

UCLA

UCLA Electronic Theses and Dissertations

Title

Path choice by Argentine ants (*Linepithema humile*) in novel and complex environments

Permalink

<https://escholarship.org/uc/item/1701x42s>

Author

Lessig, Emily

Publication Date

2019

Peer reviewed|Thesis/dissertation

UNIVERSITY OF CALIFORNIA

Los Angeles

Path choice by Argentine ants (*Linepithema humile*) in novel and complex environments

A thesis submitted in partial satisfaction of the requirements

for the degree Master of Science in Biology

by

Emily Lessig

2019

© Copyright by

Emily Lessig

2019

ABSTRACT OF THE THESIS

Path choice by Argentine ants (*Linepithema humile*) in novel and complex environments

by

Emily Lessig

Master of Science in Biology

University of California, Los Angeles, 2019

Professor Peter Nicholas Nonacs, Chair

Foragers often face difficult landscapes in terms of encountering previously unexplored areas or areas where risk is present along paths to food. Cooperatively foraging groups work both to find food and thereafter efficiently exploit or retrieve it and therefore may need to adapt their search strategies in complex environments in order to attain these goals. Previous work has focused on the simultaneous appearance of food and risk. Further, little research has been done to describe whether or not individual foragers mark their paths as they explore their environment and if they do, how this information affects subsequent foragers as they encounter the marks. Here, I examine the path choice by Argentine ants (*Linepithema humile*) in novel and complex environments. In Chapter 1, I tested if solitary exploring workers leave behind marks as they

explore novel areas and if this affects the direction subsequent individuals travel. The resulting pattern does not significantly differ from random choice, as workers are neither more nor less likely to choose the path taken by a previous ant. The time between decisions also does not predict the path chosen. This suggests that *L. humile* is more expansive in initially distributing its workers across novel ground. In Chapter 2, I examined whether colonies learn and respond to negative features of their environments and how this affects their path choice over time. *L. humile* workers respond differently depending on the type of risk present where they patrol/defend more in the presence of live competitors but decrease foraging in response to just a chemical cue.

The thesis of Emily Lessig is approved.

Noa Pinter-Wollman

Gregory F. Grether

Peter Nicholas Nonacs, Committee Chair

University of California, Los Angeles

2019

Table of Contents

Abstract ii

Table of Contentsv

List of Tablesvii

List of Figuresviii

Acknowledgementsix

Chapter 1. Exploratory behavior of Argentine Ants (*Linepithema humile*) encountering novel areas

Abstract 1

Introduction2

Methods3

Results5

Discussion5

Figures7

Literature Cited10

Chapter 2. Foraging behavior across paths that vary in risk cues and frequency of occurrence

Abstract12

Introduction13

Methods15

Results18

Discussion19

Figures22

Supplementary Materials	26
Literature Cited	28

List of Tables

Table 1-1. Ant decisions. Chi-square values equal 1.44, 0.49, and 0.21 for the First, Second, or All decisions (df = 1 for all tests; all values non-significant). For a 2x2 table-wide comparison, $G = 1.80$, df = 1 and non-significant.....	9
Table 2-1. Model comparison from time data collected across days.....	21

List of Figures

- Figure 1-1.** Experimental Setup: Start cup (C1) is located in the middle of the bottom row. Ants are then allowed to follow a course of their choice, making a series of two right vs. left decisions before finishing in one of four end cups (C5, C6, C7, or C8). The entire grid was assembled on white paper so as to ensure background consistency.....7
- Figure 1-2.** The values and logistic regression between decision time in minutes for the same (= 1) or different (= 0) path choice at the first fork (X's, black line), second fork (gray circles, gray line), or combined total fork decisions (dotted line)8
- Figure 2-1.** Diagram of the experimental grid. Cells labeled 1-4 indicated where negative stimuli were placed. All paths were of equal length.....22
- Figure 2-2.** Mean time to find food across days for all four paths by stimulus type (where colony 1-3 received FA and 4-6 received LO). There was a significant difference in time to find food between colonies that received FA as opposed to colonies that received LO ($p < 0.0001$).....23
- Figure 2-3.** Mean number of ants per day averaged across the number of observations per day (log-transformed) against risk cue (LO or FA), path traveled (0, 25, 50 or 100), number of paths with risk cues on a given day (i.e. “danger,” ranging from 1-3), and day of the experiment (divided into quartiles). Colors respond to the division of days into 4 quartiles.....24

Acknowledgements

I would like to thank my advisor, Dr. Peter Nonacs, for taking me on as a graduate student and allowing me to continue asking questions and conducting research. Working in his lab has allowed me to grow as a scientist and gain confidence in the research that I do. I am also thankful for my committee members, Drs. Noa Pinter-Wollman and Greg Grether, who offered guidance in my statistical analysis and helped me to think and write with clarity.

I would also like to thank my undergraduate assistants for volunteering their time throughout my project, especially Alejandro Martinez for collecting and maintaining ants and Daniel Rivas and Aneri Patel for helping run trials and scoring videos. I am grateful also for Descanso Gardens for allowing me to collect ant specimens and collaborating with me on numerous unique ant-related projects.

I am grateful for the support of my lab and the department. Thanks especially to Amanda Robin, as this research would not have been possible without her. Amanda coded webcams to create my “Antalyzer” which allowed me to take observations of my ants. I am forever grateful for her overall support as a lab mate and friend and cherish the time we got to work together at UCLA. Hayley Stansell first introduced me to UCLA and guided me along my journey here. I am grateful for her friendship and support throughout this journey. Julie Miller served as a mentor throughout my project from teaching me collecting techniques to answering method questions to lending supplies. I am thankful for her kindness and willingness to share her expertise.

Lastly, I would like to thank the people who first got me excited about science and conducting research. In particular, Dr. Mynhardt, who introduced me to entomology and how

fascinating studying insects can be; my parents, Tom and Leslie, for always encouraging me to pursue my dreams even if they took me far from home.

Chapter 1 is a version of Mahavni et al. 2019. Anika Mahavni contributed data and writing and editing for the manuscript. Emily K. Lessig contributed study design, facilitated data collection and writing and editing for the manuscript. Peter Nonacs also offered guidance in study design and data collection, statistical analysis and editing of the manuscript.

CHAPTER 1

Exploratory behavior of Argentine ants (*Linepithema humile*) encountering novel areas.

Abstract

When ants encounter a previously unexplored area, a variety of search patterns are possible. Previous studies have found that subsequent individuals tend to follow those that have gone before. This suggests that ants leave signature marks as they walk, independent of having encountered food, and that others recognize these and follow. This experiment tested if Argentine ants (*Linepithema humile*) also exhibit this pattern of behavior. Series of workers individually navigated through a grid consisting of two consecutive right versus left decisions. The resulting pattern does not significantly differ from random choice, as workers are neither more or less likely to choose the path taken by a previous ant. The time between decisions also does not predict the path chosen. Either *L. humile* does not mark as they walk, or workers do not use such marks in guiding their exploratory behavior. Under natural conditions, this suggests that *L. humile* is more expansive in initially distributing its workers across novel ground and less likely to have them clump together than species that do display following behavior. This has implications for how the relative effectiveness in finding versus exploiting and defending food sites might vary across species.

Introduction

Many species of ants mark their paths with pheromones that can inform and direct nestmates (Hölldobler and Wilson 1990). These pheromone trails often serve as a positive feedback mechanism for recruitment to recently found food sources (Jeanson et al. 2003, Sumpter and Beekman 2003, Jackson et al. 2006, Robinson et al. 2008, Dussutour et al 2009, Czaczkes et al. 2013). Less well-described is whether or not individual foragers marks their paths as they explore their environment, and if they do, how this information affects subsequent foragers that encounter the marks. For example, *Lasius pallitarsis* ants are significantly more likely to follow the same path as taken by a previous nestmate than an alternative path (Nonacs 1991). In *La. niger*, foragers use their own marks to reduce U-turns and therefore redundant search and the intensity of marks to indicate distance from the home site (Devigne and Detrain 2006). Workers generally prefer to explore areas saturated with cuticular hydrocarbons (CHS's) from nestmates, although such a signal in *La. niger* appears to be species-specific rather than colony-specific (Devigne and deBiseau 2012). The tendency to follow can also be negatively affected by the shape of the path itself, where ants prefer to travel in straight lines rather than to turn, even when paths are identical in length (Yates and Nonacs 2016), and paths requiring sharper turns are less likely to be followed (Forster et al. 2014). Selecting a straighter route can help to maximize mass recruitment to a food source and decrease the chance of foraging ants getting lost along the way.

Other studies across a variety of species have found both avoidance and attraction to marks left behind by non-nestmates or competing species (Binz et al. 2014; Wuest and Menzel 2017). Avoidance can be advantageous for poorly-competing species (Binz et al. 2014), while

aggressive and dominant species may gain from eavesdropping on where other species might be finding food (Wuest and Menzel 2017). This study focuses on the highly aggressive and competitive Argentine ant. Groups of Argentine ants begin by exploring a novel area randomly but as the exploratory front advances, nestmates are recruited and a trail is formed to and from the nest. Recruitment trails are typically formed between two points, while exploratory trails have no set destination (Deneubourg et al. 1990). Similar to the Nonacs (1991) study, the goal of this work is to test if solitary exploring workers leave behind marks as they explore novel areas and if this affects the direction subsequent individuals travel.

Methods

Collection

Workers were collected from the Mildred Mathias Botanical Gardens at the University of California, Los Angeles. Colonies were housed in containers filled with molded Plaster of Paris and coated on the sides with Fluon to prevent ants from escaping. Food and water was provided *ad libitum*, but food was not present during experimental trials.

Experiment

The experimental grid consisted of 8 cells (diameter 4.5 cm) connected via clear tubing (Fig. 1). At the start of each trial, 25 ants were introduced in the grid at C1. When the first ant entered the tube connecting C1 to C2, time was recorded and the tube was immediately clamped behind it after passing the halfway mark on the tube. After reaching C2, the time at which the ant entered either the tube going to C3 or C4 was recorded along with the direction chosen (L or R). This tube was clamped behind the ant after it passed the halfway mark, so as to prevent it from

returning to C2. The same procedure was conducted after the ant reached either C3 or C4. Each individual trial was complete when the ant reached an end cup (C5-8). The ant was then removed from the course with an aspirator and returned to its colony.

All paths were unclipped, and the next ant was allowed to move through the course using the same protocol. Most sets of trials ended after the fourth ant navigated the grids. In total, the experiment ran 34 sets of trials using 134 experimental ants. After a given set of time, the entire apparatus was cleaned with alcohol and dried to remove any marks left behind by the ants. One to three trial sets were run per day, and the time between trial days ranged from one to four days.

Data Analyses

The first ant encountered entirely unmarked areas. Hence, its choices were expected to be random. Each subsequent ant had a potential choice at C2 to either follow the previous ant or go the opposite direction if a mark is present and affecting path choice. Alternatively, the choice of direction could be random if not mark is present or marks have no effect on behavior (the experiment is unable to distinguish between these two possibilities). If an ant did go in a direction taken by a previous ant, then at C3 or C4 it faced a second similar choice of following the immediately preceding “leader”. However, it may not have a leader if its choice at C2 differs from all previous ants. Thus, all tested ants had a leader at C2, but the first ant going in the opposite direction would become the leader at C3 or C4 for the next decision point. Therefore, 100 ants had a leader at C2, but only 74 ants had a leader at C3 and C4. Chi-square tests at each decision point (1 and 2) determined whether or not the directions chosen deviated from a random choice. A 2x2 table-wide G-test determined if the decision pattern significantly differed across the two points.

Across trial, ants varied in the time they entered and exited through the grids, If ants mark their path with signals that dissipate over time, then a proclivity to follow or not may be negatively affected. In order to evaluate this, the effect of time between decisions was analyzed by logistic regression, with 1 for same decision or 1 for opposite direction. Time between when the last previous ant (defined as the leader) crossed the path and when the next ant reached the point was the independent variable. Times were not recorded for some trials, resulting in 88 measures at the first decision point and 65 at the second.

Results

Ants are neither significantly more or less likely to choose the path taken by the ant that immediately precedes it in the grid, either for the first or second decision or combined across both decision points (Table 1). There is no significant effect of increasing time between reaching a decision point (Fig. 2). The ants exhibited no turning bias in the experiment across the 4 cups that were the endpoints of the maze (Fig. 1: C5 = 33 ants reaching this cup; C6 = 29; C7 = 38; C8 = 34; $\chi^2 = 1.224$, $df = 3$, NS).

Discussion

Argentine ants, unlike *Lasius* species (Nonacs 1991, Devigne and Detrain 2006, Devigne and deBiseau 2012) do not tend to follow each other when encountering novel terrain. Rather, they seem to choose directions randomly, independent of the decisions of the ants before them. This could result from either workers not leaving recognizable marks behind as they walk, or other being indifferent to them. Earlier work on *L. humile* does not strongly suggest that paths

are marked and favored in the absence of any specific incentive (Aron et al. 1989). However, this result occurred when paths were being explored by large numbers of interacting workers and not by single workers navigating alone.

A proclivity to follow, such as exhibited by individuals in *Lasius* species, would result in more ants at the same place. Such a foraging strategy would not be the most systematic search pattern as not all areas might be effectively explored. However, it would have the advantage of concentrating foragers and should a food item be encountered, enable more rapid recruitment in number (Denton and Nonacs 2018). Over time, such a system has also been proposed to bias foraging effort into more rewarding locations and away from less rewarding ones (Devigne et al. 2006).

Therefore, a more random distribution of workers in space, as suggested here for *L. humile*, would more emphasize an initially broader search pattern by the first workers in a new area. In examining colony-level search patterns in this species, Denton and Nonacs (2018) found that area search was initially more evenly distributed over space when a newly-encountered area did not yield food rewards. However, when food rewards started to appear, searching foragers clumped in spatial locations that reduced food encounter rates, but likely increased recruitment to food when it was found. Overall, this suggests that *L. humile* colonies may employ a two-stage search process. At first, searching more expansively and then, once an area is scouted, concentrating on more rapid recruitment to specific sites that are more likely to produce positive payoffs.

Figures

Figure 1. Experimental Setup: Start cup (C1) is located in the middle of the bottom row. Ants are then allowed to follow a course of their choice, making a series of two right vs. left decisions before finishing in one of four end cups (C5, C6, C7, or C8). The entire grid was assembled on white paper so as to ensure background consistency.

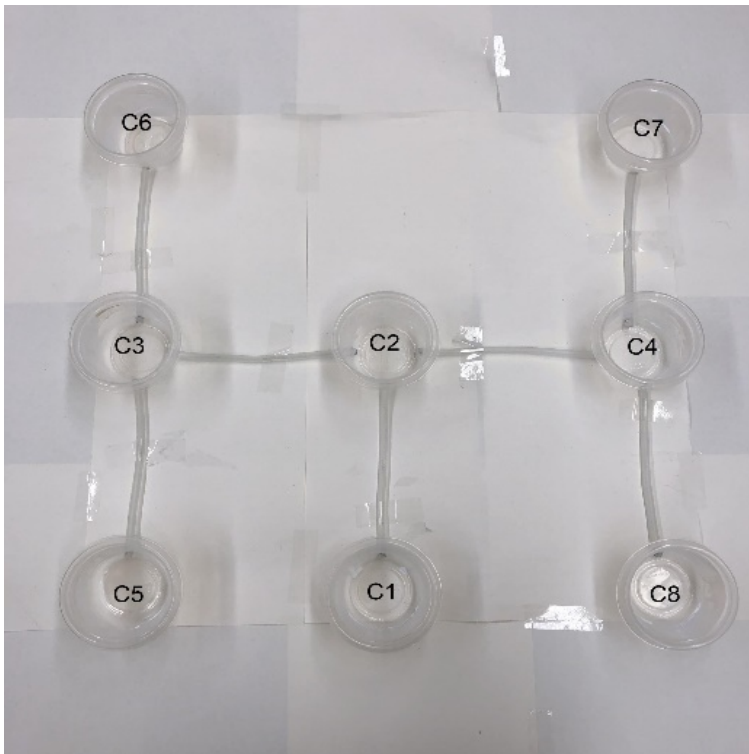


Figure 2. The values and logistic regression between decision time in minutes for the same (= 1) or different (= 0) path choice at the first fork (X's, black line), second fork (gray circles, gray line), or combined total fork decisions (dotted line).

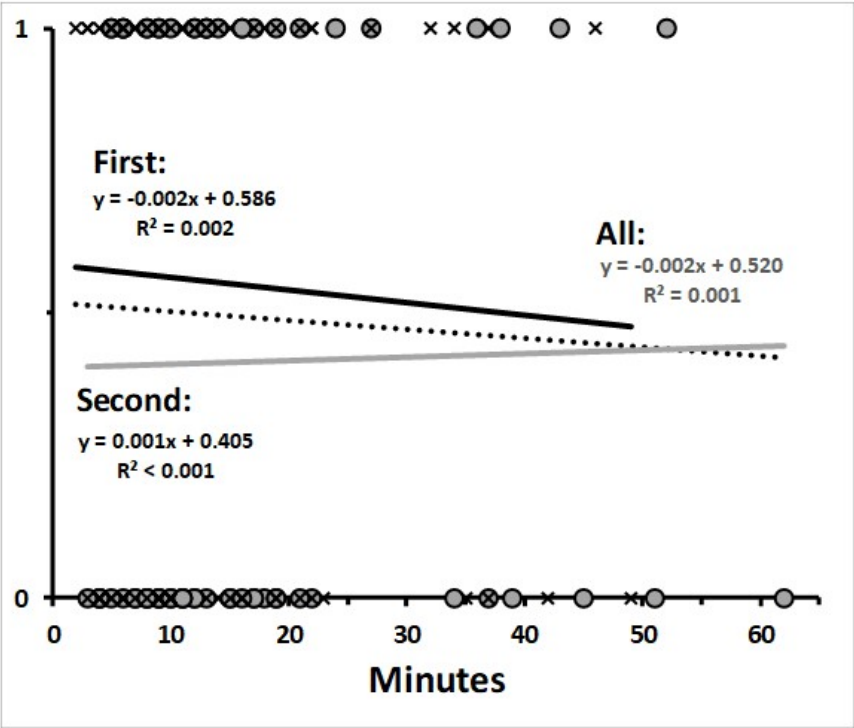


Table 1. Ant decisions. Chi-square values equal 1.44, 0.49, and 0.21 for the First, Second, or All decisions (df = 1 for all tests; all values non-significant, with $p > 0.1$). For a 2x2 table-wide comparison, $G = 1.80$, $df = 1$ and non-significant.

Follow?	First	Second	All
Yes	56	34	90
No	44	40	84
Total	100	74	174

Literature Cited

- Aron S, Pasteels JM, Deneubourg JL (1989). Trail-laying behaviour during exploratory recruitment in the Argentine ant, *Iridomyrmex humilis* (Mayr). *Biol Behav* 14:207–217.
- Binz H, Foitzik S, Staab F, Menzel F (2014) The chemistry of competition: exploitation of heterospecific cues depends on the dominance rank in the community. *Anim Behav* 94:45-53.
- Czaczkes TJ, Gruter C, Ellis L, Wood E, Ratnieks FLW (2013) Ant foraging on complex trails: route learning and the role of trail pheromones in *Lasius niger*. *J Exper Biol* 216:188-197.
- Deneubourg JL, Aron S, Goss S, Pasteels JM (1990) The self-organizing exploratory pattern of the Argentine ant. *J Ins Behav* 3:159-168.
- Denton KK, Nonacs P (2018) Habitat complexity and predictability effects on finding and collecting food when ants search as cooperative groups. *Anim Behav* 141:77-84.
- Devigne C, de Biseau JC (2012) The differential response of workers and queens of the ant *Lasius niger* to an environment marked by workers: Ants dislike the unknown. *Behav Proc* 91:275-281.
- Devigne C, Detrain C (2006) How does food distance influence foraging in the ant *Lasius niger*: the importance of home-range marking. *Insect Soc* 53:46-55.
- Devigne C, Renon AJ, Detrain C (2004) Out of sight but not out of mind: modulation of recruitment according to home range marking in ants. *Anim Behav* 67:1023-1029.
- Dussutour A, Nicolis SC, Shephard G, Beekman M, Sumpter DJT (2009) The role of multiple pheromones in food recruitment by ants. *J Exper Biol* 212:2337-2348.
- Evison SEF, Hart AG, Jackson ED (2008) Minor workers have a major role in the maintenance of leafcutter ant pheromone trails. *Anim Behav* 75:963-969.
- Forster A, Czaczkes TJ, Warner E, Woodall T, Martin E, Ratnieks FLW (2014) Effect of trail bifurcation asymmetry and pheromone presence or absence on trail choice by *Lasius niger* ants. *Ethology* 120:768-775.
- Jackson DE, Martin SJ, Holcombe M, Ratnieks FLW (2006) Longevity and detection of persistent foraging trails in Pharaoh's ants, *Monomorium pharaonis* (L.). *Anim Behav* 71:351-359.

- Jeanson R, Ratnieks FLW, Deneubourg JL (2003) Pheromone trail decay rates on different substrates in the Pharaoh's ant, *Monomorium pharaonis*. *Physiol Entomol* 28:192-198.
- Nonacs P (1991) Exploratory behavior of *Lasius pallitarsis* ants encountering novel areas. *Insect Soc* 38:345-349.
- Robinson EJH, Green KE, Jenner EA, Holcombe M, Ratnieks FLW (2008) Decay rates of attractive and repellent pheromones in an ant foraging trail network. *Insect Soc* 55:246-251.
- Sumpter DJT, Beekman M (2003) From nonlinearity to optimality: pheromone trail foraging by ants. *Anim Behav.* 66:273-280.
- Wuest M, Menzel F (2017) I smell where you walked - how chemical cues influence movement decisions in ants. *Oikos* 126:149-160.
- Yates AA, Nonacs P (2016) Preference for straight-line paths in recruitment trail formation of the Argentine ant, *Linepithema humile*. *Insect Soc* 63:501-505.

CHAPTER 2

Foraging behavior across paths that vary in risk cues and frequency of occurrence.

Abstract

Cooperatively foraging species often adjust their search strategies in complex environments to efficiently find and exploit food sources. These strategies become more complicated when food and risk can be simultaneously present and when they differ in predictability. For example, there may be multiple paths to reach a foraging site that vary in risk. This study examines how colonies of Argentine ants (*Linepithema humile*) respond to such a situation where identical-length paths differ in how they present risk. The risk cues are either a live competitor (velvety tree ants, *Liometopum occidentale* (LO)) or formic acid (FA), a defensive chemical commonly associated with formicine ant species. Across four paths to food, the presence of cues also varied from always to never present. Although the non-risky path was used more often, in no case did colonies completely avoid the paths with risk. Overall, more *L. humile* workers explored paths associated with LO than with FA. This had a significant impact on foraging ability where LO colonies were faster at finding food than FA colonies. Further, *L. humile* workers' response to FA was similar over time while declined for LO, suggesting a 'dear enemy' habituation and reduction in aggression over time. Thus, it appears that *L. humile* foragers categorize risk cues and will vary their responses in potentially effective ways.

Introduction

Efficient food acquisition challenges all animals. Foragers navigate complex environments that present a variety of costs and benefits, often associated both with the distribution or quality of food patches and with the travel associated to reach them. Patch exploitation and movements between patches can be influenced by travel distance, presence of competing species or cooperating conspecifics, food quality, and hunger levels (Alatalo and Lundberg 1986; Anderson 1984; Beckers et al. 1990; Conradt and Roper 2005; Vittori et al. 2006; Ronconi and Burger 2011; Yates and Nonacs 2016; Yamada 2017). Given that a forager's first goal is to effectively encounter potential food items, the time it takes to discover food has implications for foraging success and efficiency (Beverly 2009). In heterogeneous environments, organisms may need to change their search strategies in order to efficiently find food. For example, Argentine ants (*Linepithema humile*) appear to prioritize rapid recruitment to food once it is found, rather than maximizing food encounter rates when search areas differ in spatial complexity and food appears ephemerally (Denton and Nonacs, 2018).

Foraging strategies and spatial distributions of workers can also vary in response to the risk that is present along a path. *Lasius pallatarsis* ant colonies abandon patches with associated mortality risk to forage at "safer" patches (Nonacs and Dill, 1988). In *Formica* ants (*F. perpilosa* and *F. integroides*), smaller foragers avoid sites at which risk is present and larger foragers may spend more time at these sites in a defensive posture (Kay and Rissing, 2005; Tanner, 2008). Maximizing foraging effort therefore requires recognizing key properties of different patches and allocating foraging efforts across patches in a manner that increases time spent at "good" patches

and decreases time spent at “bad.” In heterogenous or changing landscapes, animals may then need to sample their environment and develop expectations about future encounters to efficiently exploit the area (Stephens and Krebs 1986). This process of combining older information with newer information to alter expectations is known as Bayesian updating (Valone, 2006) and has been observed in a wide variety of taxonomic groups (e.g. Lima 1984, 1985; Valone & Brown 1989; Valone 1991, 1992; Krebs & Inman 1992; Alonso et al. 1995; Olsson et al. 1999; van Gils et al. 2003; Stamps et al. 2018).

Cooperatively foraging species, such as ants, add two more aspects to foraging strategies. First, individual workers are ‘disposable’ in the sense that their deaths are expected and do not directly influence the reproduction of the colony. Such foragers gain indirect fitness by helping kin reproduce and may behave very differently from animals that risk their own reproductive success (Nonacs and Dill, 1990). Second, many ant species mark and maintain trails that can denote the optimal route to a food source; balancing trade-offs between various qualities amongst trails (Nonacs and Dill, 1990,1991, Latty et al. 2017).

In this study, we examine the path choice that ants make over time when: (1) There are multiple paths of equal length to a food source, and (2) negative stimuli (i.e., cues that a competing species may be nearby) are present along these paths in different frequencies, ranging from 0% (never present) to 100% (always present). Previous studies have focused on decisions between food and risk that are presented simultaneously, but here we focus on how colonies learn and respond over time to negative features of their environment that may vary in how often and predictably they are present.

Methods

Collection

We setup six replicate colonies of *L. humile* containing approximately 500-800 workers, 6-8 queens, and brood collected at the University of California, Los Angeles. The ants nested in open plastic containers, filled with molded plaster of Paris, which was kept moist for nest humidity, and coated on the sides with Fluon. Water was provided *ad libitum*.

Experiment

The experimental arenas consisted of four equidistant paths made of clear, plastic tubing to a single foraging arena (distance = 90cm). To measure the time it took ants to find food, we added sugar water to the foraging arena daily between approximately 12:00 and 15:00 h. Time was recorded as how long it took the first ant to find the food and was recorded up to the first 10 minutes. Food was removed two hours after placement (whether or not ants were foraging) to avoid colonies becoming satiated. To present the ants with negative stimuli, we placed cells at the midway point of each path where cues were added (Fig. 2-1). Workers from three colonies each encountered either several workers of the aggressive velvety tree ant (*Liometopum occidentale* (LO)) (Hoey-Chamberlain et al. 2013) or formic acid (FA) on the paths and depending on the treatment. LO and FA were placed along paths between 8:00 and 11:00 h. daily. They were present along paths for approximately 24 hours and not correlated with the appearance of food. The four paths differed such that LO or FA were present 0, 25, 50 or 100% of trials. Therefore, to reach food the *L. humile* foragers always had one path that never had risk associated, another that always had risk, and two that might or might not have had risk associated with them. This means that for any given trial, 1-3 paths could have had LO or FA present.

We collected LO from Descanso Gardens (Los Angeles, CA) and housed them in a similar manner to that of *L. humile*. Both *L. humile* and *L. occidentale* are in the same subfamily of Dolichoderinae. FA is the primary defensive chemical used by ant species in the subfamily Formicinae (Hefetz & Blum 1978; Blum 1978). Colonies of ants were counted every four to five days to ensure numbers did not drop drastically and workers were added if numbers dropped below 500. Trials lasted approximately 20 days to allow each colony to experience all frequencies of negative stimuli at least four times.

Risk cue: LO

We placed approximately 15-20 LO workers in cells 1-4 (Fig. 2-1). Mesh was present between the LO workers and the *L. humile* workers, to prevent physical contact (Fig. S2). However, they were able to exchange chemical cues. LO workers were removed and replaced daily to ensure their numbers and effect were consistent over the course of the experiment.

Risk cue: FA

We placed cotton pads with 100 μ L of FA (Walmart, USA) in cells 1-4 (Fig. 2-1). We covered the cells with lids for these trials to prevent the FA from completely dissipating over time but cut holes (diameter =2.54cm) into the encounter chambers for aeration and to prevent FA fumes from accumulating in deadly amounts. We removed pads and/or replaced them daily depending on the presentation schedule. If FA was to be absent for the day, we cleaned cells to remove any leftover traces.

Video Observations

We programmed webcams to take fifteen-minute recordings of path use, approximately three times per day: one at least three hours prior to food presentation (morning observation), one while food was present (afternoon observation), and one in the evening, at least three hours after food had been removed (evening observation). The number of observations across colonies varied because they were not all set up simultaneously and some observations had to be discarded due to ants escaping or the clarity of video taken. Observations were at minimum 3 hours apart and at approximately the same time daily. We later scored these videos to examine path use over time.

Video Scoring

Only the number of ants crossing the shaded portion of the experimental grids (Fig. 2-1) was recorded as this was most detectable across all cameras and colonies. To avoid double-counting an ant on a path, we kept spatial track of a forager's location and did not count it more than once when simply moving back and forth on the same path.

Statistics

We conducted linear-mixed effect models to study the amount of time ants took to find food across days. Stimulus type was included in the model as well, as *L. humile* workers could respond differently to LO or FA affecting finding times. We began by generating a null model containing colony as a random intercept, then iteratively incorporated fixed effects into the model, in order to find which combination of fixed effects generated the lowest AIC value for the model. We conducted an ANOVA to examine mean time to find food across days by stimulus type (FA or LO), all in R version 3.5.2 (R Core Team 2018).

We also conducted an ANOVA to examine path use of ants per day averaged across the number of observations per day (log-transformed) against risk cue (LO or FA), path traveled (0, 25, 50 or 100), number of paths with risk cues on a given day (i.e. “danger,” ranging from 1-3), and day of the experiment. Not every colony was tested on every day and therefore for statistical analysis, time was divided into quartiles of 6 days each.

Results

In the analysis examining time to find food across days, including stimulus type as a fixed effect significantly improved model fit (Table 2-1). This indicates that type of stimulus predicts the amount of time it takes ants to find food. Further, in the analysis examining mean time to find food across days by stimulus type, food was found significantly faster in LO treatments than in FA treatments (Fig. 2-2; $p < 0.0001$). This indicates that foraging behavior (i.e. time to find food) is influenced by stimulus type.

In the analysis examining number of ants per day against risk cue, path traveled, danger (in terms of number of paths with risk cues present on a given day), and day of the experiment, danger had neither a significant main effect nor any significant interaction effects. Therefore, it was dropped as a factor in the ANOVA. In regard to the other main effects, significantly more ants were on the paths with LO as opposed to FA (Fig. 2-3; $p < 0.0001$, $F = 154.87$, $df = 1$). The frequency of risk appearing on the paths also significantly affected their use (Fig. 2-3; $p < 0.0004$, $F = 6.205$, $df = 3$), where there was a significant difference between the number of ants using the 0 and 25% paths ($p < 0.0061$); the 0 and 50% paths ($p < 0.0001$); and the 50 and 100% paths ($p < 0.0061$). Additionally, the day of the experiment had a significant affect, where the overall number of ants declined over time (Fig. 2-3; $p < 0.0001$; $F = 9.681$; $df = 3$).

When examining the 2-way interactions, we found when FA is the stimulus the number of ants declined as risk got more frequent. However, for LO, the consistently riskiest path (100%) drew the most ants (Fig. 2-3; $p < 0.0002$, $F = 6.639$, $df = 3$). Further, the FA response was similar over time, while the LO response consistently declined (Fig. 2-3; $p < 0.0001$, $F = 10.130$, $df = 3$). Except for the 100% path, the number of ants declined over time (Fig. 2-3: $p < 0.0026$; $F = 2.872$, $df = 9$). The 3-way interaction between stimulus, path, and time was not significant.

Discussion

Cooperatively foraging species are a model system to examine how individuals with limited knowledge and cognitive capabilities can achieve complex goals such as navigating complex environments. Patch exploitation and movements between patches can be influenced by a variety of factors including presence of competing species. While a forager's goal is to successfully find food items, it also has to balance safety along paths. In heterogenous environments, organisms may need to change their search strategies in order to efficiently find food as well stay safe. Previous studies have focused on decisions between food and risk that are presented simultaneously, but here we focus on how colonies learn and respond to negative features of their environment that may vary in how often and how predictably they are present.

Different types of risk can also elicit different responses. In our experiments, we found the two types of cues about risk drew significantly different responses both in regard to foraging behavior and path use. Ants patrol/defend areas more intensely when in the presence of live LO workers than with only FA, resulting in more ants on paths with LO as compared to FA. We can

attribute the lower numbers on paths with FA due to the effectiveness of this chemical weapon.

FA as a defense in formicine species is more effective in conflicts with Argentine ants than the defenses of the dolichoderine velvety tree ants. This has been noted in *Nylanderia fulva* and other formicines where formic acid as a chemical weapon achieves competitive dominance in combat with *Solenopsis invicta* (LeBrun et al. 2014). Thus, even the occasional presence of FA may depress Argentine ant activity in that area or path. As FA is a particularly effective chemical weapon in combat, learning which trails to use or avoid FA is an important aspect of their foraging strategy.

Further, the large numerical difference in response along paths with live LO workers as compared to FA likely resulted in LO colonies finding food when it appeared significantly faster than FA colonies. This suggests that food discovery rate by Argentine ants can be directly affected by their responses to encountering different competitive species. Additionally, their response to FA is similar over time while for LO, their response declines over time. This suggests that the lack of direct contact and fights results in habituation and a ‘dear enemy’ reduction in aggression over time (Langen et al. 2000) as *L. humile* workers decline in their response to LO over time.

Additionally, the predictability of the risk along paths also had a significant effect on path use. The always safe (0%) path is used more overall than the paths with risk. Further, the overall number of ants on the paths declines over time, except for the never safe (100%) path. This suggests that ants can assess and respond to risk present along paths and they moderate their responses to paths that are not consistently risky but maintain a more consistent presence when in response to paths that are always risky.

This work demonstrates that Argentine ants are able to learn about their environment and use this information to effectively navigate and efficiently exploit their environment. Further, this work elucidates how individual ants (with local knowledge and the ability to interact) can scale up to effective organizations that optimally achieve vital, but potentially competing, objectives. This is due to the fact that learning at the colony level is often more effective than at the individual level as workers can vary in age and experience, and colonies allow for communication and an increase in numbers. These results are relevant to understanding how groups coordinate and function in a wide variety of species.

Table 2-1: Model comparison for time data across days. Model m2

was chosen based on AIC value and comparison with other models using likelihood ratio tests.

Model Name	AIC	Model Formula	Comparison	Results	P
m0	268.7	$\ln_Time \sim (1 Colony)$	N/A	N/A	N/A
m1	269.0	$\ln_Time \sim Day + (1 Colony)$	m1, m0	not significant	0.1919
m2	260.4	$\ln_Time \sim Day + Stimulus + (1 Colony)$	m2, m1	significant	0.0011

Figure 2-1. Diagram of the experimental grid. Cells labeled 1-4 indicated where negative stimuli were placed. All paths were of equal length.

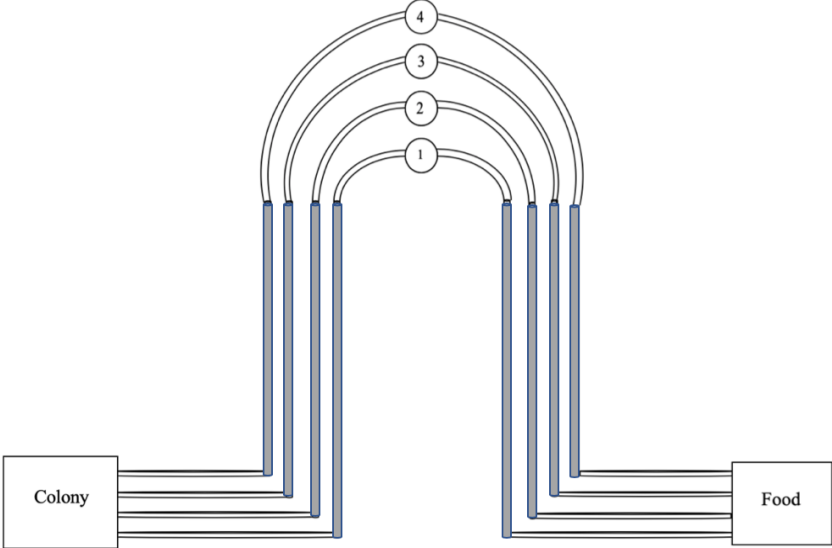


Fig. 2-2: Mean time to find food across days for all four paths by stimulus type (where colony 1-3 received FA and 4-6 received LO). There was a significant difference in time to find food between colonies that received FA as opposed to colonies that received LO ($p < 0.0001$).

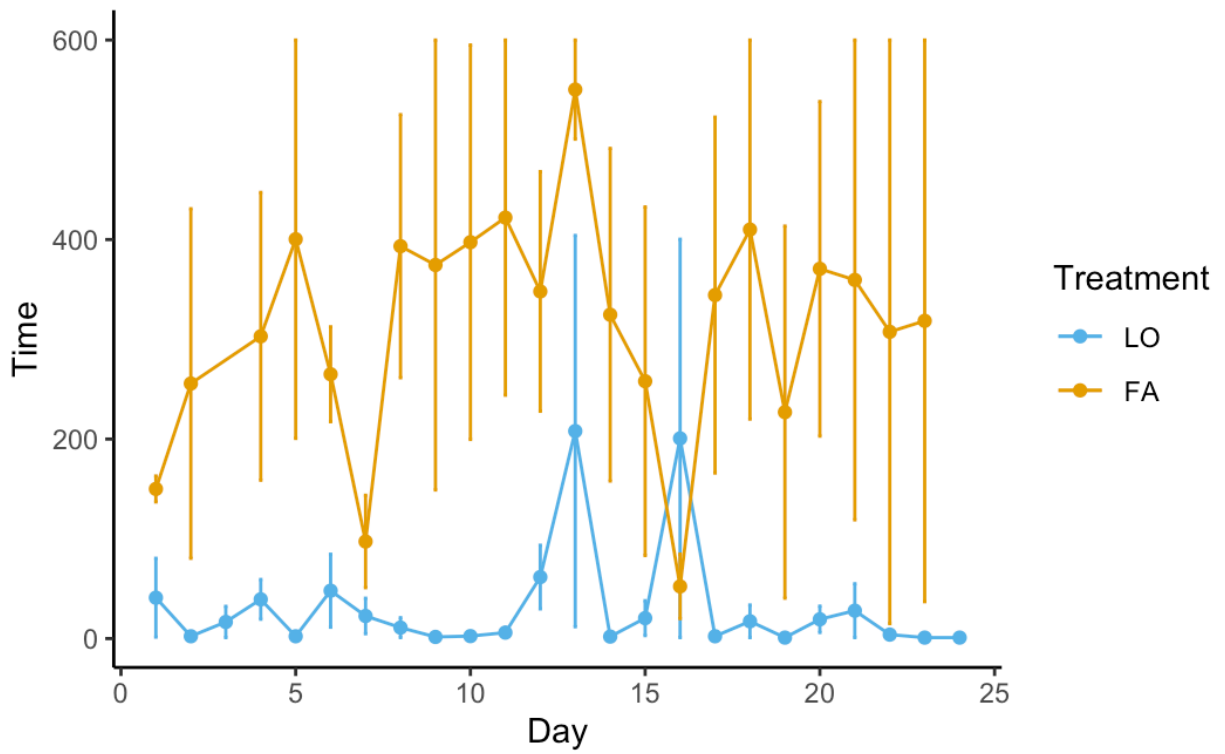
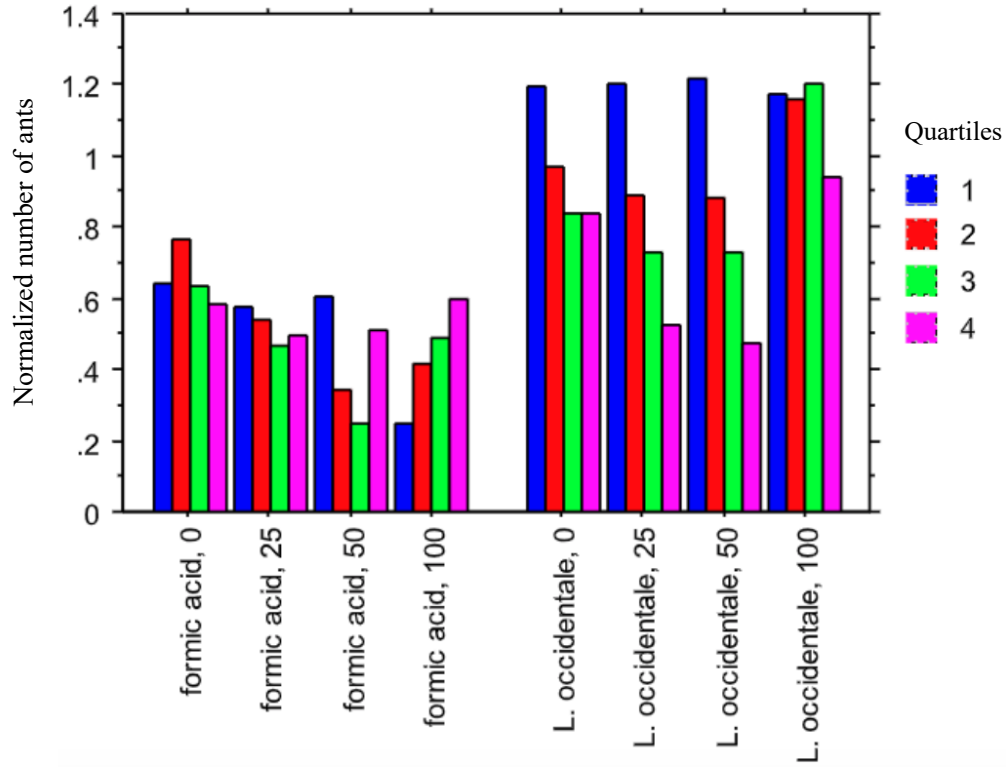


Fig. 2-3: Mean number of ants per day averaged across the number of observations per day (log-transformed) against risk cue (LO or FA), path traveled (0, 25, 50 or 100), number of paths with risk cues on a given day (i.e. “danger,” ranging from 1-3), and day of the experiment (divided into quartiles). Colors respond to the division of days into 4 quartiles.



Supplementary Information



Fig. S1: Picture of experimental setup, including paths that were scored and cells where stimuli were placed.

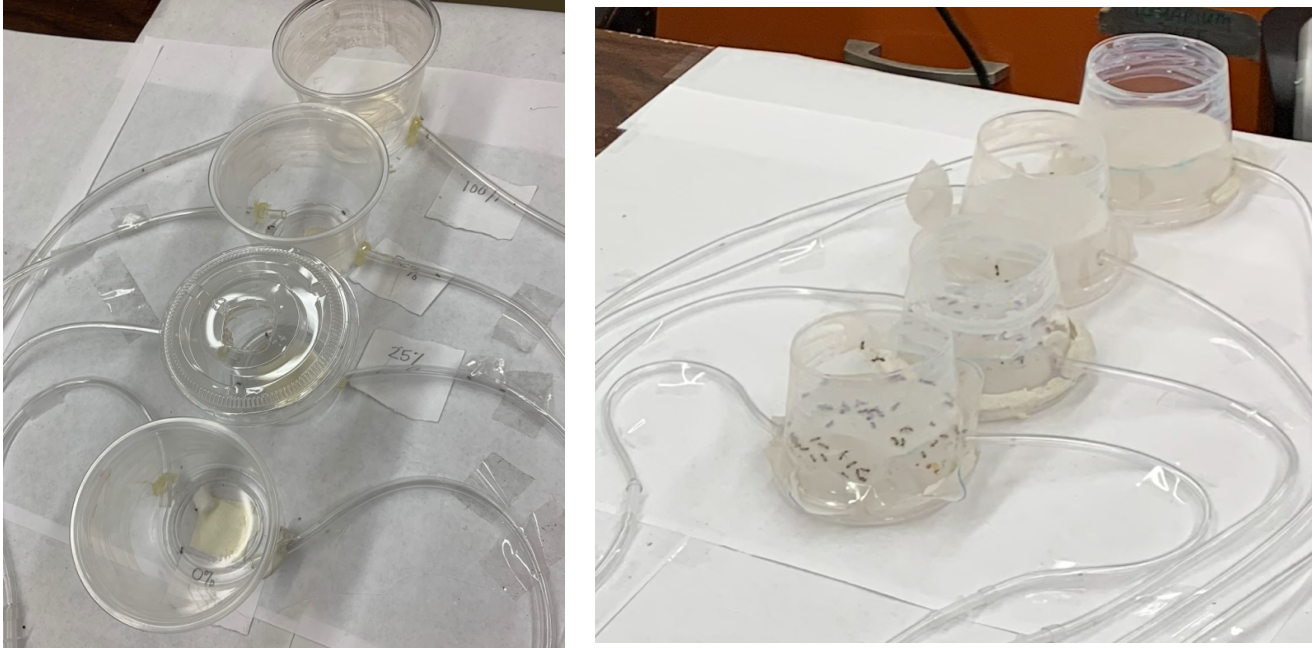


Fig. S2: Picture of cells where stimuli were placed along experimental grids. Left represents trials where formic acid was used; highlighting inside of the cell where formic acid was placed as well as the lidding of cells to prevent formic acid from dissipating over time. Right represents trials where *L. occidentale* was used. Mesh prevented *L. humile* and *L. occidentale* from coming into physical contact, but pheromones could be exchanged.

Literature Cited

- Alonso, J.C., Alonso, J.A., Bautista, L.M., & Muñoz-Pulido, R (1995). Patch use in cranes: A field test of optimal foraging predictions. *Animal Behaviour*, 49, 1367-1379.
- Alatalo, R. V. and Lundberg, A. (1986). Interspecific competition and niche shifts in tits (*Parus* spp.): evaluation of nonexperimental data. *Am. Nat.*, 127: 335-834.
- Anderson, J.R. (1984). Ethology and ecology of sleep in monkeys and apes. *Adv. Stud. Behav.*, 14:165–230.
- Beckers, R., Deneubourg, J. L., Goss, S., Pasteels, J. M. (1990). Collective decision making through food recruitment. *Insectes Sociaux*, 37(3): 258-267.
- Beverly, B.D., McLendon, H., Nacu, S., Holmes, S., Gordon, D.M. (2009). How site fidelity leads to individual differences in the foraging activity of harvester ants. *Behavioral Ecology*. 20 (3), 633-638.
- Blum, M. S. (1978). Biochemical defenses of insects. In *Biochemistry of insects* (ed. M. Rockstein). New York: Academic Press, 465-513.
- Conradt, L. & Roper, T. (2005) Consensus decision making in animals. *Trends in ecology and evolution*. 20, 449-56.
- Denton, K. and Nonacs, P. (2018). Habitat complexity and predictability effects on finding and collecting food when ants search as cooperative groups. *Animal Behaviour*. 141:77-84.
- van Gils, J. A., Schenk, I. W., Bos, O., Piersma, T., & Moore, A. J. (2003). Incompletely informed shorebirds that face a digestive constraint maximize net energy gain when exploiting patches. *American Naturalist*, 161, 777-793.
- Hoey-Chamberlain, R., Rust, M., and Klotz, J. (2013) A review of the biology, ecology and behavior of velvety tree ants of North America. *Sociobiology*, 60 (1), 1-10.
- Kay, A., & Rissing, S.W. (2005). Division of foraging labor in ants can mediate demands for food and safety. *Behavioral Ecology and Sociobiology*, 58, 165-174.
- Krebs, J. R. & Inman, A. J. (1992). Learning and foraging: individuals, groups, and populations. *Am. Nat.*, 140, S63–S84.
- Langen, T., Tripet, F., and Nonacs, P. (2000). The red and black: habituation and the dear-enemy phenomenon in two desert *Pheidole* ants. *Behav Ecol Sociobiol*, 48: 285-292.

- Latty, T., Holmes, M.J., Makinson, J.C., Beekman, M. (2017). Argentine ants (*Linepithema humile*) use adaptive transportation networks to track changes in resource quality. *Journal of Experimental Biology*, 220: 686-694.
- LeBrun, E., Jones, N, and Gilbert, L. (2014). Chemical warfare among invaders: A detoxification interaction facilitates an ant invasion. *Science*, 343: 1014-1017.
- Lima, S. L. (1984). Downy woodpecker foraging behaviour: Efficient sampling in simple stochastic environments. *Ecology*, 65, 166-174.
- Lima, S. L. (1985). Sampling behaviour of starlings foraging in simple patchy environments. *Behavioral Ecology and Sociobiology*, 16, 135-142.
- Nonacs, P., Dill, L.M. (1988). Foraging response of the ant *Lasius pallitarsis* to food sources with associated mortality risk. *Insectes Sociaux*, 35 (3): 293-303.
- Nonacs, P. and Dill, L. M. (1990), Mortality Risk vs. Food Quality Trade-Offs in a Common Currency: Ant Patch Preferences. *Ecology*, 71: 1886-1892.
- Nonacs, P. and Dill, L.M. (1991). Mortality risk versus food quality trade-offs in ants: patch use over time. *Ecological Entomology*, 16: 73-80.
- Olsson, O., Wiktander, U., Holmgren, N. M. A., & Nilsson, S. G. (1999). Gaining ecological information about Bayesian foragers through their behaviour. II: A field test with woodpeckers. *Oikos*, 87, 264-276.
- R Core Team (2017). R: A Language and Environment for Statistical Computing. [Online.] Available at <https://www.r-project.org/>.
- Ronconi, R.A., Burger, A.E. (2011). Foraging space as a limited resource: inter-and intra-specific competition among sympatric pursuit-diving seabirds. *Canadian Journal of Zoology*, 89 (4): 356-368.
- Stephens, D. W., and J. R. Krebs. (1986). *Foraging theory*. Princeton University Press, Princeton, New Jersey, USA.
- Stamps, J. A., Biro, P. A., Mitchell, D. J. and Saltz, J. B. (2018), Bayesian updating during development predicts genotypic differences in plasticity. *Evolution*, 72: 2167-2180.
- Tanner, C.J. (2008) Resource characteristics and competition affect colony and individual foraging strategies of the wood ant *Formica integroides*. *Ecological Entomology*, 33: 127-136.
- Valone, T. J., & Brown, J. S. (1989). Measuring patch assessment abilities of desert granivores. *Ecology*, 70: 1800-1810

- Valone, T.J. (1991) Bayesian and prescient assessment: Foraging with pre-harvest information. *Animal Behaviour*, 41: 569-577.
- Valone, T.J. (1992). Information for patch assessment: A field investigation with black-chinned hummingbirds. *Behavioral Ecology*, 3: 211-222.
- Valone, T.J. (2006). Are animals capable of Bayesian updating? An empirical review. *Oikos*, 112: 252-259.
- Vittori, K., Talbot, G., Gautrais, J., Fourcassie, V., Araujo, A., Theraulaz, G. (2006). Path efficiency of ant foraging trails in an artificial network. *Journal of Theoretical Biology*, 239: 507-515.
- Yamada, H. (2017). Hunger enhances consistent economic choices in non-human primates. *Scientific Reports*. 7, 2394.
- Yates, A.A., Nonacs, P. (2016). Preference for straight-line paths in recruitment trail formation of the Argentine ant, *Linepithema humile*. *Insectes Sociaux*, 63 (4): 501-505.