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RESEARCH ARTICLE

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When should bees be flower constant? An agent-based model highlights the importance of social information and foraging conditions

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Abstract

- 1. Many bee species show flower constancy, that is, a tendency to visit flowers of one type during a foraging trip. Flower constancy is important for plant reproduction, but the benefits of constancy to bees is unclear. Social bees, which often use communication about food sources, show particularly strong flower constancy.
- 2. We aimed to better understand the benefits of flower constancy in social bees and how these benefits depend on foraging conditions. We hypothesised that sharing social information increases the benefits of flower constancy because social foragers share information selectively about high-quality food sources, thereby reducing the need to sample alternatives.
- 3. We developed an agent-based model that allowed us to simulate bee colonies with and without communication and flower constancy in different foraging environments. By varying key environmental parameters, such as food source numbers and reward size, we explored how the costs and benefits of flower constancy depend on the foraging landscape.
- 4. Flower constancy alone performed poorly in all environments, while indiscriminate flower choice was often the most successful strategy. However, communication improved the performance of flower constant colonies considerably in most environments. This combination was particularly successful when highquality food sources were abundant and competition was weak.
- 5. Our findings help explain why social bees tend to be more flower constant than solitary bees and suggest that flower constancy can be an adaptive strategy in social bees. Simulations suggest that anthropogenic changes of foraging landscapes will have different effects on the foraging performance of bees that vary in flower constancy.

KEYWORDS agent-based model, communication, flower constancy

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1 | INTRODUCTION

Most flowering plant species are animal pollinated and bees, in particular, are important pollinators of wild and agricultural plants (Klein et al., 2007; Ollerton et al., 2011). Several biological features explain why bees are helpful agents of reproduction for plants, including their abundance and their often broad (i.e. polylectic) diet in combination with a tendency to specialise on a particular flower type during an individual foraging bout. The latter behaviour, called flower constancy (Bateman, 1951; Chittka et al., 1999; Darwin, 1876; Grüter & Ratnieks, 2011; Waser, 1986), reduces conspecific pollen loss and heterospecific pollen deposition, both of which can reduce plant fitness (Campbell & Motten, 1985; Chittka et al., 1999; Morales & Traveset, 2008; Waser, 1986). Flower constancy is also thought to enhance the coexistence of different plant species and, thus, shapes plant community structure (Morales & Traveset, 2008; Song & Feldman, 2014).

From a pollinator perspective, however, the benefits of flower constancy are less obvious. Ignoring potentially superior flower species appears to contradict optimal foraging theory (King & Marshall, 2022; Latty & Trueblood, 2020; Waser, 1986; Wells & Wells, 1983). Why then are pollinators flower constant? The most widely accepted view is that flower constancy is driven by cognitive limitations, which can include (i) slow learning to forage efficiently on a new flower species, (ii) an inability to memorise more than one or a few flower types, (iii) unstable short-term memories which are prone to being erased by competing information or (iv) an inability to retrieve long-term memory about different flower species fast enough to be an efficient generalist (Darwin, 1876; Heinrich, 1979; Lewis, 1986; Menzel, 1999; Raine & Chittka, 2007; Waser, 1986; for reviews see Chittka et al., 1999; Grüter & Ratnieks, 2011). These cognitive limitations are likely to cause time delays as a bee tries to extract nectar or pollen from a flower after switching from a different species and they may increase switching times (Chittka et al., 1999; Goulson et al., 1997; Lewis, 1986; Raine & Chittka, 2007).

The "cognitive limitations hypothesis" as an explanation for flower constancy is not without challenges. Given that efficient foraging is likely to be under strong natural selection due to its effects on reproductive success (Heinrich, 1979), why does natural selection not lead to the evolution of lower flower constancy in all bees? How can we explain that individual bees often show plasticity in their flower constancy, for example by being more flower constant after finding good rewards (Chittka et al., 1997; Grüter et al., 2011; Wells & Rathore, 1994; but see Hill et al., 1997) or when the distances between food sources are shorter (Gegear & Thomson, 2004; Kunin, 1993; Marden & Waddington, 1981)? Why do bee species vary in their degree of flower constancy? Social bees, in particular, are often highly flower constant (Butler, 1945; Free, 1963; Heinrich, 1976, 1979; Hill et al., 1997; Kozuharova, 2018; Pangestika et al., 2017; Rossi et al., 2015; Slaa et al., 2003; White et al., 2001; but see Martínez-Bauer et al., 2021), while flower constancy seems to be less pronounced in solitary bees (Bateman, 1951; Campbell & Motten, 1985; Eckhardt et al., 2014; Jakobsson et al., 2008; Pohl et al., 2011; Smith et al., 2019; Waser, 1986; Williams & Tepedino, 2003). Smith et al. (2019), for example, studied the composition of pollen carried by 56 bee species and found that individual social bees showed a higher degree of specialisation during a foraging bout than solitary species. Different ecological needs could explain this difference. Solitary bees need to collect all required nutrients by themselves, potentially favouring a strategy of mixing resources during a foraging trip even if this has energetic costs (e.g. Eckhardt et al., 2014; Williams & Tepedino, 2003). In social species, on the other hand, different bees from the same colony can specialise on different flower species to cover their nutritional needs.

Foragers of many social bees share information about profitable food sources, and this could affect the value of flower constancy. Honeybees use the waggle dance to indicate the odour (type) and location of profitable food sources (von Frisch, 1967) and some stingless bees lay pheromone trails (Grüter, 2020; Jarau & Hrncir, 2009; Lindauer & Kerr, 1960; Nieh, 2004). Stingless bees and bumblebees inform nestmates about the availability and odour of a good food source by means of excitatory or jostling runs inside their nest (Dornhaus & Chittka, 1999; Hrncir, 2009). Trophallaxis-food transfer between bees-is performed by honey bees and stingless bees (Farina et al., 2012; Farina & Grüter, 2009; Hrncir et al., 2006; Krausa et al., 2017; von Frisch, 1967) and is another behaviour that allows nestmates to learn the odour of available food sources (Aguilar et al., 2005; Farina et al., 2005; Lindauer & Kerr, 1960; von Frisch, 1967). These different communication behaviours share two common features. First, they depend on food quality. Dances, pheromone trails, jostling runs and trophallaxes occur at higher frequencies if the exploited food sources are of higher quality (Farina et al., 2012; Hrncir, 2009; Krausa et al., 2017; Lindauer & Kerr, 1960; von Frisch, 1967). Second, during these social interactions, nestmates can learn the odour of the exploited flower species and acquire a preference for this flower species in the field (Dornhaus & Chittka, 1999; Farina et al., 2012; Lindauer & Kerr, 1960; von Frisch, 1967).

Heinrich (1976) was probably the first to propose a link between recruitment communication and flower constancy. Since recruiting bees share information selectively about high-quality food sources, recruits can discover profitable flower types without the costs of sampling different, lower-quality plant species. This does not require that foragers are able to direct nestmates to a specific location, as in honeybees and some stingless bees, but depends more generally on foragers biasing the food search towards flower types that are more profitable. Among the social bees, bumble bees seem to be less flower constant that honey bees (Bateman, 1951; Grant, 1950; Martínez-Bauer et al., 2021; Smith et al., 2019), possibly because their communication system is less sophisticated than that of honey bees (Heinrich, 1976).

Experimental studies of the benefits of flower constancy and how they depend on social and ecological traits are logistically challenging for several reasons. For example, it is often not possible to manipulate the degree of flower constancy while keeping other

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factors constant. Agent-based simulation models can be a useful complementary tool to evaluate how biological and ecological factors affect the benefits of a behavioural strategy. We developed an agent-based simulation model to test the hypothesis that flower constancy is more beneficial in bees that communicate about profitable food sources than in bees without communication. Colonies consisting of virtual bees (agents) were either flower constant or they chose food sources randomly (indiscriminately) in environments that varied in the number and quality of food sources. Some studies have found that bees adjust the degree of flower constancy depending on the foraging conditions, being more flower constant if the rewards on offer are better (Chittka et al., 1997; Grüter et al., 2011; Wells & Rathore, 1994; but see Hill et al., 1997) and the distances between food sources (or density) are shorter (Gegear & Thomson, 2004; Kunin, 1993; Marden & Waddington, 1981). We, therefore, expected flower constancy to be more beneficial in environments with more flowers and larger reward sizes. Exploitation competition, on the other hand, is expected to favour an indiscriminate choice because greater competition increases the costs of rejecting a reward (Pulliam, 1974).

TABLE 1 Overview of the model variables and the used values

2 | MATERIALS AND METHODS

We built an agent-based model (ABM) using the programming software NetLogo 6.1 (Wilensky, 1999). The model simulates an environment with a colony surrounded by food sources. The agents ("bees") operate on a two-dimensional square grid with 400×400 patches. A single patch length corresponds to 5 m and 1 tick corresponds to 1 s. Thus, the size of the virtual world corresponds to 2×2 km. The nest of the colony is positioned in the centre of the grid (x = 0, y = 0). In the default situation, environments contained two different flower types that differed in the rewards they offered. The model is an extension of a model built to study foraging distances in bees (Grüter & Hayes, 2022).

The model allows simulating a wide range of parameter values, but for the purpose of this study we based our default parameters, such as the nest stay time (t_{nest}) , flight speed (v_{flight}) , metabolic costs of flying (M_{cost}) , and crop capacity (*Crop*) on the Western honeybee *Apis mellifera* because we have accurate information about these relevant biological parameters in *Apis mellifera*. Other values were tested (see Table 1 and section *Sensitivity analysis and model exploration*).

Variables	Description	Default values	Other values tested	Information source
Colony size	Number of foragers in a colony	100	5-300	Westphal et al. (2006) and Grüter (2020)
FS _{number}	Number of food sources per type	3000	1500, 4500	Arbitrary
FS _{size}	Size of food sources	1 patch		Arbitrary
FS types	Number of food source types	2	4	Arbitrary
Reward _{HQ}	Nectar amount per food source	5 μL	2.5 μl, 10 μL	Willmer (2011)
Reward _{LQ}	Nectar amount per food source	2.5 μL	1.25 μl, 5 μl	Willmer (2011)
V _{flight}	Flight speed	1.4 patch/tick		von Frisch (1967)
M _{cost}	Metabolic costs of flight, J/tick	0.032	0.016, 0.064	Heinrich (1975) and Willmer (2011)
$t_{\sf flower-stay}$	Time spent at food source	60 ticks	20, 180	Arbitrary
V _{nest}	Movement speed inside nest	0.1 patch/tick		Arbitrary
t _{nest-stay}	Time in nest between trips	300 ticks	150, 450	Farina (2000)
Crop _{HQ}	Crop load when flower constant to HQ food sources	50 µ L	25 μL, 100 μl	Núñez, 1966
Crop _{LQ}	Crop load when flower constant to LQ food sources	25µL	12.5 µl, 50µl	Núñez (1966)
Crop _{Random}	Crop load without flower constancy	37.5 μL	50µL	Núñez (1966)
$t_{ m refill}$	Time until food sources offer food again	0 ticks	1200, 3600	Stout and Goulson (2002)
Lévy μ	Lévy flight parameter	1.4	1.8, 2.4	Reynolds (2009)
Constancy _{HQ}	Probability to remain constant to a type after leaving a high-quality food source during flower constant simulations	100%		Grüter et al. (2011)
Constancy _{LQ}	Probability to remain constant to a type after leaving a low-quality food source during flower constant simulations	100%	90%, 95%	Grüter et al. (2011)

Foragers in social bees use different behavioural mechanisms to transmit social information and, thereby, influence the food source preferences of their nestmates (see introduction). The model does not simulate a particular behaviour, but a generic process that biases the food preferences of nestmates, which could correspond to jostling runs, trophallaxis or the waggle dance.

2.1 | Entities and state variables

2.1.1 | Bees

The default colony size was 100 agents (forager bees), which corresponds to the size of the forager pool in many species of bumble bees (Westphal et al., 2006) and stingless bees (Grüter, 2020). Agents could assume any of the following states: (1) generalists, (2) feeding forager, (3) searching forager, (4) returning forager, (5) inside-nestworker and (6) influencer (see Figure 1).

Agents begin the simulation in the centre of the nest with energy = 0 as generalists. They then move at a flying speed of 1.4 patch/tick (v_{flight}), corresponding to a flight speed of *Apis mellifera* (7 m/s, von Frisch, 1967). Their random search behaviour follows a Lévy-flight pattern (with μ = 1.4 as default; Reynolds, 2009; Reynolds et al., 2007). A Lévy-flight consists of a random sequence of flight segments whose lengths, *l*, come from a probability distribution function having a power-law tail, $P(l) \sim l^{-\mu}$, with $1 < \mu < 3$ (Reynolds et al., 2007). The speed of agents moving inside the nest (v_{nest}) was arbitrarily chosen to be 0.1 (patch/tick). Flying has a



FIGURE 1 State diagram showing the different states of the agents and the possible transitions between states. Here, yellow flowers were arbitrarily chosen to represent a lower-quality food source, therefore, the default probability that foragers visiting yellow flowers would become influencers after their return to the nest was 0. Modified from Grüter and Hayes (2022).

metabolic cost (M_{cost}) of 0.032 Joule (J) per tick in the default condition (Heinrich, 1975; Willmer, 2011). Once an agent encounters a food source, they remain on the food source for 60 ticks ($t_{flower-stay}$) under default conditions (*feeding foragers*), irrespective of whether they were choosing indiscriminately or are flower constant. Thus, we assume that the time spent handling a flower or flowers in a patch and extracting the reward is the same for flower constant and indiscriminate foragers. This was chosen as the default condition to explore whether flower constancy can be an adaptive strategy in the absence of cognitive constraints.

The agent then continues to forage (searching foragers) until its crop is full, after which it returns to the nest (returning foragers) to unload its energy and stay in the nest for 300 ticks ($t_{nest-stay}$; Farina, 2000; Seeley, 1986; von Frisch, 1967). In the default condition, only foragers visiting the high-quality food source could become *influencers* (i.e. bees that bias the food choice of other bees) upon return to the nest. *Influencers* target *inside-nest-workers* that are not yet flower-constant to the high-quality food type by changing the latter's preference if they encountered each other on same patch inside the nest. Following such an encounter, *inside-nestworkers* become flower constant for the high-quality type.

Since recruitment behaviours often depend on the food source distance (with greater foraging distances lowering the probability of recruitment), we simulated recruitment curves where the probability of becoming an *influencer* decreased with increasing distance of the last visited food patch (Figure S1).

2.2 | Food sources

In the default condition, two different types of food sources offering nectar rewards can be found in the environment, mimicking the typical situation in experimental flower constancy studies (e.g. Chittka et al., 1997; Goulson & Wright, 1998; Grüter et al., 2011; Ishii & Masuda, 2014; Wells & Wells, 1983). The food source types differ in the rewards they offer per visit. Natural bee-visited flowers offer between 0.1 and 10 μ l of nectar per flower (Willmer, 2011, p. 203). For the default condition, we chose 5 μ l (29.07 J) for the high-quality type and 2.5 μ l (14.535 J) for the low-quality type. This reward could represent an individual flower that offers a large reward or a small patch of several flowers, each offering smaller quantities, or it could represent a larger patch of flowers that is shared by several bees.

We tested different refill times (t_{refill}) for food sources: 0, 1200 and 3600 ticks (Stout & Goulson, 2002). When $t_{refill} = 0$, food sources became rewarding again immediately after the visit of a bee. This simulates conditions under which bees have a high probability of finding a reward after landing on food source, which might occasionally occur at food patches. With $t_{refill} = 3600$, a food source (flower or patch) was empty for the equivalent of an hour after it had been visited by a bee, leading to intense exploitation competition among bees. The number of food sources per type in the simulated environment varied between 1500 (low abundance) and 4500 (high abundance). Default conditions simulated even numbers of food sources for both food source types, but we also explored uneven food source abundances (Table 1). We measured the average foraging distance of bees during a simulation run to confirm that the simulated conditions led to naturally realistic average foraging distances for many social bees (271 ± 130 m; range 63-581m; N = 1800 simulations in default conditions; Kohl et al., 2020; Van Nieuwstadt & Iraheta, 1996; Walther-Hellwig & Frankl, 2000).

The energy collected by agents with a full crop was estimated in the following way: Apis mellifera can carry up to ~70µl of nectar in their crop, but they usually carry less (l'Anson Price et al., 2019; Núñez, 1966). The crop load has been shown to depend on the quality of the visited food source, with lower quality food sources leading to smaller crop loads (Núñez, 1966, 1970). Agents visiting the low-quality flower type foraged until their crop contained 25 µl, whereas agents visiting the high-quality food type collected 50µl per foraging trip. Generalist bees that choose indiscriminately have an intermediate crop load, reflecting the relative number of highand low-quality food sources in the environment. For example, in an environment with an even number of high- and low-quality food sources, they collect 37.5 µl per foraging trip. Sugar concentration of collected nectar varies considerably from c. 10%-70% (Seeley, 1986; Willmer, 2011). We chose an average sugar concentration of 35%, providing $5.814 \text{ J/}\mu\text{l}$.

Each simulation lasted 36,000 ticks (i.e. 10h), simulating a day with good foraging conditions. We measured the total energy collected by a colony during this period divided by the number of agents (Energy/bee). Our main questions were if the energy/bee depended on flower constancy (vs. indiscriminate choice), communication (vs. no communication), refill time, the number of food sources and reward size. We also tested situations when flower constancy was lower after visiting a low-quality food source (*Constancy*_{LQ}; Grüter et al., 2011), when there were 4 food source types and when indiscriminate flower choice increased the time to extract a reward from a food source (i.e. to simulate cognitive constraints; Chittka et al., 1999).

2.3 | Sensitivity analysis and model exploration

We varied a range of other factors to explore how they affected our results. These included colony size, crop load size, flower stay time, metabolic costs, nest stay time, Lévy flight μ , selectivity of communication (i.e. bees foraging on low-quality food source become *influencers* with the same probability as those foraging on the high-quality type) and the shape of the recruitment curve (see Figure S1).

We performed 30 runs per parameter combination. We do not provide *p*-values due to the arbitrariness of the simulation number but indicate 95%-confidence intervals to facilitate interpretation of effect sizes.

2.4 | Ethics approval

This study did not require ethical approval.

3 | RESULTS

3.1 | Food source abundance and refill speed

We found that communication about the high-quality flower type did not affect the collected energy if bees chose food sources indiscriminately (Figure 2). However, if colonies were flower constant, communication increased the energy collected by bees in all situations when the two flower types were equally abundant (Figure 2, see also Figures 4–8), showing an interaction between flower constancy and communication. The combination of communication and flower constancy was relatively more beneficial when high-quality food sources were easy to find, either because they were highly abundant (Figure 2c,f,i) or because visited food sources replenished quickly (Figure 2a–c). In the most favourable conditions, flower constancy in combination with communication was the most successful combination (Figure 2b,c). In all other conditions, indiscriminate choice was the most successful strategy. The relative abundance of the two food source types also played an important role. Flower constancy combined with communication was relatively more successful when high-quality food sources were more common than the low-quality flower type compared to when they were rarer than the lower-quality flower type (Figure 3). When high-quality food sources represented the common flower type, colonies with flower constancy and communication were either more successful (Figure 3b) or not much less successful than colonies with indiscriminate choice (Figure 3d). However, indiscriminate choice was considerably more successful when high-quality food sources were in the minority (Figure 3a,c). When high-quality food sources were particularly difficult to find, communication lowered the foraging success of flower constant colonies (Figure 3c). Under these circumstances, communication directs the foragers of a colony towards a rare food source, leading to long search times.

3.2 | Reward sizes

Reward quantities are known to affect flower constancy, with bees becoming more flower constant with increasing reward quantities (Chittka et al., 1997; Grüter et al., 2011; Wells & Rathore, 1994). In accordance with this observation, we found that flower constancy



FIGURE 2 Energy collected per bee (Joule) under varying food abundances (1500 [a, d, g], 3000 [b,e,h] and 4500 [c,f,i] food sources per type) and refill times (0 ticks = immediate refill, 1200 ticks = medium refill and 3600 ticks = slow refill). Colonies either showed flower constancy (constant) or they chose food source indiscriminately (random). Plots show the mean and the 95%-confidence based on 30 simulations (grey dots). Numbers show % of change compared to random choice without communication.



FIGURE 3 Energy collected per bee (Joule) when high-quality food sources were either rare (1500; a, c) or common (4500; b, d) compared to the low-quality food sources (3000). Default values were used for the other parameters (Table 1).

became relatively more successful (energy/bee) as reward sizes of both high- and low-quality food sources increased (Figure 4). However, indiscriminate choice was the most successful strategy in many tested environments.

3.3 | Time needed to collect a reward

The time bees need to extract a reward from a flower will affect the time costs of foraging decisions and, if the refill time is >0, it will affect the number of depleted food sources in the environment. Under default conditions, bees needed 60 ticks (1 min) to obtain the reward from a flower/food patch. We explored how different values for $t_{\text{Flower-stay}}$ affected the benefits of flower constancy and communication. Increasing the time needed to obtain a reward increased the relative benefits of combined flower constancy and communication compared to short reward collection times (Figure 5).

3.4 | Quality dependent flower constancy

Under default conditions, flower constancy did not depend on the quality of the food source ("spontaneous flower constancy", Hill et al., 1997). We simulated situations when bees visiting a low-quality

food source were slightly less flower constant (they had a 90% or a 95% chance to remain flower constant on the subsequent visit, as in Grüter et al., 2011). Our results show that this quality-dependent flower constancy considerably improves the energy collected by colonies following this strategy of quality-dependent flower constancy (Figure 6).

3.5 | Exploring environments with 4 food types

When environments provide four different types of food sources rather than two, flower constancy is less favourable overall (Figure 7). In other words, indiscriminate flower choice is highly beneficial in an environment where flower constancy would limit the options a forager has to a small subset (25% of all food sources) of all available food sources than with two food source types (Figure 7).

We tested situations where one, two or three of the four plant types were of high quality, while the remaining food sources were of low quality. While it was always beneficial to use communication when colonies were also flower constant, the relative benefits of communication diminished as the number of high-quality food types and the refilling time increased. Unsurprisingly, therefore, foraging in an environment that consists mainly of high-quality food sources belonging to different plant species somewhat diminishes the value



FIGURE 4 Energy collected per bee (Joule) when rewards were (a, d) smaller (2.5 and 1.25μ I) or (c, f) larger (10 and 5μ I) than in the (b, e) default situation (5 and 2.5μ I). Medium food source abundance was simulated; default values were used for the other parameters (Table 1).



FIGURE 5 Energy collected per bee (Joule) when the time required to obtain a reward from a food source was (a, d) shorter (20 ticks) or (c, f) longer (180 ticks) than in the (b, e) default situation (60 ticks). Default values and a medium food source abundance were simulated.



FIGURE 7 Energy collected per bee (Joule) when bees foraged in an environment of four flower species. In (a & d), one of the four plant types was of high quality, while the remaining three types were of low quality. In (b & e), two of four types were of high quality and in (c & f), three of flower types were of high quality.

of using communication to direct foragers towards higher-quality food sources.

bees favours flower constant colonies, especially those that also communicate the high-quality flower type to nestmates (Figure 8).

3.6 | Time penalty for non-specialists

So far, we have assumed that there are no additional time costs (e.g. as a result of cognitive limitations) for bees that do not specialise on a particular type of food source. To explore the consequences of cognitive limitations, we simulated situations when indiscriminate bees require more time to extract a reward from a food source compared to flower constant bees. A time penalty for indiscriminate

3.7 | Sensitivity analysis and model exploration

Varying colony size from 5 to 300 (Figure S2) does not greatly affect the general pattern observed for the default colony size of 100 (see Figure 2). When food sources refill immediately after a visit (Figure S2), bees do not experience exploitation competition and colony size does not affect the energy collected by individual bees. Increasing the refill time while also increasing the number of agents



FIGURE 8 Energy collected per bee (Joule) when non-specialists did not need more time than flower constant bees (a, d; default). Energy collected per bee (Joule) when non-specialists needed 50% (b, e) or 200% (c, f) more time than flower constant bees to obtain a reward from a food source. Food sources were refilling either immediately ($t_{refill} = 0$) or a medium rate ($t_{refill} = 1200$). Default values were used for the other parameters.

searching for food, on the other hand, increases exploitation competition and, therefore, lowers the energy collected by individual bees (Figure S2).

Using different recruitment curves (Figure S1) had no noticeable effect on the energy collected by bees, but non-selective recruitment (recruitment to both high- and low-quality food sources) lowers the collected energy to levels similar to those of flower constant colonies without communication (Figure S3). Changing the metabolic costs of flying has little effect on the overall pattern (Figure S4), whereas increasing the time spent inside the nest inbetween foraging trips reduces the energy collected by bees, but less so in colonies with flower constancy (Figure S5). Thus, longer nest stay times favour flower constancy. Flower constancy was also favoured when bees had smaller crop sizes (Figure S6). Changing the Lévy-flight μ (to 1.8 and 2.4) led to qualitatively similar results and is not shown.

4 | DISCUSSION

Results from our simulations suggest that flower constancy without communication is less successful than indiscriminate choice under all tested conditions. Flower constancy imposes significant costs because it (i) limits the available options to a subset of all available flowers, thereby increasing time and energy costs during foraging, and (ii) causes many foragers to specialise on a sub-optimal flower type. Communication about the high-quality flower type positively interacted with flower constancy (Figure 2) and considerably improved the foraging success of flower constant colonies. Communication allows a colony to focus on high-quality flowers, thereby reducing the second type of cost (ii). Many species of social bees have evolved mechanisms of reward-quality dependent recruitment communication, which allow *influencers* to affect the foraging decisions of their nestmates towards a particular high-quality flower type, mainly via olfactory learning (Dornhaus & Chittka, 1999; Farina et al., 2012; Jarau & Hrncir, 2009; Lindauer & Kerr, 1960; von Frisch, 1967). This, in turn, lowers the benefits of sampling alternative flower species and highlights the importance of social information use as a process of information-filtering (Grüter et al., 2010; Rendell et al., 2010). Our findings can help explain why social bees tend to be more flower constant than solitary bees (e.g. Smith et al., 2019; Waser, 1986).

The general foraging conditions had a strong effect on the value of flower constancy and the strength of its interaction with communication. Flower constancy in combination with communication was the most successful strategy when foraging conditions were very favourable, while indiscriminate choice was the better strategy when foraging options were more limited. For instance, flower constancy in combination with communication was beneficial when foragers did not encounter empty food sources (refill time of 0) and food sources were abundant (Figure 2b,c), when most food sources were of high-quality (Figure 3b) and when rewards were large (Figure 4c). These findings are consistent with empirical studies showing that bees are more flower constant when flower density is higher (Chittka et al., 1997; Kunin, 1993; Marden & Waddington, 1981) and rewards are larger (Chittka et al., 1997; Greggers & Menzel, 1993; Grüter et al., 2011). Similarly, predator-prey models show that the abundance of a prey item has a positive effect on diet specialisation of the predator (Pulliam, 1974). If food sources took time to

replenish, resulting in many empty food sources due to exploitation competition, indiscriminate choice was more successful (Figure 2di), suggesting that rejecting flowers due to flower constancy is more costly in environments that offer fewer options.

Changes in the temporal dynamics of foraging trips affected the performance of the different strategies by changing the relative costs of ignoring flowers (i) and choosing suboptimal food sources (ii). Flower constancy in combination with communication performed relatively better if bees required more time to extract a reward from a food source (Figure 5). One explanation for this is that visiting lowquality food sources, which is common with indiscriminate choice, becomes relatively more costly as the time costs of a flower visit increase. Thus, longer flower handling times, e.g. due to a complex flower morphology, favour flower constancy from both an adaptive and a constraints-based perspective (see Chittka et al., 1999 for arguments based on cognitive constraints). Flower constancy in combination with communication performed relatively better when bees stayed in their nest longer (Figure S5) and had smaller crops (Figure S6). These findings are somewhat puzzling, but one explanation could be that longer nest stay times provide influencers with more opportunities to communicate their findings to other bees. When food sources need time to replenish, longer nest stay times will reduce the number of depleted food sources a bee encounters, which favours flower constancy in combination with communication (Figure 2b,c). Similarly, when bees have smaller crop sizes, they visit fewer food sources per trip and spend a larger proportion of their time in the nest, reducing exploitation competition and the number of depleted food sources. Crop size will depend on body size and one might, therefore, predict that smaller bees are more flower constant, which is consistent with comparative data (Smith et al., 2019). However, it is unlikely that there is straightforward relationship between crop size, body size and flower constancy in nature because body size covaries with numerous other extrinsic and intrinsic factors, including foraging conditions, metabolic costs, flying speed or sensory acuity (Gervais et al., 2020; Grab et al., 2019; Spaethe et al., 2007), all of which might affect flower constancy. While our model simulates nectar foraging, we think it is likely that the general conclusions also hold if the simulated reward is pollen. Factors that reduce the number of available foraging options will often improve the relative value of indiscriminate choice, irrespective of whether a bee collects pollen or nectar.

The Western honey bee *Apis mellifera* is strongly flower constant, but there is disagreement about whether and when flower constancy depends on the profitability of visited flowers. Some studies have suggested that flower constancy is often "spontaneous", i.e. unrelated to reward size (Hill et al., 1997; Sanderson et al., 2006; Wells & Wells, 1983), whereas others have found that honey bees adjust flower constancy according to the profitability of rewards (Chittka et al., 1997; Greggers & Menzel, 1993; bumble bees: Heinrich, 1976, 1979; reviewed in Grüter & Ratnieks, 2011). Our simulations show that context-dependent flower constancy is more successful than strict ("spontaneous") flower constancy (Figure 6). When bees visiting the less profitable food type were only 90%–95% flower constant, colonies collected about 25% more energy than colonies with strict flower constancy. As is the case with communication, context-dependent flower constancy allows bees to switch from the low-quality to the high-quality flower species over time (type (ii) costs).

Human impacts have significantly affected the diversity of plant species found in some environments, especially in intensively farmed habitats (e.g. Potts et al., 2010; Tew et al., 2021), which is likely to affect the costs and benefits of flower constancy. In our simulations, flower constancy performed considerably worse when there were four rather than two flower types (Figure 7). With more plant species present, flower constant bees will ignore most of the available options and focus on a small subset of all food sources, thereby dramatically increasing opportunity costs (type (i) costs). Thus, bees should be less flower constant in more diverse foraging environments. Flower constant bees, in turn, might suffer a reduction in foraging success in more biodiverse habitats. These findings challenge the reasoning behind the "costly information hypothesis", which argues that flower constancy is an adaptive foraging strategy because acguiring information about suitable alternatives would cost too much time and energy if there are several plant species available (Chittka et al., 1999; Grüter & Ratnieks, 2011). In flower diverse environments, bees should accept even low-quality food sources if it means they can cut time and energy costs imposed by flower constancy. Empirical studies on the links between floral diversity and flower constancy provide contrasting results. While Gervais et al. (2020) and Martínez-Bauer et al. (2021) found that increasing plant diversity was associated with lower flower constancy in Bombus impatiens and B. terrestris, Austin et al. (2019) found that bumble bees became more flower constant when there are more options available. The latter finding is more consistent with a "cognitive limitations" perspective, since deciding among more options would be cognitively more challenging and flower constancy, therefore, a possible solution to avoid switching costs (see also Chittka et al., 1997; Gegear & Thomson, 2004). Decision making is often impaired as the number of choices increases (Latty & Trueblood, 2020). The different studies differ in that the first two were performed under natural conditions, whereas Austin et al. (2019) was experimental. Non-experimental surveys can be confounded by numerous factors, such as differences in rewards, clustering of flowers or management, whereas experimental studies might fail to capture crucial features of natural environments that affect decision-making (Fawcett et al., 2014).

Agent-based models have important limitations. Simulation outcomes depend on the underlying assumptions and the parameters chosen when building the model, some of which are arbitrary or simplistic. As a result, ABMs potentially miss important natural features that shape decision-making (Fawcett et al., 2014). For example, we assumed that food sources are randomly distributed, whereas natural foraging environments are often spatially heterogenous and patchy, which is likely to affect the value of flower constancy. Patchiness can lead to flower constancy "by accident" if bees forage in large patches, even if they choose flowers indiscriminately. We might, therefore, expect increasing patchiness to lead to more similar outcomes for flower constant and indiscriminate foragers. Pulliam's (1974) predator-prey model found that an increasingly clumped prey distribution favours a more specialised diet in predators and we might expect a similar finding in plantpollinator interactions. Agent-based models also have important strengths because they allow us to systematically vary factors that cannot be manipulated experimentally, such as tuning flower constancy or recruitment communication while keeping all other factors constant. ABMs should be seen as a useful tool to complement empirical studies.

One of the aims of our model was to test whether flower constancy could be an adaptive strategy per se under some foraging conditions, i.e. in the absence of cognitive constraints. If, however, switching between flower species leads to increased time costs or reduced reward sizes (Chittka et al., 1999; Darwin, 1876; Grüter & Ratnieks, 2011; Lewis, 1986; Raine & Chittka, 2007), flower constancy, with or without communication, becomes a much more beneficial strategy under a wide range of conditions (Figure 8c,f). Thus, cognitive constraints are a plausible explanation for flower constancy in some solitary pollinators (Lewis, 1986). The reasons for flower constancy in pollinators are likely to be complex and depend on both constraints and adaptive processes, to varying degrees in different species. However, our results suggest that flower constancy is likely to be more beneficial in social species due to social traits, rather than the result of poorer cognitive abilities in social bees compared to solitary bees (Amaya-Márquez & Wells, 2008; Dukas & Real. 1991).

AUTHOR CONTRIBUTIONS

Lucy Hayes and Christoph Grüter contributed to the design of the model, the generation and interpretation of the data and the drafting and revising of the manuscript. Both authors gave final approval for publication.

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CONFLICT OF INTEREST

We declare that we have no conflict of interest.

DATA AVAILABILITY STATEMENT

Data available from the Dryad Digital Repository: https://doi. org/10.5061/dryad.rr4xgxdch (Hayes & Grüter, 2022a). The model code is available here: https://doi.org/10.5281/zenodo.7385899 (Hayes & Grüter, 2022b).

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SUPPORTING INFORMATION

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