A social mechanism facilitates ant colony emigrations over different distances

Thomas A. O’Shea-Wheller*, Deraj K. Wilson-Aggarwal, Duncan E. Edgley, Ana B. Sendova-Franks and Nigel R. Franks

1 School of Biological Sciences, University of Bristol, Life Sciences Building, 24 Tyndall Avenue, Bristol, England.
2 Department of Engineering Design and Mathematics, UWE Bristol, Frenchay Campus, Coldharbour Lane, Bristol, England.

*Corresponding author E-mail: to13870@bristol.ac.uk.

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Summary statement

Within social systems, we often see ‘key’ individuals that are disproportionately active in various tasks. Here however, we show that the contribution of such individuals may be far from decisive.

Abstract

Behavioural responses enable animals to react rapidly to fluctuating environments. In eusocial organisms, such changes are often enacted at the group level, but may be organised in a decentralised fashion by the actions of individuals. However, the contributions of different group members are rarely homogenous, and there is evidence to suggest that certain ‘keystone’ individuals are important in shaping collective responses. Accordingly, investigations of the dynamics and structuring of behavioural changes at both the group and individual level, are crucial for evaluating the relative influence of different individuals. Here, we examine the composition of tandem running behaviour during colony emigrations in the ant species *Temnothorax albipennis*. Tandem running is modulated in response to emigration distance, with more runs being conducted when a more distant nest site must be reached. We show that certain individuals are highly active in the tandem running process, attempting significantly more work in the task. Contrary to expectations, however, such individuals are in fact no more successful at conducting tandem runs than their less active nest mates. Instead, it seems that when more tandem runs are required, colonies rely on greater recruitment of workers into the process. The implications of our study are that in some cases, even when apparently ‘key’ individuals exist within a group, their relative contribution to task performance may be far from decisive.

Introduction

Spanning a broad range of species and genera, behavioural modification constitutes an essential component of the adaptive repertoires of animals. Examples range from simple Pavlovian responses in mammals (Jenkins et al. 1978), to complex, multi-generational cultural changes in human societies (Kirk 1996; Inglehart 1997) and inter-species mutualisms in the social insects (Way 1963). Common to all of these strategies, is the advantage of rapid benefits when compared to the more gradual changes elicited by natural selection alone. Considerable effort has been invested into studying behavioural changes at both the group (Barsade 2002; Langridge et al. 2004) and the individual level (Shettleworth 2001). However, the interplay between the two, especially in complex networks, is less well understood (Bornstein & Yaniv 1998; Franklin & Franks 2012; Anderson & McShea 2001; Hunt et al. 2016).

Social insects display complex life history characteristics, underpinned by efficient colony organisation, and thus provide excellent models for studying how group level behavioural changes are structured through the actions of individuals. This is in no small part due to their propensity for self-organisation, whereby complex processes may emerge without the need for centralised control, an ability that many human systems lack (Camazine et al. 2003). Self-organisation can lead to the formation of adaptive, but flexible group behaviours, which often confer additional benefits to colony functioning (Goss et al. 1989). Hence, such behaviours are of interest when attempting to elucidate how multi-level behavioural change is structured. Colony relocation represents one process that is regulated at both the group and individual level in social insects, and indeed, wide
variation exists among species in relation to how migration effort is assigned. Examples range from varying
distributions of worker effort within colonies, as seen in the ant Myrmecina nipponica (Cronin 2015), to extreme
cases where a single individual is key to overall collective organisation, such as in the queenless Ponerine,
Diacamma indicum (Sumana and Sona 2013).

In the ant Temnothorax albipennis, emigration behaviour is regulated at the group level to best suit
environmental conditions, but relies on individual contributions and information sharing (Franks & Richardson
2006). Colonies of this species reside within rock cavities in a moderately heterogeneous and unstable
environment, and as such, they frequently have to emigrate to potential new nest sites in order to maintain their
protection from external threats (Dornhaus et al. 2004). Before an emigration can occur, however, scouting
workers must accumulate within a new nest site until a ‘quorum threshold’ is reached (Pratt 2004). A key
behaviour in this process is tandem running, in which workers lead their nest mates to the location of a potential
nest site or food source (Franklin & Franks 2012). During a tandem run, the leading worker will maintain
intermittent tactile contact between its gaster and a follower’s antennae, leading it to the desired resource via an
often-tortuous journey, and thus informing it of the route (Franks & Richardson 2006). Previous work has
shown that colonies of T. albipennis increase their rate of tandem running; defined as the total number of
tandem runs in relation to colony size, during emigrations where the target nest site is further away (O’Shea-
Wheller et al. 2016). This may be explained by the need to reduce emigration time and thus colony exposure
during more distant emigrations (Traniello 1989), and it is elicited by a reduction in the rate of nest discovery
(Pratt 2004). Despite being colony wide phenomena, tandem running responses are necessarily achieved by
perception at the individual level, whereby workers determine that a quorum has not yet been met in a new nest
site (Pratt 2004). As a consequence, the individual propensities of ants to engage in tandem running are likely to
influence the overall structuring of the process, yet the dynamics of this remain unexplored.

Several potential hypotheses may explain how individuals contribute to an increase in tandem running over
longer emigration distances. First, workers that would not usually be involved begin to assist in leading or
following tandem runs. Second, a ‘core’ of the same highly active workers simply lead more tandem runs
themselves, or third, a combination of both may occur. The relative likelihood and significance of such options
is uncertain, as previous studies have shown complex and differing structures of work distribution within
colonies. Much of the work conducted to date supports the concept that T. albipennis colonies have some
‘specialist’ workers in relation to certain task groups (Dornhaus 2008; Dornhaus et al. 2008). It has been noted
that task propensities are positively correlated within groups of related tasks e.g. ‘foraging’ and ‘emigration’.
Moreover, workers highly active in one task group may be less so in others (Pinter-Wollman et al. 2012),
suggesting ‘situational specialisation’. Nevertheless, some workers may still be persistently active across all task
groups (termed elites), though they constitute only a small proportion of individuals (Pinter-Wollman et al.
2012). In addition, the extent of specialisation may vary depending on colony characteristics, for example, it has
been shown that during brood carrying, smaller colonies will employ some specialist workers that are more
active than others, while in larger colonies, each active individual contributes more evenly to the process
(Dornhaus et al. 2008). Consequently, a greater understanding of how increased tandem running and other
behavioural changes occur during more distant emigrations, should help to shed light upon the underlying
processes that govern group and individual behaviour in a decentralised system.
In this study, we examined several key stages in both short and long distance emigrations at the group and individual level, in an effort to determine how behavioural changes were structured across individuals. Chiefly, we wished to determine if increases in the rate of tandem running were due to key individuals up-regulating their activity, more workers becoming involved, or indeed a mix of these two. We also tested the hypothesis that workers would alter their average tandem running return speed across emigration distances, as faster return rates would allow ants to recruit other colony members to a new nest more rapidly. Additionally, we attempted to determine if there was a link between initial scouting behaviour and a worker’s subsequent propensity to become actively engaged in emigrations.

We addressed these questions by quantifying colony level changes, including those associated with tandem running, scouting, average return speed, and workforce mobilisation. Then, in order to dissect the processes underlying such changes, we measured distributions of individual scouting, tandem running, and movement speed, having uniquely marked all workers within each of the 10 experimental colonies. Furthermore, we identified behavioural sub-groups of workers by categorising individuals in relation to their engagement in only one, or both emigration distances. We then assessed differences in attempted and successful per capita tandem running workloads between these groups, to quantify their relative importance to the system as a whole.

**Materials and methods**

**a) Colonies**

We collected ten colonies of *Temnothorax albipennis* (Curtis, 1854) from the isle of Portland, Dorset on 29th of September 2015. Colonies contained between 47 and 187 workers, between 19 and 130 brood items and each had a single queen. No specific permission was required for collection, with ants being taken from a disturbed quarry area open to the public. Additionally, no collection permit was required, as *T. albipennis* is not a protected species. In order to minimise the impact of colony removal on the local population and ecology, we used an established rota to vary collection location, based on date.

Colonies were migrated into initial laboratory nests and allowed to acclimatise for a period of 7 days after collection. We maintained them in the laboratory under established conditions (Dornhaus et al. 2004; Franks et al. 2003), with all nests being housed in plastic Petri dishes featuring Fluon coated sides. Ants were fed weekly with *Drosophila melanogaster* (Meigen, 1830), and allowed to forage for water and honey solution *ad libitum*.

**b) Marking**

The ten colonies were randomly numbered in order to determine a marking sequence. For each colony in turn, we anaesthetised all workers with CO₂ and painted them with a unique permutation of three colours, one on the thorax and two on the gaster (fig. 1A). A mounted single-hair and microscope were used in order to apply the paint with precision. Markings corresponded to an identification key, allowing rapid determination of individual identity, in addition to total colony populations (S1, Tab1). Care was taken to ensure that CO₂ exposure was kept to a minimum, and there was at least a 48h interval between marking and experimental trials, as studies have shown that insect behaviour may be affected after anoxia (Poissonnier et al. 2015). Further precautions
were taken to ensure that paint did not cover the gaster tips of workers, as this may impair the use of chemical signalling (Stuttard et al. 2015).

c) Nest design

We utilised two different nest qualities in our experiment; ‘poor’ and ‘excellent’. Colonies were initially housed in ‘poor’ quality nests, and allowed to emigrate into the ‘excellent’ nests during trials. Both designs were composed of microscope-slide lids and bases with a cardboard perimeter sandwiched between the two to create an internal cavity. The area of this cavity was constant across both nest qualities, measuring 65x35mm. ‘Poor’ quality nests had 1 mm high walls, a 6 mm wide entrance and a clear cover, while ‘excellent’ quality nests had 2 mm high walls, a 1 mm wide entrance and a red filter cover. These differences combined to make excellent nests more spacious, more secure, and darker than ‘poor’ quality nests, and thus far superior in the ants’ assessment (Franks et al. 2003).

d) Emigration trials

All colonies underwent emigration trials under ‘move-to-improve’ conditions, in which their original nests were left intact, allowing them to emigrate of their own accord to the superior ‘excellent’ quality nests (Dornhaus et al. 2004), at distances of 100 mm and 300 mm (fig. 1B,C). Twenty emigrations were conducted in total, over a period of 35 days, with each colony performing two emigrations in turn. In order to account for the potential experience effects of multiple emigrations (Langridge, Sendova-Franks, & Franks, 2008), using random assignment, five of the colonies underwent the 100 mm trial first, and five the 300 mm trial first.

Prior to the initiation of each trial, we cleaned 1000x1000 mm arenas with 70% ethanol solution, and illuminated them evenly using the laboratory lighting system (4100 Lumens, Colour rendering index: 85). Colonies housed in their original ‘poor’ nests were then transferred into the arenas and empty ‘excellent’ quality nests were placed at either 100 mm or 300 mm from them. At the initiation of experiments, the emigration process was observed and we began monitoring scout identities and numbers, tandem leader and follower identities and tandem durations, tandem run success rate at the two different distances, average ant return speed, and the total numbers of ants in the arena and nests (at 10min intervals). Additionally, HD video recording equipment (Panasonic HC-X900 series) was set up over both nest sites for the duration of emigrations, in order to facilitate further data analysis (fig. 1B, C).

e) Range of assessment factors

Analyses were broadly divided into investigations of the distribution of tandem running among workers, average return speeds, and the effect of scouting propensity on emigration engagement. Within these groups, we examined both colony level behaviour, and that of individual ants (via marking).

To quantify behavioural changes in tandem running distribution, we compared the total number of tandem runs per colony; percentage colony engagement in tandem leading or following, and the total number of tandem runs per ant. By measuring the total number of tandem runs per individual, we were able to compare the activity of ants that were involved at a single emigration distance, to those involved across both emigration distances. This division method was important for testing the hypothesis that the same key individuals would conduct significantly more tandem runs at a longer distance (i.e. those individuals comprising the ‘both’ group).
To achieve this, workers were divided into two groups; those actively engaged in only one emigration distance (the ‘either’ group), and those actively engaged in both emigration distances (the ‘both’ group). Engagement in an emigration was defined as workers conducting scouting, tandem running, or both.

We then compared the attempted and successful per capita tandem running workload of ants using the same division method, in order to account for the different sizes of these groups. Per capita tandem-running workload constituted the average amount of tandem running work (leading or following) done by each tandem-running worker, as a proportion of the total tandem running work conducted by all workers in a given emigration, and thus was a measure of individual contributions. In concert with the total absolute numbers of tandem runs conducted by workers, the use of this proportional measure allowed us to determine the relative importance of worker groups in the tandem running process at different distances. Additionally, we tested the effect of numbers of attempted tandem runs on an individual’s tandem running success.

We quantified average worker return speed across the two emigration distances, pooled by colony. When returning to their old nest, workers take a direct route between sites (Franks & Richardson 2006), and thus speed was calculated as a function of distance over time. To assess the effect of prior scouting on an individual’s propensity to engage in emigrations, we measured scouting effort (defined as the numbers of ants exploring outside the nest over time) across colonies, and the effect of prior scouting experience (defined as leaving the nest to explore) on a worker’s subsequent involvement in tandem running. New-nest discovery time, percentage colony involvement in emigrations and quorum attainment time were also analysed.

f) Statistical analysis

For all colony-level data, we used Shapiro-Wilk tests to check for normality, and hence to inform the application of appropriate statistical tests. Where the data were found to be normally distributed, we conducted either paired or independent t-tests. For measures in which the data were not normally distributed, we used a combination of Wilcoxon signed-ranks tests and Independent Sample Mann-Whitney U tests. As non-parametric tests are less powerful than parametric ones, when the significance level was between 0.05 and 0.1, we double-checked results by applying parametric methods to transformed data. In all such cases, the significance of the tests was unaltered.

When individual data were used, we employed Generalised Linear Mixed Models to account for the effect of colony and worker ID. All GLMMs included at most, two fixed predictor variables. Model selection was based on AIC and always began with the full model, involving interaction. The fit of all GLMMs was validated via analyses of Pearson residuals. All statistical analyses were performed in SPSS (Release versions 21.0.0.0, 23.0.0.0, IBM Corporation and other(s) 1989, 2012, 2016).

Tandem running at colony level

We used a combination of paired-sample t-tests for measures with normally distributed data, and Wilcoxon signed-ranks tests for measures where the data were not normally distributed.
Individual tandem running distribution

GLMMs assessing the effect of the number of leading and following attempts on successes of leading and following, used leading or following success/failure as the binomial response variables with a logit link, distance (100 mm or 300 mm) as a fixed factor predictor, number of attempted leads or follows as a covariate, and colony ID as a random factor. Worker ID was redundant, as no repeated measures were taken for any of the individuals. The GLMM assessing the number of attempted tandem runs by workers across emigration involvement groups used the total number of attempted tandem runs as a normal response variable with a log link, distance (100 mm or 300 mm) and involvement in either or both emigrations as fixed factor predictors, the interaction between the two, and colony ID and worker ID nested within colony ID as random factors. The GLMM assessing attempted per capita tandem running workload used per capita attempted workload as a normal response variable with a log link, distance (100 mm or 300 mm) and involvement in either or both emigrations as fixed factor predictors, the interaction between the two, and colony ID and worker ID nested within colony ID as random factors. The GLMM assessing per capita successful tandem running workload used successful workload per capita as the normal response variable with a log link, distance (100 mm or 300 mm) and involvement in either or both emigrations as fixed factor predictors, the interaction between the two, and colony ID and worker ID nested within colony ID as random factors.

We used paired t-tests to compare the percentage of individuals involved at both emigration distances to those involved at only one distance for each colony.

Average worker return speeds

A Wilcoxon signed-ranks test was employed to determine differences in the average return speed of workers between the two distances.

Scouting propensity and emigration engagement

The difference in the number of active scouts between distances was tested using a Wilcoxon signed-ranks test. To calculate the tandem running propensity of workers based on previous scouting activity, we measured the numbers of ants engaging in tandem runs as a proportion of the total population in each scouting category (the two scouting categories were individuals previously involved in scouting, and individuals not previously involved in scouting), within colonies. This was assessed using a combination of Mann-Whitney U tests (leading) and independent sample t-tests (following). Differences between the two distances were quantified with a Mann-Whitney U test.

Discovery times and quorum attainment

Distance-based differences in the time of quorum attainment were tested with a paired-sample t-test, while differences in new nest discovery time were assessed with a Wilcoxon signed-ranks test.
Results

\textit{a) Tandem running at colony level}

Under the longer-distance emigration treatment, colonies on average employed a higher number of tandem runs (Wilcoxon signed-ranks tests, \(U=2.194, n=10, \text{median}_{100}=8.5, \text{median}_{300}=14, P=0.028\), fig. 2A). By contrast, the percentage of successful tandem runs was reduced, on average, at the longer emigration distance, although this was not statistically significant (Paired-sample t-test, \(t=1.143, \text{df}=262, P=0.254\), fig. 2B). Additionally, at the longer emigration distance, colonies utilised a greater percentage of their potential workforce in both tandem running (Paired-sample t-test, \(t=48.00, \text{df}=9, P=0.037\), and the emigration process \textit{in toto} (scouting and tandem running) (Paired-sample t-test, \(t=-3.214, \text{df}=9, P<0.001\)) (fig. 2C,D).

\textit{b) Individual tandem running distribution}

Across both emigration distances, an individual’s probability of leading a tandem run successfully, significantly increased with the number of leading attempts made (GLMM, odds ratio=1.422, \text{df}=140, \(P=0.017\)), independent of colony ID (Colony random effect: \(Z=0.918, P=0.359\)). Furthermore, in agreement with the colony-level data, tandem leaders had a higher probability of success when leading at the shorter distance (GLMM, odds ratio=2.100, \(P=0.030\)). However, the number of following attempts had no significant impact on the probability of following success (GLMM, odds ratio=0.216, \text{df}=189, \(P=0.735\)), independent of the random factor (Colony random effect: \(Z=0.775, P=0.439\)). The probability of following success was higher, but not statistically significantly so, at the shorter distance (GLMM, odds ratio=1.284, \(P=0.427\)).

Across colonies, the percentage of individuals involved (scouting or tandem running) in emigrations at both distances was smaller than the percentage involved at just one distance (Paired-sample t-test, \(t=-4.039, \text{df}=9, P=0.001\), fig. 3). The interaction between distance and involvement group was significant in its effect on the total number of tandem runs attempted per individual (GLMM, effect of interaction: \(F_{1,262}=4.455, P=0.036\), fig. 4), indicating that worker group was important within distances. Specifically, the ‘both’ engagement group attempted more tandem runs per individual than the ‘either’ group at 300 mm (GLMM, Bonferroni post-hoc test: \(t=5.098, \text{df}=262, P<0.001, \text{mean}_{\text{either}}=1.281, \text{mean}_{\text{both}}=2.000\), but not at 100 mm (GLMM, Bonferroni post-hoc test: \(t=1.143, \text{df}=262, P=0.254, \text{mean}_{\text{either}}=1.145, \text{mean}_{\text{both}}=1.316\)), suggesting a greater role for this group at the longer distance.

Following on from this, the per capita attempted tandem running workload of ants in the ‘both’ group was significantly higher than that in the ‘either’ group across both emigration distances (GLMM, effect of group: \(F_{1,292}=18.327, P<0.001; \text{mean}_{\text{either}}=0.579, \text{mean}_{\text{both}}=0.750\)). The attempted per capita tandem running workload of the ants was lower at 300 mm than at 100 mm, regardless of their involvement in either or both emigrations (GLMM, effect of distance: \(F_{1,292}=15.149, P<0.001, \text{mean}_{100}=0.741, \text{mean}_{300}=0.586\), fig. 5A). However, the interaction between distance and involvement group was very close to the threshold of significance (GLMM, effect of interaction: \(F_{1,292}=3.817, P=0.052\)). This may be explained by the per capita workload performed by the ‘both’ group being significantly higher than for the ‘either’ group at 300 mm (GLMM, Bonferroni post-hoc test:
Our results show P=0.087. Interestingly, when only successful tandem running workload was examined, these differences were somewhat diminished; per capita tandem running workload was still lower at 300 mm than at 100 mm across both involvement groups, (GLMM, effect of distance: F_{1,292}=4.831, P=0.029, mean_{100}=0.737, mean_{300}=0.591, fig. 5B). Ants in the ‘both’ group also still had a significantly higher successful per capita tandem running workload than those involved in the ‘either’ group across both distances (GLMM, effect of group: F_{1,292}=6.037, P=0.015, mean_{either}=0.582, mean_{both}=0.748, fig. 5B), however, the interaction between distance and whether ants were involved in either or both emigrations was not significant (GLMM, effect of interaction: F_{1,292}=0.012, P=0.915). This was due to a lack of difference within distances at both 100mm (GLMM, Bonferroni post-hoc test: t=1.774, df=292, P=0.077, mean_{either}=0.654, mean_{both}=0.831) and 300mm (GLMM, Bonferroni post-hoc test: t=1.677, df=292, P=0.097, mean_{either}=0.519, mean_{both}=0.673).

c) Average worker return speeds

The average return speed of workers across colonies was significantly higher for longer emigration distances (Wilcoxon signed-ranks test, U=-3.92, n=10, P<0.001, fig. 6).

d) Scouting propensity and emigration engagement

Before emigrating over the longer distance, on average, colonies employed significantly greater numbers of scouts (Wilcoxon signed-ranks test, U=2.253, n=10, P=0.024, mean_{100}=16, mean_{300}=25). In any given emigration, previously scouting significantly increased a worker’s propensity both to lead (Mann-Whitney, U test, U=3.250, n_{scouted}=10, n_{not-scouted}=10, P<0.001, median_{scouted}=0.176, median_{not-scouted}=0.040), and to follow (Independent-sample t-test, t=-3.680, df=9, P=0.002, mean_{scouted}=0.172, mean_{not-scouted}=0.084), tandem runs, and this trend was not significantly different between emigration distances (Mann-Whitney U test, leading: U=0.338, n_{100}=10, n_{300}=10, p=0.738, median_{100}=0.071, median_{300}=0.084; following: U=1.826, n_{100}=10, n_{300}=10, p=0.068, median_{100}=0.073, median_{300}=0.137), however, it should still be noted that the median for following was much higher at 300mm.

e) Discovery times and quorum attainment

The time of quorum attainment was significantly later in emigrations over longer distances (Paired-sample t-test, t=-3.565, df=9, P=0.006, mean_{100}=39.41 min, mean_{300}=94.22 min). However, though later in the distant treatment, nest discovery time was not significantly affected (Wilcoxon signed-ranks test, U=9.811, n=10, P=0.114, median_{100}=13.37 min, median_{300}=31.54 min, checked with a paired t-test after log10 transformation, P=0.087), likely in part due to the increased rate of scouting.

Discussion

Our results show that in *T. albipennis* colonies, tandem running activity is redistributed across individuals in response to differing emigration distances. In agreement with previous work (Pratt et al. 2002; Pratt 2004; O’Shea-Wheller et al. 2016), we found that the incidence of tandem running rose significantly with longer emigration distances (fig. 2A,C, 4). Tandem success tended to decrease with distance, although this result was
statistically significant only for leaders. It was not significant either for followers, or colonies as a whole (fig. 2B). Furthermore, a significantly larger proportion of each colony was engaged in emigrations and tandem running at 300mm, and average worker return speeds were higher (fig. 2C,D, fig. 6). Crucially, we observed that a small minority of individuals attempted a greater workload (leading or following tandem runs) at the longer emigration distance in comparison to their nest mates (fig. 5A). However, the relative importance of these ants dwindled when only successful workload was considered (fig. 5B). In concert, our data show that while colonies use multiple processes to adjust emigration dynamics over longer distances, an increase in the number of active individuals is likely the key driver behind increased tandem running (fig. 2C,D).

The upregulated investment in tandem running over longer emigration distances may be explained by the lower independent discovery rate of more distant nests (O'Shea-Wheller et al. 2016; Pratt, 2004). This is because to commit to a new nest colonies must achieve a quorum threshold, which may be attained by ants individually discovering a new nest, or being recruited to it via tandem running (Pratt, 2004). Consequently, a decrease in individual discovery rate seems to require an increase in tandem running to reach such a quorum (fig. 2A).

Interestingly, average worker return speed also increased, and the explanation for this likely arises from the properties of ant locomotion; it has been shown that for any given movement, the longer the duration, the higher the average speed (fig. 6) (Christensen et al. 2015). However, while every movement is bracketed by an initiation and termination phase, these phases comprise a constant percentage of the overall movement, suggesting that ants determine movement duration before initiating a journey (Christensen et al. 2015).

Consequently, this indicates that ants up-regulate their speed when planning to move over longer distances (Hunt et al. 2016). Previous studies have found that this speed-distance rule applies to short discrete movements, yet the majority of journeys are made up of many such movements. Here, to our knowledge for the first time, we show that it is also applicable to longer, compound movements, indicative of a consistent overarching mechanism.

Differences at the colony level were underpinned by variation in individual behaviour between emigration distances. The finding that individual tandem running workload (leading and following tandem runs) decreased with the number of ants actively engaged (fig. 5A,B, fig. 2D) is consistent with observations in analogous systems, and constitutes a commonly observed phenomenon in insect societies (Karsai & Wenzel 1998; Dornhaus et al. 2008). Principally, the decrease in individual contributions when more ants are engaged may be explained by the reduced need for workers to be involved in a task, and the concomitant reduction in task stimuli (Pinter-Wollman et al. 2012). Furthermore, for longer emigrations, although more tandem runs may be required due to a reduced encounter rate (O’Shea-Wheller et al. 2016, Pratt 2004), colonies of *T. albipennis* use quorum thresholds relative to their total population (Dornhaus & Franks 2006). As such, the quorum threshold required by a colony of any given size should remain constant, and thus the task of attaining it would potentially be distributed over more workers at longer emigration distances, creating a dilution of individual effort (Dornhaus & Franks 2006).

The changes in numbers of tandem runs between distance treatments might largely be attributed to the greater numbers of workers engaging in more distant emigrations. However, in concert with this, some ants that were involved in multiple emigrations conducted many more tandem runs at 300 mm than the rest of the actively engaged worker population (fig. 4). This suggested that there were certain individuals specialising in the task, as
is the case for other task groups in this species (Robinson et al. 2009; Pinter-Wollman et al. 2012). As such, there was the potential for these individuals to contribute significantly to the tandem running process, via task threshold differences and reinforcement processes (Sendova-Franks et al. 2002; Sendova-Franks & Franks 1995). The idea of task reinforcement was supported further by the finding that more leading attempts did correlate with a higher success rate, and so we investigated the relative contributions of worker groups actively engaged in emigrations, in an effort to isolate a specialist faction.

We defined actively engaged workers as those conducting scouting, leading or following of tandem runs, or both. This definition was adopted as previous scouting activity significantly increased an individual’s propensity to either lead or follow tandem runs, a trend that remained constant across both distances, although was more probable at 300mm. We separated actively engaged workers into two groups; workers actively engaged across both distances, and workers actively engaged at only one distance. This division method allowed us to determine how individuals recurrently involved in emigrations contributed to the process. This latter group formed a minority of the total population of actively engaged workers (fig. 3), but interestingly, attempted a significantly greater number of tandem runs per individual (fig. 4). Furthermore, ants in this group attempted a significantly greater workload per individual than those in the 'either’ group, when migrating over the longer distance (fig. 5A). However, the data for successful workload performed per individual did not differ significantly between the two groups, at any distance (fig. 5B).

Our data suggest that some individuals are indeed disproportionately active in the tandem running process, specifically attempting more tandem runs per ant overall (fig. 4), and a greater per capita workload when the new nest is further away (fig. 5A). However, they do not perform a greater successful workload per capita within distances (fig. 5B), and as such, the relative importance of these ‘elite’ workers (Hölldobler & Wilson 1990) in shaping emigration dynamics is debatable. Instead, it appears that the majority of changes in tandem running at the colony level are achieved via increased recruitment of workers into the process, rather than by reliance on key individuals (fig 2C,D, 5B). Such findings are also supported by previous work, showing that the ability to utilise a larger workforce negates the need for specialists (Dornhaus et al. 2009). This highlights an interesting contradiction; it suggests that although some workers are highly active and conform to the definition of ‘elites’, they may not actually be that effective at their task, despite investing more effort than their nest mates. Hence, in this case at least, the usefulness of the term ‘elite’ is debatable. Indeed, a recent study, also in the Temnothorax genus, found that reliance on specialists in the face of dynamic environments may be disadvantageous (Jongepier & Foitzik 2016), highlighting another case in which the contribution of ‘elites’ is far from decisive.

The presence of inactive or ‘lazy’ workers has received some attention in social insects (Dornhaus et al. 2008; Beshers & Fewell 2001) and may be accounted for proximately by task thresholds, as inactive individuals are likely to be those that require a higher level of stimulus before they engage in a given task (Robinson et al. 2009). However, here we see a case of active but ineffective workers in a task that is moderately difficult to complete (only 65% of all tandem runs were successful). While the exact mechanisms underlying this are as yet unknown, there are several potential causes. One possibility, is that workers attempting a greater number of tandem runs were doing so as a compensatory reaction to failing more often. This is plausible, as interruptions to tandem runs were common in our experiment, as in others (Franks et al. 2009). It is also likely that highly
active individuals made a significant contribution by leading only partially successful tandem runs. This is because, as demonstrated in previous experiments (Franks et al. 2010), even when tandem runs end prematurely, they may still be effective in directing naive workers to a new nest (Franks et al. 2010; Pratt 2008). However, while both of these factors provide viable explanations, their relative importance will likely require further investigation.

In conclusion, we find that several factors contribute to the acceleration of emigrations over longer distances, and that both highly active individuals, and the regulation of total worker involvement play a role. Crucially, we show that the greater number of active workers employed over longer emigrations is the strongest factor linked to increased tandem running, having a greater influence than the utilisation of ‘elite’ workers. Overall, our results indicate that although key individuals have their place in the tandem running process, it is the mobilisation of the masses that likely makes the real difference when emigrating over greater distances.

**Competing interests**

The authors declare that they have no competing interests.

**Author contributions:**

TAOW and NRF conceived the study and wrote the manuscript; TAOW, DKWA and DEE conducted all experimental work; TAOW and ABSF carried out the statistical analysis. All authors gave final approval for publication.

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**Data availability**

Table S1. Raw data from the experiment.

**Ethical standards statement**

All applicable institutional and/or national guidelines for the care and use of animals were followed.

**References**


**Figure legends**

**Figure 1.** Marked *Temnothorax albipennis* workers, though some are missing paint marks, the numbering system was robust to this (A), experimental arena setup for 100 mm trial (B), and 300 mm trial (C). Dimensions indicate actual arena sizes, figure is for illustrative purposes only, and is not to scale.

**Figure 2.** Total number of tandem runs at colony level n\(_{100}\)=10, n\(_{300}\)=10, P=0.028, (A), and percentage of successful tandem runs out of those attempted n\(_{100}\)=10, n\(_{300}\)=10, P=0.329 (B), percentage of colony engaged in tandem running n\(_{100}\)=10, n\(_{300}\)=10, p=0.037 (C), and percentage of colony engaged in the emigration process as a whole (scouting, tandem leading or following) n\(_{100}\)=10, n\(_{300}\)=10, P<0.001 (D), for the two emigration distances. Coloured lines indicate individual colony changes in tandem running over distance, palette is chosen for clarity. Outliers (further than 1.5 times the interquartile range from the median) are indicated with circles.

**Figure 3.** Number of workers involved in tandem running at one, or both, emigration distances, as a percentage of the total number of workers actively engaged for both groups. Actively engaged workers are those involved in scouting, tandem leading or following n\(_{100}\)=10, n\(_{300}\)=10.

**Figure 4.** Total number of attempted tandem runs per ant. Boxplots are divided by emigration distance, and within each distance, by individuals actively engaged at only that distance; the ‘either’ group (flat boxes), or both emigration distances; the ‘both group’ (grey boxes), n=296, (n\(_{100}\):either=76, n\(_{100}\):both=53, n\(_{300}\):either=114, n\(_{300}\):both=53). Asterisked lines highlight significant differences within distances (Bonferroni post-hoc test, P<0.05). Outliers (further than 1.5 times the interquartile range from the median) are indicated with circles.

**Figure 5.** Per capita attempted (A), and successful (B), workload per ant. Boxplots are divided by emigration distance, and within each distance, by individuals actively engaged at only that distance; the ‘either’ group (white boxes), or both emigration distances; the ‘both group’ (grey boxes), n=296, (n\(_{100}\):either=76, n\(_{100}\):both=53, n\(_{300}\):either=114, n\(_{300}\):both=53). Asterisked lines highlight significant differences within distances (Bonferroni post-hoc test, P<0.05), outliers (further than 1.5 times the interquartile range from the median) are indicated with circles.

**Figure 6.** Mean return speed of workers within colonies for the two distances n\(_{100}\)=101, n\(_{300}\)=114, p<0.001, inlay displays changes in mean return speed of individuals tracked across both emigration distances, coloured...
lines indicate each individual’s change in mean return speed $n_{100}=20$, $n_{300}=20$. Outliers (further than 1.5 times the interquartile range from the median) are indicated with circles.
Figure 1.
Figure 2.
Figure 3.
Figure 4.
Figure 5.