Foraging activity and demographic patterns of two termite species (Isoptera: Rhinotermitidae) living in urban landscapes in southeastern Brazil

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Abstract. Coptotermes gestroi and Heterotermes tenuis (Isoptera: Rhinotermitidae) are important pests in southeastern Brazil causing serious economic damage. In this study we determined the demographic patterns and foraging activity of these species using mark-release-recapture and the consumption of wooden stakes. Using both the weighted mean and Lincoln index methods, population estimates ranged from ≈ 0.57 to 1.99 million individuals for C. gestroi and from ≈ 0.20 to 1.37 million for H. tenuis. Territory size of the colonies ranged from 172.5 to 5235 m² for C. gestroi and from 16 to 40 m² for H. tenuis. Our results also indicate that foraging activity was dependent on the minimum temperature; however, the existence of a compensation strategy in the foraging activities may permit foragers to exploit food sources under different environmental conditions.

INTRODUCTION

Subterranean termites (Rhinotermitidae) are known to form large colonies that may contain thousands of individuals. The colonies of these termites are composed of workers, soldiers, nymphs, and both primary and secondary reproductives. The workers are the more abundant caste and are responsible for most colony tasks including foraging, construction and repair of the nest, larvae and soldier feeding, and colony defense. Demographic parameters of termite colonies have been estimated by different methods including direct counting of the entire population, exhaustive trapping, and mark-release-recapture. Direct counting of the individuals of an entire colony has been used to determine population size in mound and tree-nesting species (Nutting & Jones, 1990). However, colonies of subterranean termites are generally located some distance from food sources (Evans et al., 1999). Furthermore, some species of subterranean termites form satellite nests and, thus, estimating their colony size by this method is difficult.

A colony of subterranean termites can simultaneously exploit several food sources, which are connected with the nest by a complex of tunnels and galleries. Consequently, the foraging population can provide a good estimate of colony size and territory range. Mark-release-recapture methods are currently used to estimate the size of the foraging population of subterranean termites (Su & Scheffrahn, 1988; Grace et al., 1989; Jones, 1990; Nutting & Jones, 1990; Haagsma & Rust, 1995). These methods make certain assumptions, which according to Evans et al. (1999) are frequently violated in studies of both tropical and temperate termites, and result in overestimates of population size (Forschler & Townsend, 1996; Evans et al., 1998).

Wood consumption by termites is influenced by both intrinsic and extrinsic factors, such as competition with other organisms, population size of the colony, food availability, and environmental conditions. Laboratory studies of wood consumption in termites reveal that termites are selective feeders and show a preference for certain types of wood (Waller & La Fage, 1987; Delaplane & La Fage, 1989). However, it is not clear whether the results observed in these studies are mediated by volume and position of the wood, environmental conditions and/or interaction with other organisms. Wood stakes are currently used for monitoring termite colonies in both prevention and control programs of subterranean termites (Su & Scheffrahn, 1996, 1998). Indeed, metabolic inhibitors, chitin inhibitors and juvenoids are sometimes applied to wooden stakes buried in areas infested with subterranean termites (Su & Scheffrahn, 1998; Lee, 2002).

The subterranean termites Coptotermes gestroi was introduced into the southeast region of Brazil in a marine cargo, probably at the beginning of the 20th century (Araujo, 1958). The Asian subterranean termite C. gestroi was accidentally introduced into the southeast region of Brazil in a marine cargo, probably at the beginning of the 20th century (Araujo, 1958).
activity of these species by measuring the consumption of wooden stakes. Such information will increase our understanding of the foraging behaviour and food preferences of these species, with the objective of developing a better pest management strategy.

**MATERIAL AND METHODS**

**Study site and species**

Two colonies of *Coptotermes gestroi* (C1 and C2) and *Heterotermes tenuis* (H1 and H2) were used in this study. The colonies were located in Rio Claro city, SP, Brazil (22°23’ S, 47°31’ W). Extensive damage caused by *C. gestroi* is recognized by the distinct foraging tubes in wooden structures (Su et al., 2000). Colony C1 were located in a residential area in Rio Claro city (C1), whereas colony (C2) was located close to the Center for the Study of Social Insects (CEIS) at UNESP. Colony C2 was located 5 km from colony C1. On the other hand, colonies H1 and H2 were attacking the trunks of dead *Eucalyptus* sp. trees in the garden of the UNESP in Rio Claro city. The foraging territory of H1 was located 500 m from that of H2. Furthermore, colonies H1 and H2 were considered to be independent as the individuals of these colonies showed strong agonistic behaviour to one another (Fig. 1).

**Mark-release-recapture**

A triple mark-release-recapture was used to estimate the size of the foraging population and territory range of the colonies of these species. The colonies were monitored for two months by checking weekly rolls of tissue buried in the ground. Once termite activity was confirmed, the tissue rolls were replaced by cardboard traps (monitoring stations) before the first marking. For *H. tenuis* colonies, new monitoring stations were established in a concentric position around the old ones in a 2 × 2 m grid. For *C. gestroi* colonies, the monitoring stations were placed around the buildings at intervals of 2 m (Fig. 1).

The mark-recapture was conducted during April and May 2001. Termites collected from the monitoring stations with the highest number of individuals were separated from the cardboard and confined in 10 cm Petri dishes. Ten reference samples of 10 workers were taken from each colony to determine individual weight. Body weight (mean ± SE) of an individual worker of colony C1 was 3.10 ± 0.10 mg and for C2 was 3.48 ± 0.10 mg. On the other hand, the body weight (mean ± SE) of workers of colonies H1 and H2 was 2.40 ± 0.10 mg. The termites were marked by feeding them for one week on filter paper impregnated with a 0.05% (weight/volume) solution of Nile blue dye. In preliminary experiments, workers of both species showed a uniform dark blue colour after the marking process, which was continued for 40 days. The mean (± SE) percentage survival of workers during the marking process was 98.5 ± 0.5% for *C. gestroi* and 98 ± 1% for *H. tenuis*. In order to avoid the possibility of the transfer of marks only dark blue coloured individuals were used for the population estimates. Mark transfer by cannibalism is reported in some termite species (Evans et al., 1998, 1999), including *C. gestroi* and *H. tenuis* (Costa-Leonardo, unpubl.), and individuals that acquire marks in this way are a lighter colour than those that fed directly on the dye.

Marked termites were weighed and released at the station where they were collected. All the monitoring stations were used for mark/recapture events and termites were collected one week later. Unmarked termites collected at the monitoring stations were weighed, stained and released. Termites were marked, released, and recaptured three times over a period of 42 days. The population estimate was calculated using the Lincoln index and weighted mean method. The Lincoln index is the simplest estimator of population size using marked individuals, involving just one release and one recapture. On the other hand, the weighted mean utilizes data collected on several days. Both methods make the same assumptions (Begon, 1979). The mean foraging population and standard error were calculated averaging the three Lincoln index estimates. Only those recaptured workers that were darkly marked were considered marked in this study. The foraging territory of the colonies of *C. gestroi* and *H. tenuis*, defined as the area encompassed by the monitoring stations, was determined by the presence of marked termites. Maximum linear foraging distance was defined as the linear distance between the outermost monitoring stations at which marked individuals were found.

Behavioural assumptions of mark-recapture were tested before determining the foraging patterns of the two species (Begon, 1979; Thorne et al., 1996; Evans et al., 1998). The assumption of complete mixing of both marked and unmarked individuals was tested by comparing the proportion of marked individuals at each monitoring station. If mixing was uniform, the number of marked individuals should increase at each subsequent recapture. Pearson correlations between the number of marked individuals and the total number of recaptured individuals were calculated. Additionally, if foraging individuals move randomly between feeding sites, an equal proportion of marked individuals would be found at all monitoring stations. Therefore, a χ²-test was used to test if the number of marked individuals varied among the monitoring stations at a particular recapture event (Sokal & Rohlf, 1995).

**Foraging activity**

Foraging activity of *C. gestroi* and *H. tenuis* was determined by the consumption of wooden stakes placed in the foraging territories of four colonies of each species. This experiment was conducted during the summer (December–March) and winter seasons (June–August). Ten *Pinus* sp. and *Eucalyptus* sp. stakes of two sizes (28 cm × 6 cm × 4 cm; 28 cm × 2 cm × 2 cm) were used in each colony. Dry weight was estimated by oven-drying the stakes at 120°C for 12 h. Afterwards, the stakes were buried 23 cm deep in the foraging territories of the termites in 4 × 4 m grids. All the stakes were collected after a three-month exposure to termite attack and oven-dried at 120°C for 12 h. Wood consumption was the difference in the dry weight before and after the experiment. The results of wood consumption were subjected to logarithmic transformation in order to improve the normality and homogeneity of the variance (Sokal & Rohlf, 1995).

An ANOVA was used to analyze mean differences in wood consumption, in terms of maximum depth and minimum depth of the attack, and the effects of the size of stakes and type of wood. Significant differences were separated using a Tukey HSD test (p < 0.05). The percentage of the wooden stakes attacked was correlated with average precipitation and minimum and maximum temperatures (p < 0.05) in order to determine if climatic variables affected foraging activity.

**RESULTS**

**Mark-release-recapture**

The number of marked termites released, number of termites captured, and number of marked termites among those recaptured during the triple mark-release-recapture are summarized in Table 1. In the colonies of both species, the mean percentage (± SE) of marked individuals recaptured was 1.27 ± 0.37% for *C. gestroi* and 1.69 ± 1.1% for *H. tenuis*. Marked workers of *C. gestroi* were recovered from monitoring stations encompassing
areas of approximately 172–5200 m² and for *H. tenuis* of 16–40 m² (Fig. 1, Table 1).

The Lincoln index estimates of *C. gestroi* colonies ranged from ≈ 0.57 to 1.81 million workers, while the weighted mean estimates ranged from ≈ 0.65 to 1.99 million workers and of *H. tenuis* colonies, from ≈ 0.20 to 0.93 and ≈ 0.28 to 1.37, respectively (Table 2). Workers of *C. gestroi* travelled a maximum distance of approxi-
mately 24 to 133 m between the outermost foraging stations and for H. tenuis it was 7 to 16 m (Table 2). Live weight of the foraging populations of C. gestroi ranged from 2.27 to 6.35 kg and from 0.68 to 3.29 kg for H. tenuis.

Workers of both species did not mix uniformly within the foraging territories. For two colonies, there was no significant increase in the number of marked individuals against the total number of individuals recaptured at the monitoring stations (C. gestroi: r = 0.2857; p = 0.583) (H. tenuis: r = −0.2636; p = 0.614). Moreover, the proportion of marked individuals was not homogenous in the foraging territories of the colonies of C. gestroi (C1: χ² = 494.24, df = 8, p < 0.0001; C2: χ² = 289.52, df = 13, p < 0.0001) and H. tenuis (H1: χ² = 584.89, df = 5, p < 0.0001; H2: χ² = 41.73, df = 3, p < 0.0001).

### Foraging activity

Foraging activity of C. gestroi and H. tenuis colonies (measured by the number of stakes attacked by termites) appeared to be dependent on temperature. On average, a greater percentage of the wooden stakes were attacked in summer (24.24%) than in winter (9.52%). Analysis of the percentage of stakes attacked and environmental factors (Fig. 2) indicated that minimum temperature was significantly related to the number of stakes attacked both in summer and winter (R² = 0.878, F(1,10) = 28.81, p = 0.006, N = 12). Maximum temperature and precipitation did not influence the foraging activity of either species of termite (Maximum temperature: R² = 0.641; F(1,10) = 7.15; p = 0.056, N = 12. Precipitation: R² = 0.086; F(1,10) = 5.16; p = 0.056, N = 12).

The total mean consumption of the two termite species, observed over a 3-month period, did not differ significantly (F(1,21) = 3.55, df = 1, p = 0.073, N = 24). The wood consumption was 32.99 ± 16.41 g for C. gestroi and 18.02 ± 4.51 g for H. tenuis. Non-significant differences were also found in total consumption in the two seasons (F(1,21) = 2.95, df = 1, p = 0.101, N = 24). Conversely, there were significant differences in the type of wood (C1: R² = 0.641, df = 1, p < 0.0001, N = 24) and size of the stakes consumed (F(1,21) = 27.43, df = 1, p < 0.0001, N = 24). The mean consumption of Pinus sp. (26.76 ± 6.86 g) was significantly greater (p = 0.001; Tukey HSD test) than of Eucalyptus sp. stakes (5.76 ± 2.50 g). Furthermore, termites of both species significantly consumed more of the large stakes (33.71 ± 9.36 g) than the small stakes (9.00 ± 2.42 g) (p = 0.0002; Tukey HSD test).

Maximum mean depth of attack did not differ significantly between termite species (F(1,21) = 1.64, df = 1, p = 0.214, N 24). The maximum mean depth of attack was 24.86 ± 2.30 cm for C. gestroi and 27.53 ± 0.36 cm for H. tenuis. Non-significant differences were found in the maximum mean depth between seasons (F(1,21) = 0.275, df = 1, p = 0.605, N = 24). The maximum mean depth of the attack in summer was 26.13 ± 1.07 cm and 28.00 ± 0.00 cm in winter. Moreover, minimum mean depth of attack was similar for both termite species (F(1,21) = 0.10, df = 1, p = 0.751, N = 24). The minimum mean depth of attack was 7.71 ± 2.74 cm for C. gestroi and 8.29 ± 1.10 cm for H. tenuis. There were no significant differences in the minimum mean depth of attack between the two seasons (F(1,21) = 0.06, df = 1, p = 0.804, N = 24). The minimum mean depth of the attack was 8.22 ± 1.54 cm in summer and 7.94 ± 1.18 cm in winter.

### DISCUSSION

Table 3 summarizes the published estimates of the size of the foraging populations of colonies of some species of subterranean termites obtained using the Lincoln index and weighted mean methods. There are differences in the population sizes, territory sizes, and maximum linear distance travelled by the workers of the different termite species. The range of colony sizes recorded for some species may indicate a range of age classes affected by environmental conditions, competition with other termites, colony health and the number of monitoring stations, or the method used to estimate the size of the foraging population (Waller & La Fage, 1987; Lenz, 1994). The popu-

### Table 1. Number of marked termites released (ri), number of termites captured (ni), number of marked termites among those recaptured (mi) and estimated territory size (ti) obtained by the triple mark-release-recapture method.

<table>
<thead>
<tr>
<th>Colony</th>
<th>ri</th>
<th>ni</th>
<th>mi</th>
<th>ti (m²)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>11659</td>
<td>8214</td>
<td>30</td>
<td>114.8</td>
</tr>
<tr>
<td>C2</td>
<td>5625</td>
<td>5819</td>
<td>194</td>
<td>256</td>
</tr>
<tr>
<td>H1</td>
<td>2601</td>
<td>6686</td>
<td>103</td>
<td>40</td>
</tr>
<tr>
<td>H2</td>
<td>4331</td>
<td>5289</td>
<td>22</td>
<td>4</td>
</tr>
</tbody>
</table>

### Table 2. Estimates of the foraging populations and maximum linear foraging distance of C. gestroi and H. tenuis, living in colonies located in urban areas.

<table>
<thead>
<tr>
<th>Colony</th>
<th>Estimator method</th>
<th>Population (+ SE)</th>
<th>Linear Distance (m)</th>
</tr>
</thead>
<tbody>
<tr>
<td>C1</td>
<td>Weighted mean</td>
<td>1995427 ± 68923</td>
<td>23.57</td>
</tr>
<tr>
<td></td>
<td>Lincoln Index</td>
<td>1811768 ± 230426</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Weighted mean</td>
<td>655127 ± 35740</td>
<td>133.33</td>
</tr>
<tr>
<td>C2</td>
<td>Lincoln Index</td>
<td>571194 ± 38560</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Weighted mean</td>
<td>286678 ± 10559</td>
<td>16</td>
</tr>
<tr>
<td>H1</td>
<td>Lincoln Index</td>
<td>200739 ± 13832</td>
<td></td>
</tr>
<tr>
<td></td>
<td>Weighted mean</td>
<td>1372923 ± 103198</td>
<td></td>
</tr>
<tr>
<td>H2</td>
<td>Lincoln Index</td>
<td>933235 ± 135706</td>
<td>7.14</td>
</tr>
</tbody>
</table>
lation estimates of *C. gestroi* and *H. tenuis* are lower than those for other *Coptotermes* species, nevertheless, the colonies of *C. gestroi* and *H. tenuis* were larger than those of the Nearctic termites *Reticulitermes virginicus* and *R. flavipes*.

Both methods used to estimate the population sizes of *C. gestroi* and *H. tenuis* using triple-mark-recapture gave different results. In general, the mean Lincoln index estimates were lower than the weighted mean estimates for all the colonies studied. The inflated Lincoln index estimates obtained for other subterranean species (Table 3) may have resulted from very low recapture rates of marked individuals (less than 1%). Lincoln index estimates are calculated for each mark-release-recapture and a low number of recaptured individuals at a single census, may increase the population estimates. Nevertheless, most of the studies using these methods do not give the number of marked individuals recaptured.

The low number recaptured could be due to some stained individuals loosing their marks. In this study the mark persisted, survival of marked individuals was close to 100% and only darkly stained individuals were used for the population estimates. Foraging behaviour in subterranean termites is a collective process, which minimizes energetic costs (Hedlund & Henderson, 1999), and depends on either the size or quality of the food source (Waller & La Fage, 1987). Moreover, age polyethism in the worker caste could also affect the number of individuals recaptured when additional medium and large workers are recruited for foraging tasks. This might also explain the absence of a correlation between the number of marked individuals and the total number of individuals recaptured at a monitoring station.

Violations of behavioural assumptions of mark-release methods have implications for the interpretation of the estimates of population size (Curtis & Waller, 1997); however, foraging patterns of species of subterranean termites are almost identical and quantitative comparisons are usually performed among species of termites in most studies. In addition, mark-release provides information on territory size and fluctuations in demography, and thus can be used to improve the control and management techniques in both urban and rural landscapes (Su & Scheffrahn, 1996, 1998; Forschler & Jenkins, 2000; Su, 2002).

Climatic factors are important regulators of foraging activity in termites in southeastern Brazil, where temperature and rainfall differ in summer and winter. The percentage of stakes attacked was positively related to the mean minimum temperatures recorded in summer and winter. In this region, differences in the mean minimum temperature between these seasons range from 12 to 14°C. This is consistent with the results of Haagsma & Rust (1995), who found that minimum temperature was the most important factor affecting activity in *R.*

**Table 3.** Published estimates of the foraging populations of subterranean termites, territory size (m²), and maximum linear distance (m) travelled by workers.

<table>
<thead>
<tr>
<th>Species</th>
<th>Estimator method</th>
<th>Population</th>
<th>Territory</th>
<th>Distance</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Reticulitermes flavipes</em></td>
<td>Weighted mean</td>
<td>970 – 1453021</td>
<td>266 – 1091</td>
<td>48 – 79</td>
<td>Forschler &amp; Townsend (1996); Grace et al. (1989)</td>
</tr>
<tr>
<td></td>
<td>Lincoln Index</td>
<td>1111 – 384615</td>
<td>–</td>
<td>–</td>
<td>Forschler &amp; Townsend (1996)</td>
</tr>
<tr>
<td><em>R. virginicus</em></td>
<td>Weighted mean</td>
<td>20109 – 156997</td>
<td>–</td>
<td>–</td>
<td>Forschler &amp; Townsend (1996)</td>
</tr>
<tr>
<td></td>
<td>Lincoln Index</td>
<td>18893 – 153945</td>
<td>–</td>
<td>–</td>
<td>Forschler &amp; Townsend (1996)</td>
</tr>
<tr>
<td></td>
<td>Lincoln Index</td>
<td>3846329 – 18834481</td>
<td>–</td>
<td>–</td>
<td>Evans et al. (1998)</td>
</tr>
<tr>
<td><em>C. lacteus</em></td>
<td>Weighted mean</td>
<td>11718537 – 33327481</td>
<td>–</td>
<td>–</td>
<td>Sornmuwat et al. (1996)</td>
</tr>
</tbody>
</table>
hesperus. However, environmental factors may have a different effect on the activity of the termites, as described for other subterranean species (Delaplane et al., 1991; Haagisma & Rust, 1995). In the present study, wood consumption was similar in summer and winter, which suggests the existence of a compensation strategy in the foraging activities in both seasons, as described for C. formosanus (Waller & La Fage, 1987). On the other hand, there is a trend that consumption differs between termite species; however, other factors such as population size, food availability or competition could influence this result.

In this study, stakes of Pium sp. (low density wood) were more readily eaten than those of Eucalyptus sp. (high density wood). Field studies of wood consumption are scarce and information on termite food preferences is limited to laboratory experiments in which termites are offered a limited range of different food sources (Lenz, 1994). These studies indicate that C. formosanus prefers low density and moist wood or wood damaged by conspecifics (Delaplane & La Fage, 1989). On the other hand, food choice experiments using Reticulitermes sp. indicate that this termite prefers woods that contain low quantities of tannin and terpenoids, which are toxic to insects (Ripa et al., 2002).

According to Delaplane & La Fage (1987), once termite foragers chose a particular food source, they exploit it to the exclusion of all others. Moreover, termit feeding seems to be associated with the size of the food source (Waller, 1988; Lenz, 1994), as we found for C. gestroi and H. tenuis, in which foragers feed significantly more on the larger stakes. The low number of stakes attacked by termites in the foraging territories of field colonies of C. gestroi and H. tenuis may be due to the presence of alternative food sources, which were considerably larger than the stakes used in this study. However, other factors could have influenced wood consumption in these species, such as the tunnel construction pattern of the foragers, soil particle size and available water.

The results obtained for C. gestroi and H. tenuis did not show differences in the extent to which wooden stakes buried at different depths were attacked. Some termite species attack mainly stakes barely buried in the soil (Shahid & Akhtar, 1989). Nevertheless, the difference in the soil depth at which they feed might be species specific or the stakes used in this study were too short.

Demographic patterns of C. gestroi and H. tenuis varied between colonies and species. Even though for subterranean termites some of the behavioral assumptions of mark-recapture methods are not met, this study showed that colonies of C. gestroi and H. tenuis are larger than those reported for Nearctic species. Even though, the foraging activity of these termites depended on the minimum temperature, wood consumption did not change between summer and winter. These results could indicate the presence of a compensation strategy in foraging, which is dependent on the minimum temperature. This information will be useful for improving the control of both species of termites. Nevertheless, further studies aimed at understanding the searching patterns of these subterranean species are being conducted under controlled laboratory conditions.

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