Individual error correction drives responsive self-assembly of army ant scaffolds

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An inherent strength of evolved collective systems is their ability to rapidly adapt to dynamic environmental conditions, offering resilience in the face of disruption. This is thought to arise when individual sensory inputs are filtered through local interactions, producing an adaptive response at the group level. To understand how simple rules encoded at the individual level can lead to the emergence of robust group-level (or distributed) control, we examined structures we call “scaffolds,” self-assembled by Eciton burchelli army ants on inclined surfaces that aid travel during foraging and migration. We conducted field experiments with wild E. burchelli colonies, manipulating the slope over which ants traversed, to examine the formation of scaffolds and their effects on foraging traffic. Our results show that scaffolds regularly form on inclined surfaces and that they reduce losses of foragers and prey, by reducing slipping and/or falling of ants, thus facilitating traffic flow. We describe the relative effects of environmental geometry and traffic on their growth and present a theoretical model to examine how the individual behaviors underlying scaffold formation drive group-level effects. Our model describes scaffold growth as a control response at the collective level that can emerge from individual error correction, requiring no complex communication among ants. We show that this model captures the dynamics observed in our experiments and is able to predict the growth—and final size—of scaffolds, and we show how the analytical solution allows for estimation of these dynamics.

To achieve resilience, biological systems at many scales, including these social insect infrastructures, utilize various mechanisms of feedback control. In a recent example, a core regulatory feedback mechanism common to many social insects was identified—the so-called common stomach—which achieves resilience through a saturation process that relies on negative feedback through integral control (12). Both positive and negative feedback have long been known to underlie self-organizing processes like the formation of ant foraging trails and collective decision making (13–15), and negative feedback has been shown to play a particularly important role among social insects in maintaining flexibility when environmental conditions change (16–18). Understanding how feedback control operates in distributed systems, like ant colonies, to facilitate resilience remains an important challenge, with implications across disciplines.

The army ant Eciton burchelli, with large colonies of more than 500,000 workers and a distinctive nomadic life cycle (19), presents an ideal model for studying resilience and distributed control in natural systems. E. burchelli is considered a top predator in the neotropical forest leaf litter community (20) and a canonical example of self-organization among social insects (21).

Significance

Human-designed infrastructures and networks relying on centralized or hierarchical control are susceptible to single-point catastrophic failure when disrupted. By contrast, most complex biological systems employ distributed control and can be more robust to perturbations. In field experiments with Eciton burchelli army ants, we show that scaffold structures, self-assembled by living ants, emerge in response to disrupted traffic on inclines, facilitating traffic flow and stemming losses of foragers and prey. Inferred by our observations, we present a theoretical model based on proportional control and negative feedback, which may be relevant to many distributed systems in which group-level properties can be modified through individual error sensing and correction. The mechanism is simple, and ants only require information about their individual state.

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Unlike most ants that maintain a fixed nest location, colonies alternate every few weeks between a stationary and a nomadic phase, during which the entire colony moves nightly to a new temporary nest site, known as a bivouac (22, 23). Most workers spend the day foraging in massive swarm raids that start at dawn, combing the forest floor to flush out a diverse array of arthropod prey from the leaf litter (24, 25). At the front of the raid, fleeing prey items are attacked en masse, with those caught dismantled for transport along a branching trail network that can stretch for over 100 m back to the bivouac (26). Raids must be conducted at a rapid pace to maintain the high rates of prey delivery needed to support the ravenous appetite of the developing brood (27). This evolutionary pressure has led to a number of morphological and behavioral adaptations, including the spontaneous formation of traffic lanes that increase the efficiency of foraging trails (10), some of the highest running speeds among all ants (28), and a specialized porter caste morphology to facilitate the transport of bulky prey items, carried under the body (29).

This suite of adaptations includes the remarkable capacity for self-assembly, by which ants can join together to create temporary structures that modify the environment (27, 30, 31). Various examples of self-assembly have been observed in other social insects (32, 33), and these structures have been described as intermediate-level parts of insect societies—adaptive units that function at a level between individual and colony (34). For E. burchellii and the closely related Eciton hamatum, self-assembled structures serve two primary functions: for shelter, in the case of the bivouac, and to facilitate the flow of traffic during raids and emigrations, with structures like plugs (27) and bridges (30, 31) that emerge along the trail network as needed. Our previous work on Eciton bridges has shown how these structures are robust to perturbations and responsive to traffic and environmental geometry, changing size and location to create shortcuts that benefit the colony.

Here we reveal another type of self-assemble in E. burchellii which we call “scaffolds,” given their function as temporary support structures. These structures have not been previously studied experimentally or clearly defined, although they have been described based on observations in previous literature without consistent terminology (35, 36). Scaffolds often form when the foraging trail crosses a sloped surface from which ants may slip and/or fall, such as a rock face, tree root, or even the wall of a building (as shown in SI Appendix, Movie S1). To form a scaffold, individual ants stop and grip the underlying surface with their tarsi, remaining stationary as traffic continues to pass. Scaffolds can be sparse, consisting of only a few dispersed ants, or extremely dense, with many ants overlapping one another in a continuous cluster (Fig. 1 and SI Appendix, Movie S2). To ascertain how scaffolds form and under what conditions, and to assess their effects on foraging traffic, we conducted field experiments with wild colonies of E. burchellii, using an apparatus that allowed for manipulation of the slope over which ants traversed (Fig. 1). We observed and quantified the growth of scaffold structures and measured traffic variables to assess their influence on scaffold formation.

Our experimental results show that scaffolds repeatedly and predictably form on inclined surfaces, and we describe the relative influences of environmental and traffic variables on their growth. Informed by these observations, we propose a theoretical model that describes scaffold growth as a control response at the group level that emerges through a simple process of error correction at the individual level. Our model describes a hypothetical mechanism underlying scaffold formation that links individual-level sensing of disruption to the adaptive collective response of scaffold formation. Comparing the model predictions with our experimental observations, we show that the model captures well the observed dynamics of scaffold growth, and we derive an analytical solution that allows for straightforward prediction of these dynamics.

**Results**

Field experiments were conducted on Barro Colorado Island, Panama, in January and February 2015. The experimental apparatus consisted of two platforms three-dimensionally (3D) printed from PLA filament, each mounted on a flexible tripod 10 to 15 cm above the ground, linked together with an adjustable hinged panel (Fig. 1). The slope of this panel could be fixed at 10° increments, from nearly horizontal (20°) to vertical (90°), using a series of removable spacers, and it was covered with sandpaper.

**Fig. 1.** Overview of experimental apparatus and video analysis procedure. (A) Photo of apparatus setup in the field during an experiment, with platform set to 90°. (B) Diagram of 3D-printed apparatus assembly, with removable spacer to adjust platform angle shown in red. (C) Frames extracted from video of experiment with platform set to 90°, at experiment start (Left), after 120 s (Middle), and after 300 s (Right). (D) Diagram of image subtraction algorithm for video analysis. Moving ants are shown in gray, with stationary ants shown in black (indicating scaffold area over time) and the final scaffold area shown in red. (E) Dimensions of adjustable platform overlaid on photo of experiment in progress, with platform set to 80° (removable spacer to adjust angle visible at bottom left of platform).
to provide a naturalistic surface that ants could grip (as tested in pilot experiments). Before each experiment, foraging traffic was interrupted as the apparatus was cleared of ants and set to one of the treatment angles. When ants reestablished traffic across the platform, video recording commenced (marking the start of an experiment) and continued for 10 min (until experiment end). The angle of the sloped platform was the only factor directly manipulated since the rates of traffic and prey delivery fluctuate throughout the course of a raid. In total, 89 experiments were conducted, with either 10, 11, or 12 trials completed at each of the eight angle treatments, from 20° to 90°. After processing the videos, 4 trials had to be removed due to issues with recording quality, leaving a total of 85 trials for the main analyses, which involved estimating the rates of traffic and prey delivery and final scaffold size. For higher-resolution analyses involving estimates of scaffold growth over time (Fig. 2A and Model Fitting, Parameter Estimation, and Validation), an additional 15 trials had to be removed due to variations in lighting (see Materials and Methods for details).

Dynamics of Scaffold Growth. Scaffolds formed more often and grew larger on steeper slopes (Fig. 2B). We defined a stable scaffold as consisting of five or more ants stopped on the platform after 10 min. Scaffolds rarely formed on slopes of less than 40° (4.76% of experiments, N = 1 of 21), and they were far more likely to form at or above this threshold (81.25% of experiments, N = 52 of 64; see SI Appendix, Table S1, for summary data from each angle).

To examine the effects of the experimental variables on scaffold size, we fitted a linear mixed-effects model, using a negative-binomial distribution with linear parameterization, with the Generalized Linear Mixed Models using Template Model Builder (glmmTMB) package for R [version 1.0.1 (37, 38)]. The mean traffic flow rate (ants per second), prey delivery rate (prey-carrying ants per second), and log-transformed surface angle θ (degrees) were examined as fixed effects, with experiment date treated as a random effect to control for between-colony variations (the proportion of bidirectional traffic was considered as another possible fixed effect but correlated strongly with overall traffic so was not included). Models containing all possible combinations of the fixed effects and their interactions were ranked using the Bayesian information criterion (BIC), and the model with the lowest BIC was selected for this analysis. The final model contained the fixed effects of surface angle θ and prey delivery rate, along with the random effects of experiment date. Residual diagnostics (SI Appendix, Fig. S1) were obtained using the simulation-based approach implemented in the Diagnostics for Hierarchical Regression Models (DHARMs) package for R [version 0.3.0 (39)]. As shown in Fig. 2C, our mixed-effects model showed that the angle of the surface, θ (log-transformed), was the strongest predictor of final scaffold size (standardized effect size = 1.36; p < 0.001), followed by prey delivery rate (standardized effect size = 1.39; p < 0.001). Model predictions of final scaffold size at different angles and rates of prey delivery are shown in SI Appendix, Fig. S3.

Effects of Scaffolds on Foraging Traffic. To assess how scaffold formation affected the flow of traffic, we counted the number of ants slipping and/or falling when crossing the experimental platform for 2 min at the beginning and end of each experiment. Fig. 3 shows the proportion of ants that slipped and/or fell entirely from the apparatus during these time windows, at each angle. To examine the effect of the environmental geometry on the proportion of ants slipping and/or falling, we fitted a linear mixed-effects model using a binomial distribution [glmTMB package for R, version 1.0.1 (37)]. Surface angle θ (degrees) and time (either beginning or end of experiment) were used as fixed effects, with the date of experiment again treated as a random effect to control for between-colony variations. Models containing all possible combinations of the fixed effects and their interactions were ranked using the BIC, and the model with the lowest BIC was selected for this analysis. The final model contained the two aforementioned fixed effects, plus their interaction. Model diagnostics were performed by visually inspecting the random effect quantiles (shown in SI Appendix, Fig. S4).

This model showed that the surface angle θ was a significant predictor of the proportion of ants slipping and/or falling (standardized effect size = 0.71; p < 0.001; see SI Appendix, Fig. S5).

![Fig. 2. Scaffold growth data and results from mixed-effects model. (A) Growth curves showing scaffold size over time, grouped by angle treatment. Points indicate scaffold size at each second, measured by the software in pixel area, then converted to estimates of the number of ants in a structure, by scaling based on manual ground-truth counts. Color scale indicates rate of prey transport for each experiment. Total N = 70, with N for each angle from 20° to 90° = 5, 7, 9, 10, 9, 10, 11, and 9. (B) Box plot showing summary data of final scaffold size, grouped by angle. Pink dots indicate final values for each experiment. Total N = 85, with N for each angle from 20° to 90° = 10, 11, 10, 11, 12, 10, 12, and 9. (C) Results from the linear mixed-effects model, showing the standardized relative effects of each of the predictors on final scaffold size (expressed as incidence rate ratios). (Left) Relative strengths of fixed effects and (Right) relative strengths of random effects (date of experiment).](https://doi.org/10.1073/pnas.2013741118)
It also revealed a significant interaction between θ and experimental time (standardized effect size = -0.39; p < 0.001), indicating that the proportion of ants slipping and/or falling increased with θ at the beginning of an experiment but did not change along with θ when measured at experiment end (SI Appendix, Fig. S6). Thus, our model revealed that the proportion of ants slipping and/or falling was no longer affected by the slope of the surface after scaffolds formed, explaining the differences observed in Fig. 3.

Theoretical Model of Scaffold Growth. Informed by these experimental results, we developed a theoretical model to explore the potential mechanism underlying scaffold growth that links individual sensing of environmental properties with the group-level adaptive properties of the surface ants traverse. This is denoted by the coefficient β interest. To test this, we first treated the slipperiness of the surface ants traverse. This is denoted by the other state variable S, a measure of how likely a particular geometric is to cause slipping (and thus increased traffic disruption). We distinguish the surface property S from the ant property P, although they map to one another, such that crossing a surface with slipperiness S correlates with a probability of slipping P for individual ants.

From P and S, we can find the overall probability of an ant slipping and/or falling was no longer affected by the slope of the surface after scaffolds formed, explaining the differences observed in Fig. 3.

Fig. 3. Effects of scaffold formation on traffic disruption. Comparison between the proportion of ants slipping and/or falling while crossing the experimental platform during the first and last 2 min of each experiment (representing the absence or presence of scaffold structures, respectively).

The overall slipperiness of the surface over time can be triggered by temporarily losing their footing (or slipping), and this can be formalized in the model as two state variables describing the relationship between scaffold growth and the overall slipperiness of the surface over time. These are denoted by the model to be fitted to the data from each trial to generate fitted estimates for the two unknown parameters P and S, measured from the rates of traffic and prey delivery, and surface angle θ. The coefficient β is a scaling factor, which can be fixed as a constant, to avoid infrequent compared to ants joining the scaffold, at least on the time scale of our experiment, the results were comparable and, in many cases, inferior to the model described here (see SI Appendix, Materials and Methods, for details).

Informed by our results that show slipping to be reduced over time by scaffold formation (Fig. 3), we propose a second differential equation to describe this reduction in slipperiness as a function of scaffold growth, where S is reduced (with respect to its current value) in proportion to scaffold size A at time t, at a rate described by

$$\frac{dS}{dt} = -\beta pSA,$$

where p is the proportion of traffic made up of prey-carrying ants and θ is the angle of the slope, in radians. The coefficient β is a scaling factor, which can be fixed as a constant, to avoid infrequent compared to ants joining the scaffold, at least on the time scale of our experiment, the results were comparable and, in many cases, inferior to the model described here (see SI Appendix, Materials and Methods, for details).

Model Fitting, Parameter Estimation, and Validation. The ODE system above was solved numerically for A and S over a time span of 600 s, corresponding to the duration of an experiment, using the ode45 solver in MATLAB (R2020b). With the observations of scaffold size over time for each trial, optimization was performed using a nonlinear least-squares approach to generate fitted estimates for P and S, with the lscurvefit function and MultiStart solver (to avoid local minima). Corresponding experimental variables for each trial were input from our observations, including p and θ, measured from the rates of traffic and prey delivery, and surface angle θ. We interpret the coefficient β as a scaling factor that can be fixed as a constant, to focus on the effects of the two fitted parameters of biological interest. To test this, we first treated β as another free parameter to be fitted. As shown in SI Appendix, Fig. S7, estimates of denoted P. Since this term captures the effects of the environment on ant traffic, it can be used as a proxy estimate for the slipperiness of the surface ants traverse. This is denoted by the other state variable S, a measure of how likely a particular geometric is to cause slipping (and thus increased traffic disruption). We distinguish the surface property S from the ant property P, although they map to one another, such that crossing a surface with slipperiness S correlates with a probability of slipping P for individual ants.
its value varied widely but only at the lowest angles, representing cases where only small scaffolds formed. Thus, we considered $\beta$ to have been overfitted in these cases to compensate for measurement noise with small or unstable scaffolds. To avoid such overfitting, we determined that the model could still perform well if $\beta$ was fixed across all experiments instead, and we fixed it at a constant value derived from the median of these estimates (for the results described here, $\beta = 0.0023$). This also allows for clear interpretation of the model results since both of the fitted parameters are biologically meaningful terms.

Since the model describes scaffold growth over time, the fitting procedure is only relevant for cases where scaffolds reached a stable size, and their growth over time could be estimated. Therefore, we fit the model to data from our experiments at and above 50°, where lighting and video quality were consistent enough for reliable growth estimates to be obtained ($N = 44$). For an initial approximation of the overall goodness of fit, we estimated $R^2$ values for each of these fits, resulting in a mean overall $R^2 = 0.48$ (SD = 0.75). Filtering out cases with extremely low estimates ($R^2 < 0.10$) revealed better fits among the remaining subset of 37 experiments (mean $R^2 = 0.70$; SD = 0.17). To complement these approximations, we examined the model fits in more detail by visual estimation. We plotted predicted growth curves along with their 95% prediction intervals over the experimental observations, using a bootstrapping procedure to generate distributions of the estimated parameters $P_{t|\theta}$ and $S_0$ to identify confidence intervals for the estimates. In addition, we generated another set of prediction intervals to capture the variation in traffic over the course of each experiment. Fig. 4A shows representative example fits at four different angles, with the results from all other experiments where the model was fitted ($N = 44$) shown in SI Appendix, Figs. S8–S12.

To validate the model, we compared its predictions of the slipperiness of the surface $S$ with estimates from our observations of the quantity $P_S$, over the first and last 2 min of each experiment. As described above, $P_S$ represents the total proportion of ants slipping over a given time, including those that join a scaffold (measured by the scaffold size at $t$) and those that do not (for which we used our counts of ants slipping, as described in Effects of Scaffolds on Foraging Traffic). If the surface property $S$ can be expressed as a proxy measure of slipperiness in the same terms as $P_S$, and if the model captures how this property relates to the experimental variables, the predicted values of $S$ output by the model should be close to our observed estimates of $P_S$, even though none of the slipping observations were used in the model fitting.

To summarize the model predictions over the first 120 s, we took the mean values of estimates for $S$ at $t = 0, 60$, and 120, for each model run. As shown in Fig. 4B, the model predictions were similar to our experimental observations for $P_S$ over the same time window, grouped by surface angle $\theta$, predicting initially higher levels of slipperiness at steeper angles. Using the same procedure, we also estimated the mean values of $S$ over the final 120 s of each experiment, using the estimates of $S$ at $t = 480, 540$, and 600 s, to compare with the experimental measurements of the total slip probability $P_S$ over this same time. As shown in Fig. 4B, our model also predicted well the reduction in surface slipperiness that we observed in our analysis of ants slipping while crossing the experimental surface and how this varied depending on the angle $\theta$. Thus, the model predicted well the initial slip potential of the surface $S_0$ for each angle $\theta$ and how $S$ was reduced over time by scaffold growth.

Analytical Solution. In addition to the numerical procedure above, the system of differential Eqs. 1 and 2 that constitute the model can be solved analytically, with the solution given by

\[
\begin{align*}
S(t) &= \frac{S_0}{1 + \frac{P_{t|\theta}}{P_{0|\theta}}}, \\
S_0 &= \frac{P_{0|\theta}}{P_{t|\theta}},
\end{align*}
\]

Fig. 4. Model predictions and analytical solution. (A) Examples of growth curves predicted by the model (red lines), with the corresponding experimental observations (blue circles), for one representative trial at each of four angles ($\theta = 60, 70, 80, 90$, as noted). Results from all experiments for which the model was fitted ($N = 44$) are shown in Figs. S8–S12, grouped by surface angle $\theta$. Dark pink bands indicate 95% prediction intervals with the fitted estimates for $S_0$ and $P_{t|\theta}$, assuming constant traffic flow at the mean rate for each replicate. Light pink bands show 95% prediction intervals over the entire range of traffic, with lower bounds indicating predictions using the lowest rate as an input, and upper bounds using the highest. (B) Model predictions of $S$ compared with experimental observations of $P_S$ (mean $S$ (0–120 s) and $P_{480–600}$ s) and $P_{0–120}$ s, showing slippingness, $S$, and scaffold size, $A$, over time.
The parameters are as described above, with the initial conditions \( A(0) = 0 \) and \( S(0) = S_0 \). This solution can be used to generate straightforward predictions of some aspects of the system behavior and model output given certain experimental observations or estimates. For instance, the overall size of a scaffold tends to a final value \( A_F \), which is given by

\[
A_F = \sqrt{\frac{2 \mu S_0 P_{\||S}}{\rho \beta}} \tanh \left( t \sqrt{\frac{\mu p \beta S_0 P_{\||S}}{2}} \right),
\]

and

\[
S = S_0 - S_0 \tanh \left( t \sqrt{\frac{\mu p \beta S_0 P_{\||S}}{2}} \right)^2.
\]

The model predicts that scaffolds grow most rapidly at the start, before slowing down and settling at \( A_F \), and the probability of ants slipping (and potentially joining) decreases as they grow, tending to zero as scaffolds approach their maximum size. This is consistent with the following phase portrait analysis of the system, which demonstrates that such behavior must necessarily arise in any solution of the model. First, we combine both parts of [1] and [2], which gives

\[
d\bar{S} = \frac{\beta \theta A}{\mu P_{\||S}},
\]

to produce a phase portrait describing the relationship between \( A(t) \) and \( S(t) \) as the system evolves (Fig. 4C). Each trajectory of the system begins at some \((A, S) = (0, S_0)\), and as the system evolves, scaffold size increases, and the probability of slipping, and thus joining, decreases. The trajectory tends to some value \((A, S) = (A_F, 0)\), where the probability of slipping is 0, and the size is therefore fixed at some \( A_F \). This analysis makes clear that adjusting the value of \( S_0 \), maintaining all other model parameters, has the effect of reducing the final scaffold size.

**Discussion**

Our experimental results reveal that *E. burchelli* army ants consistently form scaffold structures when crossing surfaces inclined beyond a threshold of around 40°, above which ants begin to lose their footing. Scaffolds are responsive to local environmental geometry and specific traffic conditions: steeper slopes and higher rates of prey delivery and traffic are all associated with the formation of larger structures. When they form, scaffolds exhibit a saturating growth response, growing rapidly at first, then settling to a final size, which is a predictable function of the environmental variables. We also identify some of the functional benefits that scaffolds provide for the colony, including reducing losses of foragers and prey and alleviating traffic disruption.

Informed by these results, our theoretical model offers a simple and plausible mechanistic explanation of scaffold formation. It describes scaffold growth as a response at the collective level, akin to a form of distributed control, which results from a process of individual error correction. As observed in our experiments, ants crossing steeper inclines are more likely to slip, and we hypothesize that ants stop to join scaffolds with a conditional probability that depends on slipping, responding to this stimulus by attaching to the surface and remaining stationary. Our experimental observations quantify how scaffold formation reduces the probability of slipping over time, and the model predicts the initial slipperness of a surface at a given slope and how this is reduced over time as scaffolds grow. These predictions emerge from the model even though none of our experimental observations of ants slipping are used as inputs for the model or for the model-fitting procedure. The model captures the growth dynamics of scaffolds across a range of traffic conditions, providing insight into how particular combinations of traffic and environmental geometry influence their growth.

The growth rate and ultimate size of scaffolds are emergent properties that can arise from individual sensing and error correction, without the need for complex communication, enabling a system-level response that is suited to each particular configuration of traffic and environmental geometry. The error in this system is detected at the individual level, and individual error correction leads to the emergent group-level response of scaffold growth. This individually sensed error may be ants responding to their own slipping, experiencing disruption due to other ants slipping at an increased rate, or some combination of such stimuli. In any case, this error is reduced by the formation of scaffolds and is detected at the individual level; therefore, we consider this a distributed form of proportional control.

In an early description of army ant self-assembly from 1874 (40), the naturalist Thomas Belt asked, “Can it not be contended that such insects are able to determine by reasoning powers which is the best way of doing a thing, and that their actions are guided by thought and reflection?” Indeed, many dramatic examples of collective animal behavior were long thought to result from sophisticated reasoning capabilities or even unknown mechanisms involving thought transfer. However, research on collective behavior has revealed common principles that explain complex group-level behaviors as the result of relatively simple individual rules and local interactions (41, 42). A key challenge in understanding these principles is ascertaining how inputs from individual-level sensors are combined and filtered to produce group-level outputs (43), and such insights have come to inform the design of complex engineered systems that rely on distributed forms of control (44). Thus, while the experiments described here involved ants crossing an inclined surface, our results may be relevant to other systems in which group-level properties can be modified via individual error sensing and correction.

As human technological and social systems increase in complexity, the need for simple, robust mechanisms for error correction to rapidly respond to systemic disruption at multiple scales without relying on complex communication or oversight is paramount. Such simple mechanisms can avoid some of the pitfalls that often arise from communication within groups like biases (45, 46) that result from the oversharing of correlated information (47, 48), and the model presented here may inform approaches to addressing these. Our model and experimental results should provide insights for the field of swarm robotics as well (49–51), in which increasingly complex group behaviors, including self-assembly (52), are often constrained by technical limitations on both individual sensors and communication capabilities (53).

In this context, our model relates to previous models of self-organized aggregation that have informed the development of robotic systems, such as models of cockroach aggregation in which individuals assess others already in a cluster (50, 54–56) or corpse piling in ants, where individuals respond to corpses already deposited, and to other ants (57, 58). However, the mechanism we have identified requires less complex sensing since the ants need only to respond to their own slipping and do not need to assess the size of a structure in place. This describes scaffold growth as a rapid, on-demand response that alleviates disruption that would otherwise result from the dynamic environmental conditions. With minimal requirements for sensing and information processing, this finding may also be relevant at smaller scales, for the design of self-healing materials (59–61) and future developments in biofabrication (62, 63).
Materials and Methods

**Experimental Protocol, Site, and Study Organism.** Field experiments were conducted in the seasonal tropical forest of Barro Colorado Island, Panama, in daylight hours during the dry season in January and February 2015. Three diamictropic colonies of *E. burchelli* foreli (Mayr), all in the nomadic phase, were located and used in experiments for 2 or 3 d each. Experiments were performed along the primary raiding trail, where traffic is most consistent throughout the day, within 50 m of the bivouac (temporary nest site). Each morning a suitable location along this trail was found, with enough space to set up the experimental apparatus and recording equipment. A break was created in the existing trail by removing sticks and leaves from the substructure, and the apparatus was inserted into the resulting gap. A field compass with spirit level was used to adjust the side platforms to be level and in plane with one another when setting up the apparatus. The flow of ant traffic was redirected up onto both ends of the apparatus using the displaced sticks and leaves as makeshift ramps, which were covered in trail pheromone from preexisting traffic. An experiment was considered to begin once the first ant had reestablished a continuous flow of traffic over the apparatus, and each experiment was recorded for 10 min from this point.

**Data Collection and Extraction.** Videos were filmed with a high-definition camcorder (Panasonic HDC-300) at 30 fps. The camera was mounted on a tripod and positioned perpendicular to the surface being filmed, at a distance of approximately 60 cm. After initial processing of the video footage, 4 experiments were discarded due to issues with the experimental setup or recording, leaving a total of 85 experiments used for the analyses in *Dynamics of Scaffold Growth and Effects of Scaffolds on Foraging Traffic*. These involved measuring the traffic variables as described below and final size of scaffolds. Out of these, 70 experiments were used for further analyses of scaffold growth over time using the automated method described below, with 15 videos that could not be analyzed at higher temporal resolution due to wide variations in lighting (although we could still estimate the final scaffold size and traffic counts, so these could be used for the first part of the analysis).

The rate of ant traffic for each experiment was measured by visual counts taken of each video, slowed to 0.25× speed using VLC media player. The total number of ants crossing the edge of the experimental surface in each direction was counted during three 30-s windows over the course of each 10-min experiment—from 00:30 to 01:00, from 05:00 to 05:30, and from 09:00 to 09:30. Counts from both directions were combined, and these totals were divided by 30 s to give an estimate of the traffic rate at experiment start, middle, and end, in ants per second. The mean of these three rates was taken as an estimate of the overall traffic rate for each experiment. The proportion of bidirectional traffic was calculated by dividing the traffic flow in the nondominant direction by the overall traffic flow. The rate of prey delivery was estimated in a similar way, by manually counting the number of prey items transported for the duration of each experiment (at full speed). The total number of prey-carrying ants was divided by 60 s, giving a value for the prey delivery rate in prey-carrying ants per second.

Visual counts were also made of the number of ants that slipped and/or fell from the platform when crossing. These counts were performed for the first 2 min and last 2 min of each experiment, to estimate the effects of the presence or absence of scaffold structures (in those cases where one formed). A slip was defined as a sudden vertical displacement of more than one body width, in which an ant crossed a line estimated as the lower boundary of the trail across the platform (or the structure, if one formed). A fall was counted if an ant fell completely from the platform.

The size of scaffolding structures over time was measured using a custom image subtraction algorithm in MATLAB R2016b (version 9.0). Subsequent frames of the videos were compared to detect and record whenever an ant remained stationary (joined a structure) rather than moving with the flow of traffic (Fig. 1D). Output from the software was given in pixel area, then these estimates were rescaled by comparison with manual ground-truth counts to estimate the number of ants within a scaffold (see SI Appendix, Materials and Methods, for details of the scaling procedure).

**Probability Estimates for Theoretical Model.** To derive the probability estimates underlying the theoretical model, we first make the simplifying assumption that all ants joining a scaffold have slipped. This does not require an explicit definition of what constitutes a slip, so we can state more generally that all scaffold ants have experienced some slipping-related stimulus. Since scaffolds do not form on level surfaces and only rarely form on shallow slopes, as our data show, we consider this a reasonable assumption. This assumption can be expressed as the conditional probability *P* (slip|join), or the probability that an ant has slipped (or experienced some slipping-related effect, e.g., from traffic disruption or collisions) given it has joined a scaffold. Assuming this is the case for all joining ants, the conditional probability *P* (slip|join) will be equal to 1. From this value, we can estimate the inverse conditional probability *P* (join|slip), or the probability that an ant will join a scaffold, given it has slipped. Based on our observations, we hypothesize that this individual-level response may be crucial to the mechanism of scaffold growth, so we use Bayes' theorem *P* (A|B) = *P* (B|A) *P* (A) / *P* (B) to estimate it as follows:

\[ P(\text{join}|\text{slip}) = \frac{P(\text{slip}|\text{join}) P(\text{join})}{P(\text{slip})} \]  

Since the other conditional probability, *P* (slip|join), is assumed to be equal to 1, we can simplify and rearrange the above equation in terms of a quantity we can relate directly to our data, the term *P* (join), which describes the overall probability of an ant to join a scaffold:

\[ P(\text{join}) = P(\text{slip}|\text{join}) P(\text{slip}) \]  

From this relationship, we derive Eq. 1 of our model of scaffold growth directly, in its simplest form, by incorporating a traffic flow rate of ants, a proportion of which are induced to join a scaffold when crossing the surface, given by *P* (join). This results in a rate of ants joining over time, corresponding to the change in scaffold size, dA, in our model. We rename the conditional probability *P* (join|slip) to *P* s for simplicity and reformulate the term *P* (slip) as a measure of the effect of environmental geometry on ant traffic (S, or slipperiness), rather than an internal ant property. However, the two can still be compared in the same terms (as a proportion of ants slipping), and we retain the term *P* (slip) (shortened to *P* s) when referring to a property of ants.

*P* s describes the overall probability of an ant slipping at a given time. Therefore, this accounts for ants that join scaffolds (assumed to have slipped) as well as those that slip and continue walking, both of which can be estimated from our experimental observations. To compare the observed estimates of *P* s from our data, obtained by this method, correspond quite well with model estimates of the mean values of *S* over the same time windows (output from numerically solving the model's differential equations) to offer strong support for the model's validity.

**Data Availability.** All study data are included in the article and SI Appendix. All code used in the study for data analysis are available at GitHub, https://github.com/matthewlutz/army_ant_scaffolds.

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