Sweeping the house clean: burrow architecture and seasonal digging activity in the East African root rat from Tanzania

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Abstract
We investigated changes in burrow architecture and fractal dimension across seasons and between the sexes in the solitary East African root rat Tachyoryctes splendens over an entire calendar year. The basic burrow system comprised a main tunnel reticulating into foraging tunnels, a nest consisting of food store chamber, latrine and sleeping area, and a bolt hole. Main tunnel length was strongly affected by sex, and contrary to expectations, it was longer for females than for males (during both the dry and the wet seasons). The number and the length of foraging tunnels were affected by both sex and season, with females’ burrows having more foraging tunnels than males in both the dry and the wet seasons. Females also had burrows with higher fractal dimension than males, while fractal dimension increased with burrow length for both sexes. We suggest that unlike the solitary bathyergid mole-rats, male T. splendens do not construct larger burrows than females in the search for mates, but rather females have larger burrows with more foraging tunnels resulting from the increased need for provisioning of their young.

Keywords
Tachyoryctes splendens; burrow architecture; subterranean rodent; mole-rat; fractal dimension; root rat.
Introduction

The East African root rat, *Tachyoryctes splendens* (Family: Spalacidae) is a solitary dwelling fossorial rodent, widely distributed across East Africa. The Spalacidae are classified within the Myomorph suborder of Rodentia, and are divergent from better-known African mole-rats, (Family: Bathyergidae, Suborder: Hystricomorpha). While overlapping geographical ranges with several bathyergid mole-rat species, *Tachyoryctes* has convergently adopted a subterranean lifestyle, and investigation of comparative distributions and habitat use are of broad interest. As with other rodent moles, *T. splendens* feeds on underground plant storage organs and roots (which will therefore influence burrow structure and dynamics), but it is also known to occasionally forage above ground (Jarvis & Sale 1971). Where they are associated with farmland, *T. splendens* is a major pest species (Kokiso & Bekele, 2008). Previous excavation of small numbers of adult burrows in Kenya have show that the typical architecture comprises of a main tunnel with a number of superficial foraging tunnels branching from this, a deeper bolt hole and a centrally-positioned multi-purpose nest chamber which comprises a food store, latrine and sleeping area (Jarvis & Sale 1971). Following a study of two adult and two juvenile burrows during flooding, Hickman (1983) also suggests the bolt hole may also act as a drainage sump during heavy rains. Although a common subterranean species in areas where present, no comprehensive study of *T. splendens* has been carried out to investigate the detailed structure and geometry of burrows with respect to the sex of the occupant, and seasonally-induced changes (e.g. over wet and dry seasons).

Most burrowing activity in subterranean species occurs following rainfall when there is increased soil moisture that makes the soil more friable and workable, and it is also easier to extrude the soil workings onto the surface as mounds (Miller, 1957). Other factors that may also
be related to rainfall and which can seasonally influence burrowing patterns are mate acquisition and foraging (Miller & Bond, 1960; Hickman & Brown, 1973). Thus, in species occurring in regions of marked wet and dry seasons, as is the case in many African mole-rats (Family: Bathyergidae), some aspects of the architecture of the burrow system change seasonally and are influenced by both biotic and abiotic factors. Burrows may be extended for: a) the primary purpose of food acquisition, especially in the case where food resources are geophytic and occur below ground; or b) tunnelling may be triggered for the purpose of mate searching, or during dispersal, for example, when a juvenile leaves the maternal burrow system as is the case in solitary species, or when emigration occurs from a parent colony in social species (Bennett & Faulkes, 2000).

Among solitary bathyergid mole-rats, males seek out mates and as a consequence they construct longer tunnel systems, communicating with and attracting females in the subterranean niche via seismic signals (Bennett & Jarvis, 1988; Herbst & Bennett, 2006; Le Comber et al., 2006; Thomas et al., 2009, 2012). We therefore predicted that the burrow systems of male and female *Tachyoryctes* would show sex differences reflecting sex-specific mate seeking behaviour. We further predicted that the burrow systems of both sexes of *T. splendens* would be longer and more reticulated during the wet season as they actively search for food items.

The complexity of a burrow system and the exploration efficiency of the surrounding environment, or reticulation of the system, can be examined through fractal dimensions (Kenkel & Walker, 1996). Fractal dimension provides a measure of burrow shape since it provides a measure of the extent to which a one-dimensional structure fills a plane, with a low fractal dimension (i.e. close to 1.0) describing a burrow that explores relatively little of the surrounding area whereas a high fractal dimension (i.e. close to 2.0) designates a burrow which permeates the
surrounding area more thoroughly (Romañach & Le Comber, 2004). Fractal dimension is thus a valuable measure of burrow shape when the burrow is used for foraging (Le Comber et al. 2006). While there are no differences in the general burrow architecture between seasons in the solitary bathyergid, the silvery mole-rat, *Heliophobius argenteocinereus*, their burrows had higher fractal dimension at the peak of the dry season which suggests the burrow occupant increases its search for food (Sumbera et al., 2003). Similarly, in the social giant Zambian mole-rat, *Fukomys mechowii*, total burrow length did not vary with season, but the fractal dimension was greater during the rainy season than the dry season. This suggests greater exploratory tunnelling without an overall increase in total tunnel length, which increases the chances of finding food resources (Sichilima et al., 2008).

Thus in subterranean rodents, while the overall home range remains relatively constant, burrow systems are usually in a state of dynamic flux in response to environmental cues and mate acquisition. This is the first study to undertake an extensive survey of *T. splendens* in Tanzania, excavating burrow systems on a monthly basis over an entire calendar year. The objectives were (i) to test the prediction that burrow size and fractal dimension were larger in males, and (ii) to quantify seasonal differences with the prediction that increases in burrow length, the number of foraging burrows and fractal dimension occur in the wet season.

**Materials and methods**

*Study site*

This study was conducted over 12 consecutive months from January to December 2011, in farmlands at Mamba Komakundi village, located on the slopes of Mount Kilimanjaro in the Moshi Rural District, north-east Tanzania (03°16.54' S, 037°32.49' E; 1495 m above sea level).
The site is characterized as an agro-ecological zone with very fertile volcanic soils supporting a variety of food and cash crops (Kilimanjaro Regional Profile (KRP), 1998). The area has two rainy seasons with a mean annual rainfall of 1250 mm. The long rainy season is between March and June, while the short rainy season is between October and December. The dry season spans from January to early March and from July to early October. Annual average temperature ranges from 15-30° C with high humidity during October, March, April and May (KRP, 1998).

Excavation of burrow systems

Burrow systems were excavated manually with hoes to expose the tunnels along their entire length. A total of 60 burrows were excavated (i.e. five burrow systems per month). During each month we captured *T. splendens* the animals were weighed and sexed prior to excavating the burrow system. Equal numbers of each sex were collected (15 males and 15 females) from wet and dry seasons respectively. Animals caught and removed from burrows were used as samples for other on-going parallel studies.

The lengths of the tunnels and their dimensions and shape were recorded *sensu* Thomas *et al.* (2009; 2012) and measured to the nearest 0.1 cm using a tape measure as follows: the depth from the ground surface to the top of the tunnel (recorded approximately every 2 m), height and width of the tunnel, the length of the main tunnel and the length of foraging tunnels. The number of foraging tunnels was also noted, and burrow fractal dimension calculated as detailed below. Tunnels were defined as either: deep, semi-permanent (> 20 cm) or shallow, foraging (< 20 cm) tunnels (determined by the depth of the bulbs and roots of the plants in the sites reached). The location and dimensions of any nests, food stores, bolt holes and latrines were also recorded. Nests were defined as chambers with only a single entrance and filled with nesting material. Food stores were blind-ended tunnels filled with either bulbs or roots and the contents of these
caches were recorded as percentages of food type. Bolt holes were steep-angled tunnels (almost vertical) that were > 30 cm in length used as either anti-predatory function, thermoregulation or as drainage sumps (Hickman, 1983, 1990; Nevo, 1999). Latrines were blind-ended tunnels packed with soil and faeces. The study was approved by the Animal Ethics Committee of the University of Pretoria (Ethics clearance number ECO47-10), the Tanzania Commission for Science and Technology (COSTECH) (Ref. 2011-44-NA-2010-204) and the University of Dar es Salaam, Tanzania.

Data analyses

To quantify the geometric configuration of the burrow in a way that was independent of burrow size we calculated fractal dimension. This provides a measure of the extent to which the burrow explores the surrounding area, particularly in subterranean mammals where the burrow is primarily used for foraging (Le Comber et al., 2002, Šumbera et al., 2003; Romañach & Le Comber, 2004). All previous studies on mole-rat tunnel systems have assessed 2-D rather than 3-D structures, since the vertical variation is very slight (perhaps around 1 m). Since this study was primarily concerned with the burrow and its relation to foraging, we also only quantified how the burrows are distributed in 2-D. The geometry of the burrow systems was mapped to a scale of 1:100 on graph paper for estimation of fractal dimension (Le Comber et al., 2002; Sichilima et al., 2008; Thomas et al., 2009, 2012). Each drawing was then digitized separately and subjected to image-processing software GIMP version 2.6.12 (http://sd-cf.en.softonic.com) and Inkscape version 0.48.3.1(http://www.flyplayerpro.net), for re-draw and to convert the burrow system image into a readable format for 2-D fractal dimension calculation using Fractal Dimension Calculator version 1.2.0 (Thomas et al., 2012).
Data from the short and long rainy periods and the two intervening dry seasons respectively were combined to produce “wet” and “dry” season comparisons, to simplify the analysis and increase sample size. Other environmental factors such as temperature do not differ between the short and long rainy/dry periods. All data were then analysed using R (R Core Team, 2012). The effects of sex and season on burrow length (i.e. length of the main tunnel), number of foraging tunnels and foraging tunnel length were analysed using a two-way analysis of variance, including a sex*season interaction. A similar approach was taken to study factors affecting burrow fractal dimension, but in this case burrow length, number of foraging tunnels and foraging tunnel length were included as additional predictors in a general linear model. Data were summarised in boxplots generated in R, showing ‘hinges’ around the median (versions of the first and third quartiles), and notches showing 1.58 times the interquartile range / √n which approximately correspond to 95% confidence limits (R Core Team, 2012).

**Results**

*Burrow structure*

A single animal was sampled in each burrow system excavated, except for five adult lactating females that had pups with them in their burrows. The burrow systems comprised a bolt hole, multi-functional nests and foraging side branches ending at mole hills diverging from a main tunnel (Figure 1). The food store chamber was located on the side of the nest and connected to the nest by one exit point. In the food store, the composition of the cache was rich in roots, tubers, bulbs and rhizomes, as well as grasses and shoots of forbs that may either have been pulled into the burrow tunnel, or harvested during foraging above ground.
Several nest chambers were excavated within individual burrow systems of *T. splendens* but only one was active at any particular time, with other nest chambers back-filled with soil and no longer in use. An active nest was connected to the main tunnel by a single entry/exit point. Within the nest chamber, dry plant materials were found proximal to the entrance, while the latrine was found in the distal section. Nesting comprised of 90.4% non-cultivated vegetation such as grasses and forbs, 5.5% crop materials, 4% faecal pellets and 0.09% invertebrates. Invertebrates observed in the nest chamber included spiders (54.4%), termites (20%), earthworms (7.1%), flies (6.5%), ants (6.2%) and beetles (5.8%). In the same section of the tunnel, disused nests contained dry plant material and old faecal pellets mixed with soil.

Foraging tunnels originated from the main tunnel (Figure 1), and in some burrow systems they branched into several peripheral tunnels and terminated at mole mounds. These mole mounds were characterized by both old and freshly excavated piles of soil, the latter indicating recent burrowing activity. The diameter of the foraging tunnels was large enough to allow easy movement of the occupant without excessive digging. The burrow system of *T. splendens* also contained a single bolt hole which was located at the end of the main tunnel, and at a greater depth than the nest chamber.

*The effects of sex and season on burrow structure*

Burrow length (total length of the main tunnel) was strongly affected by sex, but not season (ANOVA: sex: F_{1,56} = 14.80, p = 0.0003; season: F_{1,56} = 0.65, p = 0.424; sex*season: F_{1,56} = 0.15, p = 0.70), and was higher for females than for males (Figure 2). Means ± SD (range): male
wet: 6.64 ± 1.12, (5.00–8.50), male dry: 6.16 ± 1.06, (4.51–8.50); Female wet: 8.17 ± 2.17, (5.85–13.51), female dry: 7.99 ± 2.16, (5.85–11.50).

The number of foraging tunnels was affected by both sex and season (ANOVA: sex: F1,56 = 20.46, p < 0.0001; season: F1,56 = 6.50, p = 0.014; sex*season: F1,56 = 0.17, p = 0.68). Females’ burrows had more foraging tunnels than males’ burrows in both the dry and wet seasons, while overall burrows had more foraging tunnels in the wet season than the dry season (Figure 3). Means ± SD (range): male wet: 5.13 ± 0.92, (4.00–6.00), male dry: 3.93 ± 0.96, (3.00–5.00); Female wet: 6.80 ± 2.08, (4.00–10.00), female dry: 5.93 ± 1.94, (4.00–8.00).

Foraging tunnel length showed a slightly different pattern, with significant differences due to sex and sex*season (ANOVA: sex: F1,56 = 38.15, p < 0.0001; season: F1,56 = 0.79, p = 0.378; sex*season: F1,56 = 5.83, p = 0.019). Tukey post-hoc tests showed that overall females had burrows with longer foraging tunnels than males, in both the wet and dry seasons (Figure 4). Means ± SD (range): male wet: 4.96 ± 2.36, (2.52–7.63), male dry: 3.35 ± 1.46, (2.52–6.19); Female wet: 6.79 ± 1.45, (5.21–8.63), female dry: 7.53 ± 2.10, (5.18–9.82).

Fractal dimension was strongly affected by both sex and burrow length. Females had burrows with higher fractal dimension than males (Figure 5), while fractal dimension increased with burrow length (Figure 6; results from GLM: sex: F1,53 = 59.49, p < 0.0001; season: F1,53 = 0.35, p = 0.56; sex*season: F1,53 = 2.43, p = 0.13; burrow length: F1,53 = 77.29, p < 0.0001; number of foraging tunnels: F1,53 = 0.03, p = 0.87; foraging tunnel length: F1,53 = 1.56, p = 0.22). Means ± SD (range): male wet: 1.19 ± 0.04, (1.15–1.25), male dry: 1.17 ± 0.04, (1.14–1.24); Female wet: 1.23 ± 0.04, (1.18–1.29), female dry: 1.24 ± 0.04, (1.16–1.27).
A comparison of overall burrow fractal dimensions (FD) between *T. splendens* (mean ± SD = 1.21 ± 0.08) and other previously studied solitary bathyergids including the silvery mole-rat (*Heliophobius argenteocinereus*) = 1.32 ± 0.05, the Cape mole-rat (*Georychus capensis*) = 1.27 ± 0.08, and the Cape dune mole-rat (*Bathyergus suillus*) = 1.35 ± 0.12 (Le Comber *et al.* 2002) showed fractal dimension to be similar.

**Discussion**

Our prediction that burrow length and associated burrow architecture would differ between the sexes was supported, but interestingly, and in marked contrast to the bathyergids, we found that females, rather than males, had longer burrow systems (length of the main tunnel), together with more foraging tunnels of greater length and higher fractal dimension (Herbst & Bennett, 2006; Thomas *et al.*, 2009, 2012). We speculate that sexual dimorphism in burrow geometry of *T. splendens* is most likely attributed to maternal care of pups and the associated increase in food provisioning (and therefore digging) that is required when the female has juvenile young co-habiting the burrow. The greater fractal dimension of the burrows of female *T. splendens* is noteworthy since in other solitary mole-rats such as *Bathyergus suillus* (Thomas *et al.*, 2009) and *Bathyergus janetta* (Herbst *et al.*, 2006), and solitary subterranean rodents *Thomomys bottae* (Reichman *et al.*, 1982) and *Ctenomys minutus* (Gastal, 1994), it is the male burrows that have greater complexity, which may reflect mate searching strategies. A recent study has shown that in *T. splendens*, seismic signalling using head raps against the tunnel roof may be used to communicate for purposes such as mate attraction (Hrouzkova *et al.*, 2013). Such a mechanism, coupled with close proximity of male and female burrows, may mitigate against mate searching by burrowing alone. Seismic signalling has also been suggested as a mate attraction mechanism
in the solitary bathyergid, *Georychus capensis*, which uses drumming of the hind feet to create seismic vibrations (Bennett & Jarvis, 1988). While the overall burrow geometry of *T. splendens* is similar to the solitary bathyergids in terms of range of values recorded for fractal dimension, reflecting aspects of their similar lifestyles, interesting differences (e.g., a more complex nest and the aforementioned sex differences) undoubtedly reflect their convergent exploitation of the subterranean niche.

The effect of sex on burrow architecture that we observed tended to override the predicted seasonal differences, but the general trend was as expected, in that digging activity was greater during the rainy seasons. Such differences between wet and dry periods were clearer for male burrows, possible because increased provisioning by females was also necessary during the dry seasons. (Figures 2-5). Pregnancies were observed to peak during both rainy seasons and were occasionally recorded in the dry period in June and July (Katandukila *et al.*, 2013), thus females may be found either lactating or provisioning young over much of the year, with the accompanying need for increased foraging. Both males and females dug more foraging tunnels in the wet periods, although this was only reflected in a greater mean foraging tunnel length for males in the wet versus the dry periods. Greater burrowing activity during the wet season resulting in an increasing number and length of foraging tunnels has been noted previously for subterranean rodents (Andersen, 1987; Šumbera *et al.*, 2003; Šklíba *et al.*, 2009) as the soil is more workable (Jarvis & Sale 1971; Vleck 1981; Reichman & Smith, 1990; Antinuchi & Luna, 2006; Šklíba *et al.*, 2011) and the food quality and quantity increases from sprouting and regeneration following the rains (Bennie, 1991; Barber, 1995; Pregitzer & King, 2005). It has been found that during foraging, straight paths are more efficient for finding food resources (Zollner & Lima, 1999). However, once resources are located, increased reticulation may result
as a function of the vegetative propagation of tubers, bulbs and corms (Bennett & Faulkes, 2000). Increased fractal dimension of the burrow system has been linked to the period of rainfall in the solitary mole-rats, *B. suillus* and *B. janetta*, (Herbst *et al.*, 2006, Thomas *et al.*, 2009, 2012) as well as in the social giant Zambian mole-rat, *F. mechowii* (Sichilima *et al.*, 2008).

The extensive survey of *T. splendens* burrows undertaken for this study confirms earlier reports with respect to their general structure, which was consistent across all samples (multi-function nest, main tunnel, foraging tunnels and bolt hole; Jarvis & Sale 1971; Hickman, 1983; Kokiso & Bekele, 2008). The burrow systems of *T. splendens* are inhabited by a single individual, with plural occupancy either during breeding or when females have young. In this study, the burrow systems of five females were found to include young individuals. Although adopting a similar lifestyle and found across similar ranges to bathyergid mole-rats, the burrow system of *T. splendens* shows some marked differences, particularly with respect to the multi-functional nest incorporating a sleeping area, food store chamber and latrine area. The food store chamber in the burrow system of *T. splendens* is located within the sides of the active nest chamber and is comprised mainly of grasses, forbs and underground plant organs including tubers, roots, rhizomes, and bulbs. Similar observations were recorded for *T. splendens* in Kenya and Ethiopia (see Jarvis & Sale, 1971; Hickman, 1983; Kokiso & Bekele, 2008). Interestingly, in the abandoned food stores some bulbs and tubers were found sprouting. The storage of geophytes and crop materials suggests that *T. splendens* have similar dietary needs to other herbivorous subterranean rodents such as bathyergids, geomyids, ctenomyids and Asian spalacids (Keith *et al.*, 1959; Jarvis & Sale, 1971; Huntly & Reichman, 1994; Bennett & Faulkes, 2000; Rosi *et al.*, 2000; Spinks *et al.*, 2000; Šumbera *et al.*, 2004; Romañach, 2005; Herbst & Bennett, 2006; Kokiso & Bekele, 2008; Šklíba *et al.*, 2009; Šklíba *et al.*, 2011).
A series of shallow disused nests were observed in the tunnel before the active nest chamber. In fossorial animals, the presence of multiple nests along a tunnel is common and has been reported in the burrow systems of silvery mole-rats, *Heliophobius argenteocinereus* and Cape dune mole rats, *B. suillus* (Hickman, 1977; Šumbera et al., 2003; Šklíba et al., 2009, 2011; Thomas et al., 2009). Multiple nests are reported to be a mechanism for reducing ecto-parasite infestation within burrows. The active nest chamber is identified by the presence of dry grasses and the husks of geophytes, located at the proximal part of the nest entrance. Within the nest chamber, all adjacent above ground geophytes had been consumed. This indicates that foraging also occurs from within the nest chambers of *T. splendens*, as reported for *Geomys* sp. and bathyergids (Jarvis & Sale, 1971; Hickman, 1983; Andersen, 1987; Bennett et al., 1988; Bennett & Faulkes, 2000).

Apart from the food store and sleeping site in the nest chamber, accumulations of faecal pellets were recorded at the distal part of this chamber, forming a latrine. The fermenting faeces and decomposition of nesting material in the burrows of *T. splendens* raises the temperature in their underground chambers (Jarvis & Sale, 1971; Flynn, 1990; Nevo, 1999; Nowak, 1999) and concurrently, these decomposed faecal pellets, food remains and nesting add to soil fertility. Thus the nest chamber in *T. splendens* functions as a sleeping site, a latrine and also as a feeding chamber, as has also been reported in the blind mole rat, *Spalax ehrenbergi* (Nevo, 1961; Jarvis & Sale, 1971; Hickman, 1983; Kokiso & Bekele, 2008). Other subterranean rodents such as bathyergids, geomyids and ctenomyids differ in that they have a distinct toilet chamber (Jarvis & Sale, 1971; Hickman, 1977; Šumbera et al., 2003, 2004; Cutrera et al., 2006; Šklíba et al., 2009, 2011; Thomas et al., 2012).
In conclusion, our findings for *Tachyoryctes* show common features with other rodent moles in terms of increased burrowing following rains, but interesting differences in sexual dimorphism of burrow architecture. Species-specific differences in burrow structure were also evident, particularly with respect to their multi-function nest chamber. Across its range, *Tachyoryctes* overlaps with species of the solitary bathyergid mole-rats *Heliophobius* and the social *Fukomys*, especially in Tanzania (Beolchini & Corti, 2004; Faulkes et al., 2010, 2011). Despite this, so far there are no confirmed published reports of any of these three genera cohabiting the same area, and neither did this study find such evidence. Whether this is a result of stochastic or historical processes, or differences in habitat use, remains to be determined.

**Acknowledgements**

We acknowledge funding from the University of Dar es Salaam (UDSM) World Bank Fund (C1B1 of the University of Dar as Salaam, Tanzania. NCB acknowledges funding from the South African DST/NRF SARChI Chair for Mammal Behavioural Ecology and Physiology, University of Pretoria.

**References**


Figure 1: Representative burrow systems of *Tachyoryctes splendens* at Mamba Komakundi, Kilimanjaro Region, Tanzania. A1 = typical female burrow system; A2 = typical male burrow system; B1 = female burrow system during the wet season; B2 = male burrow system during the wet season; C1 = female burrow system during the dry season; C2 = male burrow system during the dry season; Continuous thick line ( ) = main tunnel; E = entrance; broken line (  ) = foraging tunnel;  = mole mound;  = active nest;  = old/disused nest; L = latrine; S = sleeping area, F = food store chamber, and B = bolt hole.

Figure 2: Box plot of burrow length (length of main tunnel as illustrated in Figure 1) versus sex of burrow occupant for wet and dry seasons. Each boxplot shows “hinges” (boxes) which are versions of the first and third quartiles around the median (bold horizontal lines), and “notches” (vertical lines) showing 1.58 times the interquartile range / √n, which approximately correspond to 95% confidence limits. Outliers are represented by circular plot symbols (R Core Team, 2012).

Figure 3: Box plot of the number of foraging tunnels versus sex of burrow occupant for wet and dry seasons.

Figure 4: Box plot of foraging tunnel length versus sex of burrow occupant for wet and dry seasons.

Figure 5: Box plot of the fractal dimension versus sex of burrow occupant for wet and dry seasons.

Figure 6: The relationship between fractal dimension and burrow length (main tunnel length) for sexes and seasons (male wet: ■, dry ●; female wet: □, dry: ○).
Figure 2

The box plot shows the distribution of burrow length (m) by sex and season. The box plot indicates that female burrow length is generally greater than male burrow length. The dry season has a higher burrow length compared to the wet season for both sexes.
Figure 3

The graph shows the number of foraging tunnels by sex and season. The x-axis represents sex (female, male), the y-axis represents the number of foraging tunnels, and the graph includes data for both dry and wet seasons.
Figure 5

Box plots showing the fractal dimension for female and male sex across dry and wet seasons.

- **Fractal Dimension** range from 1.15 to 1.25 for both sexes.
- **Female** tend to have a higher fractal dimension in the dry season compared to the wet season.
- **Male** show a similar trend, with the dry season having a higher fractal dimension.

**Seasons**:
- **Dry**
- **Wet**

**Sex**:
- **Female**
- **Male**
Figure 6

- dry, male
- dry, female
- wet, male
- wet, female