, one around 80 and the other at −100 for the variation of 0° or 5°. In contrast, when the variation was 15°, equal distribution (no prominent peak) was found in the histogram (Fig. 4).

For further analysis of the switching manner, we calculated the degree of advantage by waggle dances (see Methods) for each combination of variation and distances. When the variation was 0°, 5°, or 10°, a degree of advantage of 4 was found for all combinations. When the variation was 15° (Fig. 5a), a degree of advantage of 4 was found only for the 400-m feeder condition and a degree of 0 was found only for the 2000-m feeder condition. In other five combinations, the degree of advantage, the degree of advantage ranged between 1 and 3.
Figure 3 | Visit rates coded by color. (a) and (b), visit rates of all simulation results (20 runs for each simulation condition) for the two-feeder condition in phase 1 (a) and all conditions in phase 1 and 2 (b). Only the waggle dance colony clearly showed high visit rates for feeder 1 in phase 1, meaning that a greater number of bees in the waggle dance colony visited the most profitable feeder and that bees in other colonies visited each of the feeders with almost equal frequency. In phase 2, the random-search colony, no-communication, and random-information dance colonies showed almost no switching, although the random-information dance colony exhibited some switching under the two-feeder conditions. Thus, only the waggle dance colony succeeded at switching visited feeders in response to environmental changes. (c) and (d), visit rates by the waggle dance colony under 7-feeder conditions (c) and under the variations of 0° and 5° with high magnification (d). In phase 1, feeder 1 attracted the largest number of visits (c). In phase 2, although feeder 2 was expected to attract the largest number of visits, in some cases feeder 1 remained the most frequently visited feeder, indicating that switching was not successfully carried out in all cases (the white arrows in (d) indicate the representative cases). F1-10, feeder 1–10; NC, no-communication colony; RD, random-information dance colony; RS, random-search colony; WD, waggle dance colony.
We found that the number of negative switching indices increased as the distance to the feeders increased (Fig. 5b). The prominent failures in switching, i.e., the cases in which the switching index was nearly −100, were pronounced when the feeders were located 1500 m or longer away from the hive for a variation of 10° or less. When the variation was 0° or 5°, such prominent failures in switching began to be found at a shorter distance (1000 m). However, when the feeders were located 400 m away, the switching indices usually took large positive values. Because all the combinations described above were assigned a degree of advantage of 4, there might be no or little relationship between the degree of advantage and the switching manner. Interestingly, for a distance of 1750 m or longer and a variation of 5° or 10°, the histograms showed only one peak at −100. This means that bees very often failed to visit the selected feeder.

Discussion

We created an individual-based Markov Model of honeybee dance-guided foraging, iMoAD-f. iMoAD-f was validated by several lines of results that were consistent with previous findings from biological and theoretical experiments. Most importantly, iMoAD-f expressed bee-foraging behavior similar to that in Seeley’s experiment20. It is known that information error (variation) decreases with increases in feeder distance4,5,33, and that colonies respond to changes in feeder conditions34. The waggle dance was found to be advantageous when fewer feeders were available, i.e., when the feeders were difficult to find, both in biological experiments17–19 and modeling studies30–32,34,35.

Although numerous modeling and simulation studies of honeybee foraging have been reported to date30–34–40, many, if not all, have used quantitatively and physically unmeasurable parameters, e.g., homing motivation and foraging motivation38. This kind of partial lack of representation of real-world conditions has prevented a truly comprehensive understanding of the functions and mechanisms of the waggle dance. Here, we used only physically measurable parameters, which enabled us to evaluate iMoAD-f directly by using experiments with real bees. Furthermore, the iMoAD-f could be improved quickly by incorporating any additional features that we found necessary—for example, the diurnal change in variance of dance direction change in the source of the day35, characteristics of the outbound flight of the foragers with the dance information (social information) or individual memory (private memory)35, and the visiting pattern of the foragers, such as the flower constancy34 and the cross visit66. Further experiments on the followers would facilitate improvement of iMoAD-f.

Figure 4 Switching indices by the waggle dance colony for all simulation runs. Switching indices of all single simulation runs were color-coded. The column at the far right shows the occurrence histograms of switching indices of the corresponding row. When the error was 0° or 5°, high-risk, high-return foraging occurred. SI: switching index.

Figure 5 Details of switching. (a), The best colonies when the error was 15°. (b), Occurrence histograms of switching indices for each variation-distance combination shown in Figure 4. The position of each histogram corresponds to the position in the color-coded panels shown in Figure 4. The color of each histogram indicates the degree of advantage; red, magenta, and blue histograms indicate degrees of advantage of 4, 1–3, and 0, respectively. D.A., degree of advantage; NC, no-communication colony; RD, random-information dance colony; RS, random-search colony; SI, switching index; WD, waggle dance colony.
Noise and error are a major problem in biological systems, and often reduce the accuracy of these systems and degrade performance. Nevertheless, in the case of bee colonies, providing information with no noise/error was also not always suitable. IMoAD-f, in which the dance probability was defined only by a successful forager that evaluated the profitability of the food source, showed that dances with the smallest variation yielded the greatest number of visits to a feeder with the highest sucrose concentration (Supplementary Fig. S5). However, many foraging bees completely failed to switch feeders (Fig. 4). The finding, under the conditions we simulated, that extremely accurate directional information (overly narrow tuning) in waggle dances recruited the most bees to the advertised feeder suggested that the bees had a much lower chance of finding another feeder. Consequently, the colony had to abandon its flexible response to abrupt changes in the foraging environment.

We have shown the possible role of information error in waggle dances, but further questions must be answered for a full understanding of the function and mechanism of these dances. Does a follower bee use information from all waggle runs it attends or only from selected waggle runs? Is the dance efficiency a result of the error in dance information or simply a by-product of inaccurate dances? We believe that the improvement of a colony’s flexibility by dance inaccuracy is not simply a fortunate by-product because the degree of imprecision of the directional information depends on the target, i.e., we speculate that bees might actively exploit error for foraging. To examine this hypothesis, the variation in dance information must be controlled in a real-world setting. One possibility is to perform biological experiments using a robot bee that could recruit nestmates to a food source by the waggle dance rules. Another possibility is to use a computer simulation. The combination of biological experiments and theoretical research will provide new insights and a fuller understanding of the waggle dance.

Noise sometimes contributes to an improvement in system performance. For example, mechanosensory neurons can detect small signals using noise. In ants, noise can improve collective decision-making. Hence, a specific range of error in the waggle dance can be expected to be of significance. Efficient food collection is likely to have selective benefits. Food collection must consist of two major activities: acquiring food (choosing the effective foraging strategy in our work) and responding quickly to changes in the environment relating to food collection (switching the visited feeder). Although we cannot exclude the possibility of physical constraints, taken together with the dependency of precision in the directional information on the target and the smallest variation ranges, our present results suggest that a 10–15° variation in the conveyance of food-source information might be optimal for guaranteeing foraging success under a dynamically changing environment. Therefore, this range of error might be the result of a naturally selected trade-off between risk and success.

Methods

Video analysis. To obtain parameters for modeling and simulation, behavioral analysis was performed as described in our previous work. Briefly, about 1500 honeybees, *Apis mellifera*, were kept with a queen in an observation box under fully natural conditions at the campus of Hokkaido University (Sapporo, Japan). The experimental location was a flower-rich area, but there were fewer flowers than in spring. This situation sufficiently induced waggle dances. There were 1–10 dancing bees at one time. The bee behavior on the vertical comb was videotaped at 30 frames/second (GR-HD1; JVC, Tokyo, Japan) from 8:30 am to 4:00 pm on several days in August and September, 2006 (temperature, 25–36°C on most days). All analyzed data were obtained off-line and frame-by-frame with a computer algorithm obtained from our own analysis of videotapes or from the published literature (Supplementary Fig. S1 and Supplementary Table S1). In addition to probability, states also changed according to state-specific rules. For example, if a wandering bee had moved closer to a dancing bee than the distance necessary to find a dancer (Par-38 in Supplementary Table S1), the state of the wandering bee transitioned from wandering to following, irrespective of probability. On the other hand, if the following bee exceeded that distance, this bee was automatically considered to be wandering. These transitions were incorporated into our simulation based on our preliminary observation that most bees within one body length (ca. 1 cm) appeared to orient to a dancing bee reflexively, and those spaced to lose their dancer when the dancer changed walking direction rapidly, when a follower was obstructed momentarily by another bee, or when the dancer was out of range (data not shown). For more details about modeling, see the Modeling and simulation section in the Supplementary Methods.

Simulations. A single-day (8-hour) of activity was simulated (time step = 1 sec, thus 28,800 time steps in total). The hive contained 1000 worker bees. The 2, 5, 7, or 10 feeders were located radially around the hive at 400, 750, 1000, 1250, 1500, 1750, or 2000 m away from the hive. All feeders were set at the same distance in a single simulation. In the experiment with two feeders, the concentration of sucrose solution in feeder 1 was 2.5 M and that in feeder 2 was kept constant at 2.0 M for 4 hours (phase 1). In the second 4-hour period (phase 2), the concentration of the sucrose solution in feeder 1 was 0.75 M and that in feeder 2 was 2.5 M. This concentration treatment was the same as in Seeley’s experiment, which was used for the validation of IMoAD-f. The feeders were always placed in opposite directions from the hive (Supplementary Fig. S7). In other scenarios, the sucrose concentrations in feeder 1 and 2 were the same as those in the two-feeder simulations, and the sucrose concentrations in the remaining feeders (feeder 3–10) were kept constant at 1.0 M throughout the experiment. The feeder closest to feeder 1 was designated feeder 3, and that closest to feeder 2 among the remaining feeders was feeder 4. Although we saw essentially the same results with a simple reverse of the concentrations when the 2 feeders were located 400 m away from the hive, we kept Seeley’s concentration conditions in order to permit an easy comparison of results throughout all simulation experiments. The directions of feeders from the hive were randomly distributed for every run, but the same set of 20 feeder-distribution conditions was used for the same number of feeder conditions (Supplementary Fig. S7). Thus, we were able to eliminate unexpected spatial effects of feeder distribution and investigate only the influence of distance. Feeders were always located more than 120 m (twice the radius of the search area for a flying bee) apart in order to prevent a flying bee from finding more than one feeder at the same time. We performed the simulation once for each set of 20 feeder distributions. To execute simulations under conditions similar to those of Seeley et al., each feeder was known by 10 bees as an initial condition.

Analysis. For the comparison with Seeley’s experiment, we excluded initially-informed bees from the analysis and counted only the number of recruited individuals, not visits, every 30 minutes. For other analyses, we used all 1000 bees to evaluate the benefits to an entire colony. We counted the total number of visits to all feeders in each trial, and then calculated the average number of visits. Using energetics information (Supplementary Table S1), we also calculated the colonial energy balance by subtracting the energy out from the energy in, and then compared the results among virtual colonies. The advantage of dancing behavior was examined by comparing the number of visits in feeders and energy balance over a single day among four virtual colonies as follows (for details see Supplementary Methods): a random-search colony in which bees did not communicate to memorize the location of the feeders or perform dances; a no-communication colony in which bees memorized feeder locations, but did not communicate with each other (no dance was performed); a random-information dance colony in which bees memorized the feeder location and danced, but transferred meaningless directional information about the feeder; and a waggle dance colony in which bees memorized feeder locations, danced, and transferred significant information with variations of 0, 5, 10, 15, 30, or 60°. The colony with the highest value of energy balance was referred to as the best colony. If a colony or colonies were not significantly different from the best colony, these were also classified as best. For quantitative analyses of switching, the switching index was calculated for every run. Under our simulation, the most profitable feeder always changed after 4 experimental hours. Thus, we calculated the visit rate at 4 and 8 hours separately. The visit rate was obtained by dividing the number of visits to each feeder during each phase by the total number of visits to all feeders during the corresponding phase, and then the obtained value was multiplied by 100. Next, the switching index was used for examining colony responses to changes in the foraging environment. The switching index was calculated by subtracting the visit rate for the previously most profitable feeder, feeder 1, from that for the currently most profitable feeder, feeder 2, at the end of...
the simulation. Thus, the switching index ranged between −100 and 100, and if the bees correctly switched feeders, the switching index was positive. The occurrence histograms for the switching index were created by counting the number of cases for every switching index of 10 (for variation) or 20 (for a combination of variation and distance). For the detailed analysis of switching indices, switching was separately analyzed for each combination of variations and distances. All switching indices from the 2-, 5-, 7-, and 10-feeder condition for a particular variation-distance combination were pooled. The degree of advantage conferred by the waggle dances for each combination was calculated by counting the number of sub-conditions (each combination consisted of 4 sub-conditions, because simulations were done for 4 different numbers of feeders) under which the waggle dance colony was the best. If the waggle dance colony was the best under all sub-conditions for a particular combination, the degree of advantage was 4. In contrast, a degree of 0 indicated that there were no sub-conditions under which the waggle dance colony was the best for that combination.

Simulations and analysis were carried out using all possible combinations of the 4 different numbers of feeders, 7 distances, and 6 variations. Full Methods and all associated references are provided in the Supplementary Methods.

Statistics. All statistical analyses of colony effects were performed using Analysis of Variance (ANOVA) and a post-hoc Tukey-Kramer test unless otherwise mentioned.

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Additional information
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