Chapter 1: Introduction

Rationale and background
Fire is an often observed phenomenon in ecosystems across the world, with up to 40% of the earth’s land surface consisting of fire-prone systems (Chapin et al. 2002), including grasslands, boreal forest, mediterranean shrublands and savannas. Although much control of ecosystem functioning is attributed to climatic factors and resource availability, fire is a very real, yet underestimated, driver of processes related to such functioning (Bond et al. 2005). It is analogous to herbivory in many aspects, acting as a ‘generalist herbivore’ consuming vast amounts of grass and tree biomass and in turn modifying vegetation structure and composition (Bond & Keeley 2005). In savannas in particular, it has been attributed a pivotal role in maintaining the co-existence of trees and grass (Higgins et al. 2000; van Langevelde et al. 2003) and has also been shown to influence other aspects of ecology such as grazing patterns (Archibald & Bond 2004; Archibald et al. 2005; Parrini & Owen-Smith 2009), nutrient cycling (Hobbs et al. 1991; Van de Vijver et al. 1999) and aboveground net primary productivity (Buis et al. 2009).

Savannas are considered to be highly variable and diverse ecosystems (House et al. 2003) and as a result fire plays differing roles of importance in these systems. In arid savannas, climatic variables often set limits to the amount of canopy cover because there is insufficient moisture to form a closed canopy system. However, where rainfall is sufficient for this to happen, fire and other disturbances such as herbivory are thought to limit woody cover and thus maintain a savanna structure of trees and grass (Bond & Keeley 2005; Sankaran et al. 2005). Effects of fire on other ecosystem processes and potentially fauna are therefore expected to differ somewhat in different savanna types.

Knowledge on faunal responses to fire is, however, limited. This was highlighted in a review by Parr and Chown (2003) who critiqued the literature on fire and fauna studies in southern Africa and found it to be limited. Certain groups of animals, notably reptiles, amphibians and invertebrates were strikingly under represented in fire studies. In addition to this, the authors suggested that most of the current studies are incomparable due to vastly different experimental approaches and poor reporting. Furthermore, many of the studies were observational reports and few were based on long-term burning
experiments, focusing rather on once off fire events which are quite different to fire regimes. If ecologists and managers of conservation areas are to better understand the role of fire, more studies, based on sound experimental design, need to be conducted, especially in relation to long-term burning regimes (Parr & Chown 2003; Chapter 2).

This is of much relevance in relation to invertebrates. Although a fair amount of work has investigated insect responses to fire (see Swengel (2001) for a review), little of this is specifically related to savannas globally or to the southern Africa region (Parr & Chown 2003). Yet insects form the bulk of species diversity and biomass on earth and are critically important for ecosystem functioning around the globe (Wilson 1991) and therefore cannot be ignored when seeking to understand ecological processes or planning conservation objectives.

One such insect group is termites (Blattodea: Termitidae), which are of great ecological importance in the tropics and sub-tropics. They are considered to be ecosystem engineers in that they modify the mineral and organic composition of soils, as well as affecting soil drainage (Jones et al. 1994; Konaté et al. 1999). They also improve soil infiltration rates and porosity, while decreasing bulk density (Mando et al. 1996). In tropical forests and savannas, termites are considered to be the dominant arthropod decomposer (Collins 1981; Holt 1987; Schuurman 2005) and in African savannas have been shown to have a biomass comparable to ungulates and mega herbivores (Dangerfield et al. 1998). As decomposers, their role in nutrient cycling and distribution can also not be ignored (Holt & Coventry 1990; Konaté et al. 1999; Lepage et al. 1993; Scholes 1990).

Termitaria constructed by termites have also been shown to affect the ecological dynamics of savannas in a variety of ways (see e.g. Fleming and Loveridge (2003) on how *Macrotermes* mounds provide habitat for small mammals). Moe et al. (2009) found a significantly higher number of woody plant and forb species occurring on mounds compared to the surrounding matrix, which suggests that mounds can increase the patchiness and heterogeneity of savanna vegetation by forming nutrient hotspots. By concentrating nutrients, termitaria also provide key resource areas including quality forage that is intensely used by browsers and grazers (Grant & Scholes 2006; Mobæk et
The presence of this forage may enable the persistence of herbivore species within a landscape that would otherwise offer lower forage quality.

Despite their importance in these ecosystems, knowledge on termites is limited, with even basic natural history information lacking for many species (Dangerfield & Schuurman 2000; Dawes-Gromadzki 2003). Taxonomically, many genera are also in need of revision (Uys 2002), and many new species are surely awaiting discovery and description. Standardised sampling protocols, especially for savannas, are also lacking (Dawes-Gromadzki 2003) which contributes to the problem. In terms of the effects of fire on termites, much the same scenario arises, with little data or comparable studies (see Chapter 2 for a review and suggested future research directions). But if we are to better understand savannas and effectively manage conservation and agricultural land within them, more knowledge on interactions between fire (an often used management tool) and termites (a critically important organism) is necessary.

Objectives

The objective of this study was to enhance the scientific knowledge and understanding of termite-fire interactions in southern African savannas. To this end sampling and analysis of termite communities subjected to different long-term burning regimes was conducted using sites in two South African reserves, Kruger National Park (22° 25' - 25° 32' S and 30° 50' - 32° 02' E (hereafter KNP) and Hluhluwe-iMfolozi Park (28° 01' - 28° 25' S and 32° 15' - 32° 26' E (hereafter HiP)). Since effects of fire on vegetation, and likely fauna, vary in savannas of differing climatic and other abiotic variables (Sankarran et al. 2005, Bond and Keeley 2005), these interactions were tested in four different savanna types situated along a rainfall gradient. In addition, several methods and techniques were used to adequately sample the termite communities, allowing an additional investigation of sampling method efficiency to be conducted, with the aim that this will enhance future studies related to termite ecology and diversity.

Study area

The KNP is the largest protected area in South Africa, covering an area of 1 948 528 ha (van Wilgen et al. 2007) while HiP is considerably smaller at approximately 90 000 ha.
I. Introduction

(Archibald et al. 2005), but still one of South Africa’s largest parks. Both parks are located in the eastern part of the country and have a sub-tropical climate with a distinct summer rainfall pattern. Although highly variable throughout the park, the existing fire regime in KNP is generally characterised by a concentration of fires during the late dry season (late winter), with a mean fire return interval of 4.5 years (van Wilgen et al. 2000). At HiP, fire regimes have been highly variable with a shorter mean fire return interval of 3.8 years, although this is highly skewed toward shorter return intervals, the median being only 1.3 years (Archibald et al. 2005).

Between 1948 and 1956, the fire management policy in the KNP was aimed at reducing the impacts of fire (van Wilgen et al. 2007), but in 1953 an uncontrolled fire burnt approximately a quarter of the park (Biggs et al. 2003). This raised concerns over the impacts of fire on the flora and fauna of the park and in response to this a series of experimental burn plots was established in 1954 spanning much of KNP’s latitudinal gradient (Fig. 1). This experimental burn plot trial therefore provides an ideal platform to examine questions related to long-term burning regimes and rainfall seeing that it has been in place for several decades and is situated along a rainfall gradient of approximately 450 mm to 700 mm. Consequently, it has become an invaluable savanna burning experiment, and although initially established to test effects of fire on vegetation, the experiment has since been used to study many other aspects of savanna ecology and fire interactions including faunal responses (see van Wilgen et al (2007) for a review of the experiment).

The KNP experiment was set up in four of the major vegetation types in the park, within the Mopani, Satara, Skukuza and Pretoriuskop areas (Fig 1), and within each of these vegetation types it was replicated four times in the form of strings consisting of 12 – 14 plots, each of which are subjected to a different burning regime. These burning regimes consist of different season (dry, wet and transitional) and frequency (from annual through to sexennial) combinations and importantly, each string has a control plot which is prevented from burning (see Fig. 2 for more information on the different burning regimes used in this study). Each plot measures approximately 380 x 180 m (7 ha) and is protected from each adjacent plot by a single fire break. Each string (replicate) is in turn protected from the surrounding vegetation by a double fire break road (Biggs et al. 2003).
For this study, I made use of only three of the four KNP vegetation types available because the fourth set of plots (the Skukuza plots) are located across differing soil types (Venter 1999) which could possibly have led to confounding results. Moreover, I made use of only three of the four available strings of plots (replicates) in each vegetation type seeing that the fourth replicate is on a different soil type to the other three (Venter 1999) and therefore considered to be unrepresentative and was not utilised. Thus three replicates, situated 10 – 20 km apart, were chosen in three vegetation types.

In order to extend the rainfall gradient, I also sampled termites at HiP, where rainfall is approximately 900 mm. However, no such burning experiment exists here and so fire records were utilised in order to match up sites with similar fire histories to those of the burn plots in KNP. Even though there is not such a neat experiment in HiP, it was considered useful to add to the study because the higher rainfall enabled me to assess termite communities in a more mesic savanna. Here, three sites of each burning regime, as described in Figure 2, were selected to match the replication used in KNP, with the exception of the burning regime used to match the triennial August burns as an adequate third site was not found. Although these sites were initially chosen in order to test effects of different fire regimes on termites, it was decided to only use the HiP sites in combination with the KNP sites to test for differences in termite communities across savanna types (i.e., the overall rainfall gradient component of the study), due to uncertainty in the HiP fire histories and thus uncertain conclusions been drawn.

Within these two parks, the study was therefore carried out in four distinct savanna vegetation types, Mopane woodland (Mopani area of the northern KNP), Acacia savanna (Satara area of the central KNP), Terminalia woodland (Pretoriuskop area of the southern KNP) and mesic Acacia savanna (Hluhluwe section of HiP), arranged along this rainfall gradient. The three replicates used in the Mopani section of KNP were Tsendse (23°41'S 31°31'E), Mooiplaas (23°34'S 31°27'E) and Dzombo (23°26'S 31°22'E); the replicates used in the Satara region were N’wanetsi (24°26'S 31°51'E), Marheya (24°32'S 31°46'E) and Satara (24°24'S 31°45'E); and the replicates in the Pretoriuskop area were Kambeni (25°15'S 31°26'E), Numbi (25°13'S 31°20'E) and Shabeni (25°12'S 31°23'E). In HiP, all sites were located in the northern Hluhluwe section of the reserve. The following four burning regimes were selected in each of the vegetation types: August
annual, August triennial, December triennial and the unburnt control plot (Fig. 2). These
were, as best as possible, matched in HiP with sites of similar fire history. August burns
represent late winter fires which are at the end of the dry season and are intense, hot fires
while December burns represent early summer fires at the beginning of the wet season
and are cooler, less intense fires. The use of these burning regimes enabled me to test the
effects of fires of differing season (late dry and early wet), frequency (annual and
triennial) and intensity (hot dry season fires and cool wet season fires) (Fig. 2).

The Mopani plots were situated in the Mopane shrubveld, which is a mopane
(\textit{Colophospermum mopane} J. Kirk ex J. Léonard) dominated habitat with few other
woody species present (Low & Rebelo 1996). The mean annual rainfall is 450 to 500 mm
and the area ranges in altitude from 300 to 340 m a.s.l. (Gertenbach 1983). In the Satara
region, the plots were situated in a mixed knobthorn \textit{Acacia nigrescens} Oliv. and marula
\textit{Sclerocarya birrea} (A. Rich. Hochst.) savanna in the sweet lowveld bushveld (Low &
Rebelo 1996). The area ranges in altitude from 240 to 320 m a.s.l. and has a mean annual
rainfall of 550 mm (Gertenbach 1983). The Pretoriuskop region is sour lowveld bushveld,
consisting of open tree savanna dominated by silver clusterleaf (\textit{Terminalia sericea}
Burch. ex DC), but bushwillow (\textit{Combretum collinum} Fresen.) is also common (Low &
Rebelo 1996). The region has a mean annual rainfall of 700 mm and the altitude varies
from 560 to 640 m a.s.l. (Gertenbach 1983). The sites situated in HiP were located within
Natal lowveld bushveld, dominated by various acacia species, notably \textit{A. tortilils} and \textit{A.
karoo}, but several other species are also common (Low & Rebelo 1996). The mean
annual rainfall for the entire park is 600 to 1000 mm and the altitude ranges from 40 to
750 m a.s.l. (Archibald \textit{et al.} 2005), with the Hluhluwe region being in the wetter and
higher altitude range of the park. Both the Mopani and Satara regions have basalt-derived
clay soils while the Pretoriuskop region has granite-derived sandy soils (Gertenbach
1983). At Hluhluwe the underlying geology is predominantly sandstone and shale
resulting in sandy soils (Archibald \textit{et al.} 2005).

\textbf{Structure of the thesis}

Following this introduction, Chapter 2 provides a review of current knowledge on
termite-fire interactions. I show that currently general conclusions cannot be drawn and
provide suggestions for future research. Building on this, Chapter 3 examines the effects of long-term burning regimes on termite diversity and activity. Chapter 4 makes use of some of the sampling methods used in Chapter 3 to assess the efficiency of sampling methods and how this may change across savanna types situated along the rainfall gradient. The thesis ends with a general conclusion of the overall thesis findings in Chapter 5, with reference to management and conservation implications.

References
1. Introduction


I. Introduction


I. Introduction


Figure 1: Location of the replicates of the experimental burning plots in the Kruger National Park, South Africa. Hatched and striped areas indicate the four major vegetation types the experiment is located within while the unshaded area represents granitic based geology while the shaded area shows the basalt areas. Taken with permission from van Wilgen et al. (2007)
Figure 2: A schematic diagram of the experimental design that was followed in each savanna type. Four burning regimes, replicated three times, were sampled in each savanna type in order to examine a) different fire seasons (late winter and early summer), b) fire frequencies (annual vs. triennial), and c) burnt plots vs. long unburnt plots. Fire intensity was also assessed along a gradient of very hot intense fires at triennially burnt August plots to cool low intensity fires at triennially burnt December fires.
Chapter 2

Termites and fire: current understanding and future research directions for improved savanna conservation

Abstract

Termites are considered to be major ecosystem engineers in tropical environments. However, they are often neglected and under studied, especially in grassy systems. The interaction between termites and fire ecology is one example of this. Given the importance of both fires and termites in savanna systems, it is critical for an improved ecological understanding of savanna biodiversity that the interaction between them is better understood. In this paper, we highlight the lack of published literature on fire ecology and invertebrates in austral systems, with particular focus on termites. We review the six studies conducted in savannas spanning the southern hemisphere which investigate this interaction, summarizing their findings and limitations. We suggest areas of future research related to fire and termites which will improve our ecological understanding of savannas and the dynamics that structure these systems.

Keywords: burning regimes, invertebrates, research directions, savanna, termites.
 Importance of fire and invertebrates

While tropical savannas and grasslands typically attract less attention than temperate systems (see e.g. Gaston et al. 1998 and references therein indicating the general lack of biological data at low latitudes), they represent one of the dominant biomes of the southern hemisphere, and indeed the world, with savannas covering an estimated 12.5% of global land area and over half of Africa and Australia, 45% of South America and 10% of India and Southeast Asia (Scholes & Archer 1997). Fire is considered one of the most important disturbances in these grassy systems being both frequent and widespread across Australian, African and South American savannas (Bond & Keeley 2005). The exceptional diversity and biomass of invertebrates has been long recognized, as has their functional importance in ecosystem services across the globe (Wilson 1987). Fire and invertebrates are thus key components determining the functioning and dynamics of savannas (Parr et al. 2004, van Wilgen et al. 2007), as well as other systems around the world (e.g. boreal forests – McCullough et al. 1998).

Yet, generally scientists, and more especially conservation managers, have a poor understanding and limited predictive capacity of the way in which invertebrate communities respond to fire and the implications for diversity and functioning in savanna systems. This was highlighted for the southern hemisphere by Parr & Chown (2003) who reviewed fire and invertebrate studies in southern Africa. A search on Web of Science using the keywords ‘insect*’ and ‘Africa’ with either ‘fire*’ or ‘burning’ revealed that since the review by Parr & Chown (2003) only three additional papers and one book chapter dealing with fire and invertebrates in Africa have been published (Parr et al. 2004; Axmacher et al. 2006; Uys et al. 2006; Underwood & Christian 2009); this indicates little has changed since 2003. Similarly and for the same time period, work conducted at the global scale is also limited in its contribution to our broader understanding of the ecological processes involved when studying fire and invertebrate interactions. The lack of knowledge on invertebrates and fire is thus an important shortcoming to understanding how savanna systems are structured and function, and also when managing for the conservation of biodiversity, especially since savannas are often subject to intense fire management (van Wilgen et al. 2007).
In tropical and sub-tropical areas, including savannas, termites are considered key ecosystem engineers, altering the mineral and organic composition of soils, their hydrology, drainage (Jones et al. 1994) and infiltration rates (Mando et al. 1996), as well as influencing decomposition, nutrient cycling and distribution (Holt & Coventry 1990; Scholes 1990; Lepage et al. 1993; Konaté et al. 1999). Their vast biomass alone makes them an important consideration in tropical and sub-tropical ecology (Josens 1983) and they are considered the dominant arthropod decomposer in tropical forests and savannas (Collins 1981; Holt 1987; Schuurman 2005). Termites are also a diverse and varied group comprising of several functional groups, including wood feeders, grass harvesters and soil feeders (Josens 1983). Although it is somewhat surprising that relative to other invertebrate groups termite work is scarce with even basic natural history information lacking for many species (Dangerfield & Schuurman 2000; Dawes-Gromadzki 2003), this is likely because there are sampling and taxonomic difficulties associated with the group (Josens 1983).

Here I use examples of studies conducted in savannas across the globe to highlight what I see as a critical research gap: the lack of published information on the interactions between termites specifically (Blattaria: Termitidoidae) and fire. Furthermore, I propose that if protected areas in savannas are to uphold their mandate of conserving biodiversity in its entirety, then a better ecological understanding of interactions between fire and termites is crucial for achieving this. The aims of this paper are therefore to highlight what I perceive as a paucity of studies on the topic, to demonstrate that critical information is still lacking, and to look towards potential future research directions. I compare studies from several continents and suggest ways in which the termite fauna of these continents could vary in their response to fire and some resultant ecological implications of this, particularly for the southern hemisphere.

Current knowledge status
Since fires in savannas are frequent, and that in these systems they represent a disturbance that is often controlled (e.g. determining season, fire size and frequency, Govender et al. 2006), it is important to have basic information on how burning influences diversity and processes within savannas. Basic research questions that will
advancing ecological understanding of termite-fire interactions include: what is the effect of different fire regimes (e.g. fires of differing frequency, season and intensity) on termite diversity and activity? How do the effects of repeated fires vary from single fires? And, how do different functional groups of termites respond to variation in burning regimes, e.g. are grass harvesters, for example, more susceptible than other groups?

Following on from this and given the importance of fire and the role of termites in savannas it is concerning that to our knowledge, only six studies focusing on fire-termite interactions in savannas have been published to-date: Trapnell et al. (1976), Benzie (1986), Ferrar (1982), DeSouza et al. (2003), Dawes-Gromadzki (2007) and Traore & Lepage (2008). These constitute four from Africa, and one each from South America and Australia. Even when other biomes are included, there has only been one additional study conducted on fire and termites in the southern hemisphere, this in the Mediterranean-type vegetation in Western Australia (Abensperg-Traun & Milewski 1995). Although I acknowledge the scientific value of these studies, an understanding of the effects of fire on termites is, to-date, clearly limited in scope and geographic coverage, with large areas such as east Africa and India being poorly studied. Single studies in Australia, South America and southern Africa also cannot be seen as a true representation of fire-termite interactions on these continents due to the high levels of variability within and across savanna landscapes and at larger scales (e.g. continental and sub-continental scales).

Of the six studies conducted in savanna ecosystems, three dealt with fire regimes (Trapnell et al. 1976; Benzie 1986; Traore & Lepage 2008) and three with single fire events (Ferrar 1982; DeSouza et al. 2003; Dawes-Gromadzki 2007). The three studies dealing with fire regimes (all conducted in Africa) considered only two extreme regimes: that of fire exclusion and of annual burns. In all three, only mound building termites were considered and the experiments in the Trapnell et al. (1976) and Benzie (1986) studies were not replicated. Results were complementary for the two studies carried out in west Africa (Benzie 1986; Traore & Lepage 2008); in both these studies Trinervitermes mound densities decreased significantly under fire treatment, and Macrotermes and Cubitermes mounds remained unaffected by fire. However, in the study by Trapnell et al. (1976), mound densities of Cubitermes termites increased under fire treatment, with no results for other genera reported. The latter study was conducted on plots which had been
subjected to experimental burning for much longer (23 years) than the Benzie (1986) and Traore & Lepage (2008) studies (3 and 10 years respectively). Difference in habitat (Zambian miombo woodland and west African guinea and Sudanian woodland), and likely also species, may explain some differences in the results for *Cubitermes* mound density changes, but no clear patterns can be detected with such limited data.

Of the three studies which investigated effects of a single fire event on termites, one was conducted in Africa, one in Australia and the other in South America. The Australian study (Dawes-Gromadzki 2007) was replicated and all functional groups were considered along with other soil macroinvertebrate fauna. Termite abundance was found to decrease significantly after fire was applied to the experimental sites (this despite the relatively low intensity of the fires), but effects on species richness were less apparent. In South America (DeSouza et al. 2003), fire had no effect on the termite assemblages but the relatively coarse level of resolution may have masked any species-level effects and experimental replication was poor. The African study was conducted in a savanna area of South Africa (Ferrar 1982), and was an opportunistic study after a low intensity fire burnt a study area being used to look at more general termite ecology. Cellulose baits consisting of toilet rolls were laid out on the site post fire to sample termites present over time as the vegetation recovered from the fire. The results of the study show that termite attack of the baits was much reduced compared with similar sites which had not been burnt. Ferrar (1982) concluded that this was probably the result of the protective grass cover being removed by the fire, leading to greater insolation and desiccation of the soil, making it unfavourable for termites, however, no pre-fire data were available. The study documents a slightly delayed increase in termite activity post-fire, although this post-fire time interval is not given.

The above studies suggest that fire can have a significant effect on termite densities and activity levels, especially it seems when the extremes of fire treatment are examined, that of fire exclusion and annual burning. However, even though some invertebrate groups have shown resilience to burning and others have not (see Swengel 2001 for a review on insect responses to fire), I propose that there are insufficient studies to provide a clear picture of the relationship between fire ecology and termites, particularly because only extreme fire regimes have been studied. Fire exclusion and
continuous annual burning are both unrealistic and arguably unnatural management options for savannas, and since these are the only two regimes investigated, the above studies are somewhat limited in their contribution to our understanding of fire ecology. Furthermore, the four African studies focused only on one suite of termites – mound builders in west and central Africa (Trapnell et al. 1976; Benzie 1986; Traore & Lepage 2008) and wood feeders attracted to cellulose baits in southern Africa (Ferrar 1982). Experimental replication was also limited in three of these four studies, with suitable replication only occurring in the Traore & Lepage (2008) study. The South American study did take more functional groups into consideration, but here replication was again poor and the study had a coarse taxonomic resolution. The Australian study avoided both these problems, but only gives us a snapshot of the ecological process of termite recovery after fire since it was based on a single fire event and termites were only sampled once after the fire (after two weeks). Finally, the range of responses detected from these studies means that general patterns at this stage are impossible to detect.

Future research directions

In order to improve current knowledge on fire-termite interactions, and consequently savanna ecology, I highlight the urgent need for further research and recommend that it should especially be focused on long term experiments. Research on long term burning regimes in addition to single-fire events is important because the effects of single-fire events can be markedly different to those from repeated burning (Parr & Chown 2003); the effects of fire regimes on biodiversity need to be understood in order to implement ecologically relevant fire management strategies. In addition, a wider range of fire regimes spanning different intensities (e.g., seasons) and frequencies need to be investigated rather than only fire exclusion and annual burns which are not realistic or practical to implement. At the same time, it is also important to better understand the ecological impacts of single fire events since accidental or anthropogenic unnatural fires occur frequently in savannas across the globe. To this end, studies of single fire events will be valuable especially where accompanied with pre-fire data.

Another aspect of termite ecology which warrants urgent attention is that of their diversity. Because the studies to date focus on a limited suite of functional groups the
ecological effects of fires at the assemblage level are unclear. I predict that the various functional groups will respond differently to repeated burning events. For example, wood feeding termites may be more likely to be affected by intense fires or repeated burning which consume large amounts of dead wood. Although no data are available for African savannas, in Australian savannas, on average 27% of standing dead wood is consumed per low intensity fire (Fensham 2005); this figure is likely to be significantly higher with more intense fires. In contrast, less severe fires which only remove the grass layer may be more likely to affect grass harvesters. This may potentially result in a shift in functional groups under different fire regimes which may have a significant influence on system functioning.

Interactions between termites and fire on different continents are expected to differ somewhat. In African savannas, for example, these interactions are expected to differ from those elsewhere due to the high diversity of ungulates and the resultant effects they have on ecosystem functioning. These effects include maintaining a balance between grass and woody biomass and increasing the rate of nutrient cycling (Du Toit & Cumming 1999), a process also critical to and facilitated by termites and their ecology. The large degree of savanna spatial heterogeneity in Africa which is driven by variations in rainfall, topography and soil fertility (Du Toit & Cumming 1999) could provide many opportunities to study termite fire interactions in differing habitats. These studies would also be useful to understanding savanna ecology elsewhere, since many processes operating in African savannas are common to savannas elsewhere and this could lead to worthwhile comparisons being drawn between continents; similarities include high seasonality and the importance of fire in these systems (Bourlière & Hadley 1983).

Since termites originated in African rainforests (Aanen et al. 2002), this may also lead to differences in response to fire between Africa and other continents due to consequent higher termite diversity. Evolutionary history could also be used to explain findings on other continents because traits carried over to these continents could be ancestral and evolved under African conditions. Similarly, the biogeographic origin of different species could possibly be used to predict responses of species to fire. For example, there may be a range of savanna-adapted genera that are less affected by fire compared to genera that have a forest-oriented biogeographic history such as
Ancistrotermes and indeed most of the Apicotermitinae subfamily (Eggleton et al. 2002). These forest-associated genera may be more susceptible to frequent fires that open up woody vegetation and alter microclimatic conditions and food availability. Climatic variables may also be of importance in predicting termite responses since more arid areas have been shown to be more resilient to fires than mesic areas with regards to other invertebrates (see e.g., Parr et al. 2004 for ants). Much work needs to be conducted before these hypotheses, related specifically to termites, can be supported or falsified.

Another avenue of future research could focus on how termites might, in turn, influence fire patterns in savannas. Termites are herbivores which often forage sporadically in a patchy pattern dependent on environmental conditions (Wood 1978). This could have important implications for fire spread in savannas; for example, areas where harvester termites have removed grass provide short-grass patches which could prevent the spread of fire and create unburnt refuges for other taxa to utilize. Understanding of this relationship could provide answers to how termites as herbivores interact with fires to shape savannas.

I hope that this paper will inspire future work on termites and fire ecology, despite the difficulties associated with it. Knowledge at a species, as opposed to generic, level would be especially helpful in providing a more complete picture of responses and ecological implications. Although this is not an easy task given the state of termite taxonomy, understanding how fire ecology and termites interact is of critical importance for improved ecological understanding and furthermore for ensuring effective management and conservation of savanna ecosystems.

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Chapter 3

Savanna termite responses to long-term burning regimes

For submission to *Ecology* as: Andrew B. Davies, Paul Eggleton, Catherine L. Parr and Berndt J. van Rensburg
Abstract

Fire in savanna systems is regarded as a major and necessary disturbance for the maintenance of biodiversity. However, most fire ecology studies have focused on vegetation dynamics with little attention given to other taxa, especially invertebrates. Termites are considered to be major eco-system engineers in savannas, but are often neglected due to sampling and taxonomic difficulties. Hence very little is known on the interactions between termites and ecological processes such as fire. But if we are to conserve biodiversity, interactions between ecological processes and drivers need to be well understood. I made use of a long-term burning experiment initiated in 1954 in the Kruger National Park, as well as sites in Hluhluwe-iMfolozi Park, South Africa, to test the effect of different fire regimes on termite assemblages. I sampled across four distinct savanna types situated along a rainfall gradient, and across three seasons. Termites were comprehensively sampled using a variety of methods in order to sample a large suite of functional groups. Assemblages differed significantly across savanna types with higher levels of diversity in mesic sites. Diversity was lowest at the most arid site but certain feeding groups peaked at Satara, a savanna with intermediate rainfall. Differences between these savannas are attributed to broad underlying changes in net primary productivity, temperature and soil type, with the role of mammalian herbivores also being considered. Seasonality was also examined and it was found that termite activity peaks in the wet and transitional seasons and is significantly lower in the dry season. Termites were overall highly resistant to fire, but assemblage composition was affected, this being more pronounced with increasing annual rainfall. Assemblage changes are linked to changes in vegetation structure caused by fire. Vegetation is little affected by fire in drier areas, but in wetter areas fire plays a major role in shaping its structure and composition, this same pattern seems to be reflected with termites. Although managers can afford to place more emphasis on other taxa when deciding on a burning policy, in mesic savannas more caution should be exercised.

Keywords: assemblage composition, conservation, fire ecology, invertebrates, resistance
3. Termite responses and fire

Introduction

Fire is a frequent and widespread disturbance in African savannas, where it acts like a generalist herbivore modifying vegetation structure and composition (Bond and Keeley 2005) and facilitates the co-existence of trees and grass (Higgins et al. 2000, van Langevelde et al. 2003). Fire has also been shown to influence grazing patterns (Archibald and Bond 2004, Archibald et al. 2005, Parrini and Owen-Smith 2009), nutrient cycling (Hobbs et al. 1991, Van de Vijver et al. 1999) and aboveground net primary productivity (Buis et al. 2009).

Although the effects of fire on savanna vegetation are well documented (see e.g. Bond 1997, van Wilgen et al. 2003, van Wilgen et al. 2007), for animals, and especially invertebrates, knowledge is scarce (Parr and Chown 2003, Parr et al. 2004, Chapter 2). This is concerning since fire is used extensively in savanna systems across the globe as a management tool for livestock farming, and often to achieve conservation goals in protected areas (Biggs et al. 2003, van Wilgen et al. 2004, Andersen et al. 2005). Few studies have been conducted on invertebrates, and results thus far have been mixed, with some studies concluded that fire does have some affect on invertebrates (Ferrar 1982, Zieger et al. 1998) while others (Parr et al. 2002, Parr et al. 2004) found fire to have little affect. Furthermore where studies have been carried out, they have focused predominately on single fire events or burning regimes which do not reflect natural regimes. Experiments which make use of long-term burning regimes that more closely resemble natural fires are thus likely to provide more meaningful ecological understanding and allow more informed management decision-making (Parr and Chown 2003).

Sankaran et al. (2005) introduced the idea that African savannas tend to be either stable or unstable depending on the mean annual precipitation (MAP) received. That is, below a threshold MAP of ~650mm savannas are considered to be more stable, with disturbances such as fire and herbivory having little effect on woody vegetation cover. However, savannas with a MAP greater than ~650mm are considered to be unstable because they have enough moisture to potentially form closed canopy forests. In these areas, disturbances such as fire and herbivory are crucial for maintaining an open canopy that allows grasses to grow, and thus these disturbance types play critical roles in
determining woody vegetation structure and composition. Whether the effect of these disturbances can also significantly influence the fauna remains to be formally examined. However, if faunal response to fire is linked to changes in vegetation structure, as has been suggested (Catling et al. 2001, Parr et al. 2004), I would expect limited or no response in arid, more stable savannas and a significantly greater response in more mesic savannas (>650 mm). Results from a study by Parr et al. (2004) looking at fire effects on ant assemblages across savanna systems suggested this may indeed be the case: in low rainfall (‘stable’) savannas there was little or no change in ant richness, abundance and composition with burning, whereas in higher rainfall (‘unstable’) savannas responses were greater. I aimed to test this idea more fully by determining how termite communities responded to fire at a range of savanna sites along a rainfall gradient. I investigated long term burning regimes across this rainfall gradient which encompassed both ‘stable’ and ‘unstable’ African savannas. Effects of fire were thus investigated at, and compared between, savannas of markedly different rainfall patterns.

Termites (Blattodea: Termitidoidae) are an invertebrate group which has been largely neglected in terms of fire research (see Chapter 2), even though they are considered to be ecosystem engineers and crucial to savanna system functioning (Jones et al. 1994). They are capable of altering soil composition and hydrology, as well as drainage (Jones et al. 1994) and infiltration rates (Mando et al. 1996). Termites are also considered to be the dominant invertebrate decomposer in tropical forests and savannas (Collins 1981, Holt 1987, Schuurman 2005), and thus also play an important role in nutrient cycling and distribution (Holt and Coventry 1990, Scholes 1990, Lepage et al. 1993, Konate et al. 1999). Although termites are notoriously difficult to sample comprehensively, especially in savannas, understanding how they are affected by fire in savanna systems is critical for a more effective understanding, and conservation, of those systems.

The aims of this study were to investigate how termite diversity (richness, abundance and composition) and activity (i) vary between major savanna habitats situated along a gradient of increasing MAP; (ii) vary with season, and (iii) are influenced by long-term burning regimes. I will test whether termite responses to fire can be fitted within the stable-unstable savanna conceptual framework. This is the most
comprehensive study to date on savanna termites not only in terms of the spatial and temporal extent of the study but also in the diversity of sampling methods employed.

**Methods**

**Study site**

The study was conducted in four distinct savanna vegetation types along a rainfall gradient in two of South Africa’s major reserves: Kruger National Park (KNP, 22° 25' - 25° 32' S, 30° 50' - 32° 02' E) and Hluhluwe-iMfolozi Park (HiP, 28° 01' - 28° 25' S, 32° 15' - 32° 26' E). These parks are located in the eastern part of the country and experience a sub-tropical climate with a distinct summer rainfall pattern. The existing fire regime in KNP is generally characterised by a concentration of fires during the late dry season (late winter), with a mean fire return interval of 4.5 years, although this does vary substantially throughout the park (van Wilgen et al. 2000). In HiP, fire regimes have been highly variable with a shorter mean fire return interval of 3.8 years, although this is highly skewed toward shorter return intervals, the median being only 1.3 years (Archibald et al. 2005) (for more details on these parks see Archibald et al. 2005 and van Wilgen et al. 2007).

In KNP termite sampling was carried out in three dominant savanna vegetation types in the park using a large-scale (spanning ca. 250 km), long-term burning experiment established in 1954 (Biggs et al. 2003, van Wilgen et al. 2007). Within each savanna type, sampling took place on a series of burn plots (7 ha each in size), replicated three times. Replicates within a savanna type are situated 10 – 20 km apart. The three savanna types consisted of Mopani woodland (Mopani area of the northern KNP), Acacia savanna (Satara area of the central KNP) and Terminalia woodland (Pretoriuskop area of the southern KNP). In order to test the effects of fire season (late dry and early wet), frequency (annual and triennial) and intensity (hot, dry season fires and cool, wet season fires), four burning regimes were selected within each replicated plot namely August annual, August triennial, December triennial and the unburnt control plot. August burns represent late winter fires which are at the end of the dry season and are intense, hot fires while December burns represent early summer fires at the beginning of the wet season and are cooler, less intense fires.
Because no such experiment exists in HiP, fire map records were used to identify sites in the landscape with fire histories that, as best as possible, match those of the burn plots in KNP. Despite using only fire histories, this savanna site was investigated because it is in a higher rainfall area and thus provides a more mesic savanna for comparison. Three sites of each burning regime were selected in mesic *Acacia* savanna (Hluhluwe section of HiP) to match the replication used in KNP, with the exception of the burning regime used to match the triennial August burns where only two sites were selected at HiP as an adequate third site was not found. Because there is less certainty about the fire histories of the selected HiP plots, they were later excluded from any fire ecology analyses, and instead only used for overall savanna comparisons. The total rainfall gradient represented by the four savanna types (i.e. KNP and HiP combined) ranged from 450 mm (Mopani woodland in KNP) to 900 mm (mesic *Acacia* savanna in HiP) (see Supplementary material S1 for more information on the replicated sites in each of the four habitat types).

**Termite sampling**
Termites were sampled using a variety of methods in order to sample all feeding groups adequately: cellulose baits, active searching, and mound counts. Sampling was conducted in three seasons during 2008 and early 2009: March to May 2008 (the austral autumn, transitional season), June to September 2008 representing the dry season (the austral winter), and November 2008 to February 2009 representing the wet season (the austral summer).

During each season, 40 cellulose baits in the form of toilet paper rolls (500-sheet single-ply, unscented - hereafter referred to as baits) were placed on each plot (see Le Fage et al. 1973). They were arranged in a grid of five by eight baits, spaced five metres apart at the centre of each plot to prevent edge effects. Half (20) of the baits were placed directly on the soil surface, any litter present was brushed aside to ensure direct contact was made with the soil and the base of the bait (Dawes-Gromadzki 2003). These baits were wrapped around the middle with packaging tape to prevent unravelling and secured to the soil surface using an aluminium tent peg placed through the centre gap of each bait. The other half of the baits were buried below the soil surface at a depth of approximately
2 cm which ensured they were adequately covered with soil. Each line of eight baits consisted of four buried and four surface baits with the position of the baits alternated. Baits were checked for termites, and termite attack assessed, after predetermined intervals of 7 (5 days during the wet season), 14, 28 and 56 days. At each of these time intervals, five surface and five buried baits were removed, and any termites present were sampled (at least ten of each caste were sampled when present to ensure a representative sample, Uys 2002). The baits were then scored according to the intensity of attack (IA). This involved giving each bait a score of one to six according to how much of the bait had been consumed, which was assessed visually. Baits which were completely intact with no evidence of termite activity were assigned a value of 0, subsequent to this baits were scored according to the following scale: 1 = 1 – 25% consumed, 2 = 26 – 50% consumed, 3 = 51 – 75% consumed, 4 = 76 – 99% consumed and 5 = 99 – 100% consumed (Dawes-Gromadzki 2003). Baits which had been disturbed by vertebrates or otherwise affected (e.g. no longer in direct contact with the soil surface or completely destroyed) were excluded from analysis.

During the transitional and dry seasons, two belt transects of 50 by 2 metres were walked by two people in each plot, during which any termite activity was noted and termite searching was conducted. In addition, any course woody debris (CWD) found along the transects was sampled and later weighed and searched thoroughly for termites. During the wet season, representing peak termite activity (Dawes-Gromadzki and Spain 2003), this method was augmented to be a modified version of the standardised belt transect method described by Jones & Eggleton (2000) (see Supplementary material S2 for more information on this modified method). Two 50 by 2 metre belt transects were laid out on each plot, with each sub-divided into ten 5 x 2 metre blocks and each block searched for a period of ten person minutes. During this time period the entire block was searched for termites; this included excavating soil and searching dead wood, termite nests and other microhabitats. Vegetation up to a height of one metre was searched.

Mound counts were conducted during the dry season of 2008. Two blocks of 50 by 50 metres were laid out on each plot and searched thoroughly by three people for any mounds present. These mounds were then broken open to sample any termites found. The
3. Termite responses and fire

dry season was selected for this because grass cover at this time is less dense, allowing more effective searching.

Termite Identifications
Termite samples were identified at the Natural History Museum, London, U.K., the University of Pretoria and the National Collection of Insects, Pretoria, South Africa. Specimens were identified to species were possible using soldier castes whenever available (when soldiers were not present, workers were used). Where this was not possible, genera were separated into morphospecies (e.g. for Odontotermes), and where even this was problematic the specimens were pooled to genus (this occurred, for example, with the genera Microtermes, Microcerotemes and Astalotermes). Many termite genera are in need of taxonomic revision (Uys 2002) and it was considered better to treat these taxa at a genera level only instead of risking the possibility of finding differences in community assemblages when species thought to be different were actually the same. Although I refer to ‘species’ throughout the manuscript, due to the above taxonomic constraints, this refers to genera in cases where a genus could not be easily split into its constituent species. A reference collection will be lodged with the National Collection of Insects, Pretoria.

Environmental data
A variety of methods were used to quantify and describe the vegetation present on each plot. The percentage cover of each vegetation type was assessed using twenty 1-m² quadrats placed ten metres apart in a 5 x 4 grid. The percentage cover of grass, bare ground, dead grass and leaf litter, forbs, woody vegetation, dead wood, dung and rock was estimated for each quadrat and the percentage of canopy cover was also estimated. Data on the dominant grass and tree species for each plot were provided by South African National Parks (KNP) and the Zululand Tree Project (HiP).

Temperature readings were recorded using iButtons at selected sites which provided a representation of the various burning regimes. A single iButton was placed on each site for the most centrally located string of plots in each savanna type during each season and also on two sites for each adjacent string. In HiP, a similar setup was
followed. At these sites, temperature readings were recorded at hourly intervals. Soil type and associated data were provided by South African National Parks.

**Analyses**

Species accumulation curves were constructed in EstimateS for each savanna type in order to assess sampling adequacy (Gotelli and Colwell 2001). This was achieved by using the nonparametric incidence coverage estimator (ICE) (Chazdon et al. 1998), Chao 2, Jack 2 and Michaelis-Menten richness estimators (Colwell and Coddington 1994), all estimators recommended for small sample sizes (Colwell and Coddington 1994, Magurran 2004). If the sample-based rarefaction curves and these estimators converge closely at the highest observed species richness, this observed richness can be considered representative (Longino et al. 2002, Magurran 2004). Multiple estimators were used to assess sampling adequacy because each one is influenced differently by species abundance data and it is difficult to determine which one is the least biased as it involves factors such as community evenness and sampling intensity (Brose et al. 2003). This analysis showed that sampling was incomplete at Satara and Pretoriuskop, but satisfactory at the other sites. However, although sample-based rarefaction curves (i.e. observed richness) together with the richness estimators started flattening off for most study regions (Supplementary materials, Fig. S3), typically they did not reach an asymptote (based on singleton and doubleton values), especially at Satara. Additionally, when each season’s species richness estimates were considered separately, an asymptote was not reached. I therefore opted to use the raw species richness values in subsequent analyses because the estimators were considered unreliable and hence the observed richness values were considered to be the best available estimate of richness.

Observed species richness and overall abundance (the latter was only obtained for the wet season sampling) were compared between the savanna types, seasons and burning regimes using one-way analysis of variance (ANOVA) when data were parametrically distributed. Where data were non-parametrically distributed and could not be easily transformed, appropriate non-parametric tests were applied such as a Kruskal-Wallis ANOVA. Due to difficulties associated with collecting and counting all termites found at a bait or in a block of a belt transect, the number of encounters for each method was
considered as a surrogate for termite abundance. Abundance is defined as the total number of encounters for each termite species in a transect or grid of baits. An encounter is an occurrence of a population of termites of one species at one point (bait or block of transect) (Davies et al. 2003).

Termites were classified into feeding groups based on the classification of Donovan et al. (2001), which places termites into four groups dependent on the humification of the feeding substrate, which is a measure of the stage of decomposition this feeding substrate is in and ranges from living plants and trees to organic material in the soil (Donovan et al. 2001). Group I comprises the non-termitids, all of which feed on dead wood and grass; Species in groups II-IV are all Termitidae. Group II has a range of feeding habits, which include dead wood, grass, leaf litter and micro-epiphytes; group III feed in the organic rich upper soil layers (broadly ‘humus-feeders’); and group IV contains the true soil-feeders which ingest mineral soil. Different termite feeding groups have different ecological roles and so understanding how these groups differ between savanna types, seasons or burning regimes is important ecologically.

Differences in termite assemblage composition, based on the data collected for each sampling method respectively, were analysed between savanna types, seasons and burning regimes using multivariate analyses in Primer v. 5.2 (Clarke and Gorley 2001). Bray-Curtis similarity matrices of the data were constructed in order to assess overall compositional differences. The data were square root transformed before similarity matrices were constructed in order to weight common and rare species equally (Clarke and Warwick 2001).

Analysis of similarity (ANOSIM, Clarke 1994) was used to test if there were significant differences between the assemblages; this is a non-parametric permutation procedure applied to rank similarity matrices which underlie sample ordinations (Clarke and Warwick 2001, Parr et al. 2004), in which a significant global R-statistic of close to one indicates distinct differences between the assemblages compared. Negative values of R indicate that there are more differences between sites of the same treatment than between treatments, and R can take on any value between -1 and 1.

The R-value itself is not unduly affected by the number of replicates in a sample, however, the significance level of this R-statistic is. This can result in a situation where
the R-value is, for example, very low, which indicates that assemblages are barely separable, but this value may be significantly different from zero, perhaps leading to an interpretation that the assemblages in question are indeed different. However, this significance level is more a reflection of a high number of replicates or samples, and the R-value is of greater importance for interpretation (Clarke and Warwick 1994, Parr et al. 2004). The opposite may also be true with an R-value being close to one. Similarity percentage analysis (SIMPER) from Primer (Clarke and Warwick 2001) was then conducted on the wet season data in order to identify which species were contributing most to differences between these assemblages.

Termite activity levels were also assessed and compared between savanna types and burning regimes indirectly as a percentage of the baits attacked as well as the intensity of attack on these baits. These data were compared using ANOVA where the data were normally distributed. For data which could not be easily transformed, non-parametric tests, Mann-Whitney U and Kruskal-Wallis ANOVA were applied to the data. Since animal disturbance resulted in a different number of baits been available for termite sampling, the number of encounters and richness values obtained were corrected for the number of baits per site (the number of encounters or species were divided by the number of baits present, giving the number of encounters or species per bait, and this total multiplied by 40 for each site, giving the number of encounters or species per 40 baits).

Results

Across savanna habitats and seasons

Species richness and abundance
Termites from 18 genera and 5 subfamilies were collected during the study; the genus Anenteotermes being recorded from South Africa for the first time (Appendix A). When data from all sampling methods and seasons were pooled, there was a significant difference in species richness across the four savanna habitats (ANOVA, $F_{3,43} = 5.016$, $P < 0.005$). Mopani had significantly lower species richness (mean of 4.50 species plot$^{-1}$, SE = 0.47) than Satara and Pretoriuskop. Satara (mean of 6.75 species plot$^{-1}$, SE = 0.39) and Pretoriuskop (mean of 7.08 species plot$^{-1}$, SE = 0.48) had higher richness while Hluhluwe had an intermediate species richness (mean of 5.27 species plot$^{-1}$, SE = 0.81).
An examination of species feeding groups indicates however, that termites at Mopani and Satara differ to those at the mesic sites (Pretoriuskop and Hluhluwe): Feeding group II dominates the assemblage at Mopani and Satara, whereas the mesic sites are more evenly split between Feeding groups II and III, with Feeding Group IV also being present (Fig. 2).

Using bait data, there is a significant seasonal effect on observed species richness in all savanna types except Mopani (Kruskal-Wallis ANOVA, $H_{2,27} = 1.584, P > 0.05$) with the wet season, in general, being most species rich (Fig. 3, see also Appendix A). Termite abundance also differed significantly across the four savannas (Fig. 4a; ANOVA, $F_{3,43} = 14.488, P < 0.000001$), with the highest number of encounters at Satara. Considering active searching data, Mopani had significantly fewer occurrences compared with the other three savannas (Fig. 4a; ANOVA, $F_{3,39} = 4.049, P < 0.05$).

**Assemblage composition**

Considering all possible comparisons across savanna types (pair-wise tests), analysis of similarity (ANOSIM) revealed differences in termite assemblages between savannas were most pronounced in the wet season compared to those in the dry and transitional seasons (Table 1). Pair-wise tests between Mopani and Pretoriuskop contributed most to the overall assemblage differences across seasons followed by Satara vs. Pretoriuskop.

**Frequency of attack**

For all seasons the highest percentage of baits attacked was at Satara (Fig. 5a, ANOVA, $F_{3,43} = 15.089, P < 0.000001$; $F_{3,43} = 83.988, P < 0.000001$; $F_{2,33} = 12.526, P < 0.0001$ for the wet, dry and transitional seasons respectively), and was highest in the wet and transitional seasons and lowest in the dry season for all savannas (Fig. 5a, ANOVA, $F_{2,33} = 5.445, P < 0.01$; $F_{2,33} = 28.442, P < 0.000001$; $F_{2,33} = 25.828, P < 0.000001$; Mann-Whitney U, $U = 16.500, P < 0.01$ for Mopani, Satara, Pretoriuskop and Hluhluwe respectively). Post-hoc Tukey HSD tests revealed that while the reduction in frequency of attack from the wet to the dry season was significant for all the savanna types examined, it was especially pronounced at the more mesic sites (Pretoriuskop and Hluhluwe) (Fig. 5a).
3. Termite responses and fire

*Intensity of attack*

In all three seasons, the intensity of termite attack was significantly higher at Satara than in the other savannas, which showed very similar intensity levels (however see Mopani when all seasons are combined where the IA is significantly higher than that at Hluhluwe) (Kruskal-Wallis ANOVA, $H_{3,779} = 87.048, P < 0.0001, H_{3,691} = 91.059, P < 0.0001, H_{2,592} = 73.194, P < 0.0001, H_{2,2062} = 219.079, P < 0.0001$ for the wet, dry, transitional and all seasons pooled respectively, Fig 5b).

Intensity of attack also differed significantly across seasons at all four sites (Mopani: Kruskal-Wallis ANOVA, $H_{2,503} = 19.054, P < 0.005$; Satara: Kruskal-Wallis ANOVA, $H_{2,604} = 55.543, P < 0.0001$; Pretoriuskop: Kruskal-Wallis ANOVA, $H_{2,634} = 107.996, P < 0.001$; Hluhluwe: Mann-Whitney U, $U = 8757.5, P < 0.000005$), with the dry season having significantly lower levels of attack than the wet and transitional seasons which did not differ from each other (except at Mopani where the transitional season does not differ from either the wet or dry seasons) (Fig. 5b).

*Fire effects*

*Species richness and abundance*

There were few clear differences in species richness and occurrence of termites across different fire regimes within each savanna type (Figs. 1, and 4b & c for richness and occurrence respectively). For species richness, the only significant difference was at Pretoriuskop where the August 3 plot had more termite species than the August 1 plot. In terms of the effect of fire on termite abundance, there were only two significant differences, one at Mopani for the AS data (wet season), where the December 3 plot has significantly more termites than the August 1 and Unburnt plots (Fig.4b) and one at Pretoriuskop for the bait data (wet season) where the August 1 plot has significantly more termites than the Unburnt plot (Fig. 4c).

*Assemblage composition*

When examining differences in termite assemblage composition with burning regime, the results vary with sampling method. For the active searching data, assemblage
composition differences between the burning regimes become less pronounced as sites become more mesic: there are three pairs of regime that differ significantly at Mopani (Unburnt vs. August 2, December 3 vs. August 1 and August 2), two at Satara (Unburnt vs. August 1 and August 3) and only one at Pretoriuskop (August 3 vs. August 1) (Table 2). In contrast, when the bait data are considered, differences become more pronounced as sites become more mesic: there are three pairs of regime differences at Pretoriuskop (Unburnt vs. August 1 and December 3, December 3 vs. August 1), one at Satara (December 3 vs. August 1) and none at Mopani. During the dry season, assemblage composition is unaffected by burning regimes, and the transitional season also produces few differences, with the exception being at Mopani (Table 2).

ANOSIM reveals significant difference in assemblage composition with sampling method (active searching and baits) at Mopani and Pretoriuskop (R = 0.601, p = 0.001 and 0.498, p = 0.001 for Mopani and Pretoriuskop respectively). The differences are less pronounced at Satara (R = 0.247, p = 0.001). The sites where the assemblages are more dissimilar are also the savanna types which show more differences in assemblage response to fire, suggesting that different suites of termites (i.e. Feeding Groups) sampled by the different methods respond differently to fire (Table 3). Active searching targets a broader range of Feeding Groups while baits attract Feeding Group II primarily. In addition, using active searching along the rainfall gradient demonstrated there is a shift in Feeding Group with savanna type: Feeding Group III and IV were sampled primarily at Pretoriuskop, while at Mopani only Feeding Group I and II were sampled. SIMPER showed that most differences were driven by FG II, indicating differences in abundances are more important, however, this did vary with savanna type and sampling method (Supplementary material, Table S4).

Frequency and intensity of attack
There is no significant difference in the percentage of baits attacked by termites between burning regimes at any of the four savanna types in any of the three seasons. However, with intensity of attack, in KNP the response to burning regime becomes more pronounced as sites become more mesic (Fig. 6); there is no difference in termite attack
between any of the burning regimes at Mopani (Fig. 6a), compared with one difference found in Satara (Fig. 6b) and three differences found at Pretoriuskop (Fig. 6c).

Environmental variables

Data for the environmental variables are shown in Table 3. Grass cover generally increased as fire intensity and frequency decreased as did canopy cover. Bare ground showed the opposite relationship, increasing with fire intensity and frequency. Temperatures decreased with increasing rainfall and latitude.

Discussion

Diversity across savanna habitats

Termite diversity differed significantly along the rainfall gradient in terms of species richness, abundance and assemblage composition. This corresponds to extensive variation in rainfall, vegetation and soil type. In semiarid areas such as South Africa, precipitation sets limits to net primary productivity (NPP), giving rise to a strong association between these two variables (see O'Brien 1998, O'Brien et al. 2000, van Rensburg et al. 2002). It is therefore reasonable to assume that changes in rainfall result in changes in NPP across the study sites, with corresponding impacts on termite diversity, as predicted by species-energy theory (see Rosenzweig and Abramsky 1993, Wright et al. 1993). Indeed, Mopani (the most arid site, and presumably with the lowest NPP) had significantly lower species richness and abundance than all other sites. This is in keeping with Eggleton et al. (1994), who found a moderately strong relationship between NPP and termite diversity within biogeographic regions.

Although termite diversity is highest in the warm, wet tropics (Eggleton et al. 1994, Eggleton 2000), I found species richness was highest at Pretoriuskop, before declining at Hluhluwe, the wettest site. Although Hluhluwe has the highest rainfall, the higher elevation and generally increased cloud cover and mist at this site result in lower temperatures than those along the gradient further north, perhaps explaining to some extent the decrease in species richness. Termites are susceptible to temperature changes (Ueckert et al. 1976, Ohiahu 1979, Smith and Rust 1994, Korb and Linsenmair 1998), and soil-feeders are reported to be excluded by low temperatures (Bignell and Eggleton...
2000, Donovan et al. 2002, Palin et al. in press), hence a decrease in diversity and proportion of soil feeders (relative to Pretoriuskop) at Hluhluwe. Independent of burning regimes, interactions between rainfall, NPP and temperature thus seem important for termite diversity responses in savannas.

The clearest trend across the investigated rainfall gradient is that the termite assemblage changes, especially in functional group composition. Because soil feeders are especially susceptible to habitat desiccation and require malleable, carbon-rich soils (Bignell and Eggleton 2000) they tend to be excluded from drier sites; none were sampled at Mopani and although a few species occurred at Satara their abundances were relatively low. Throughout the tropics closed canopy forest has a higher diversity of termites, this is likely to be due to environmental stability and resource availability in tropical rain forests (Eggleton et al. 1994, Eggleton 2000). Similarly, savannas with higher rainfall and more canopy cover are likely to be more favourable for termites. It is therefore unsurprising that the mesic sites (Pretoriuskop and Hluhluwe) have higher levels of termite diversity. Eggleton et al. (1995) and Eggleton et al. (1996) found that in tropical forests when disturbance occurs which opens up the canopy, there is a decrease in termite diversity, particularly of soil feeders (FG III and IV). The absence of these FGs in the drier savanna sites would suggest that this pattern is also evident on the landscape scale.

Variation in soil type across the savanna types (clay at the two drier sites and sandy soils at the mesic sites) is also likely to influence termites since they are considered to be soil animals (Holt and Lepage 2000). Termites have a profound affect on soil characteristics (see Holt and Lepage (2000) for a review), and are, in turn, strongly influenced by soil properties (see Lee and Wood (1971) for a review). Close inspection of the assemblages sampled suggests that sites with a similar broad soil type are more similar, and this factor cannot be ignored when seeking to understand why these termite communities differ. The ability of mound building and subterranean termites to construct mounds and tunnels respectively is influenced largely by the clay, silt and sand content of soils (Lee and Wood 1971). Changes in these ratios observed in this study will no doubt affect termite diversity.
3. Termite responses and fire

Only the driest site, Mopani, had significantly lower abundance when considering the active searching data (mirroring the species richness results). When baiting data are considered, Satara (low to intermediate rainfall) stands out with significantly higher abundance than all other sites. Because baiting attracts termites primarily from Feeding Group II, my results suggest that at Satara these termites dominate, being replaced by other FGs as rainfall increases, perhaps as a result of competition with fungi in wetter savannas (Aanen and Eggleton 2005). Termite activity (primarily from baiting and hence a reflection of FG II) reflected this same pattern, with significantly higher levels of attack frequency and intensity at Satara. As the range of humification of the feeding substrate decreases, humivorous termites (FG III and IV) disappear and are replaced by fungus-growers and wood-feeders (FG II); a pattern which has been shown previously in African and Asian savannas (Buxton 1981, Bignell and Eggleton 2000). Satara also has comparably higher mammalian herbivore densities than the other study sites, resulting in higher concentrations of dung in these areas. This higher dung concentration may lead to higher nutrient concentrations in the soil as well as to more available food for termites, which feed on drier dung material, especially termites from FG II (Freymann et al. 2008). The processing of dung material by termites may in turn facilitate nutrient cycling, increasing plant productivity (and possibly grass nutrient quality) making the area more favourable for herbivores, thus creating a positive feedback loop. Similar patterns have been documented by Gosling (unpublished) in Hluhluwe-iMfolozi Park: higher levels of termite activity (of FG II) were recorded in the drier iMfolozi region than the mesic Hluhluwe. At iMfolozi, herbivore densities are also higher, presumably providing more nutrients for the termites. The feedback link between herbivore and termite densities remains, however, to be tested.

Seasonal effects
The higher levels of termite activity and species richness during the wet and transitional seasons follow the general trend for insect activity (Wolda 1988) and similar findings for termites recorded elsewhere (e.g. west Africa: Ohiagu (1979), Western Australia: Abensperg-Traun (1991), northern Australia: Dawes-Gromadzki and Spain (2003), but see Dibog et al. (1998) where tropical forest termites in Cameroon were largely
unaffected by seasonality). This is likely to be a reflection of greater food and moisture availability during the wet season and the effect extends into the transitional season due to a lag effect on vegetation growth and also because this season may be better suited to termite foraging than the wet season (e.g. see Dawes-Gromadzki and Spain 2003). Those authors recorded higher levels of termite activity during the transitional season and deduced that this was because of heavy monsoonal rainfall in northern Australia during the wet season, causing soil to become waterlogged and hence unsuitable for termite foraging. Dibog et al. (1998) also found a negative relationship between termite activity and rainfall, especially within a two day period following a rainfall event, and concluded that this was possibly due to flooding of termite subterranean galleries. In this study, the mesic savanna at Pretoriuskop showed higher activity levels during the transitional season, possibly also due to waterlogging of soils and above-ground seepage during the wet season, which was apparent during sampling. In the more xeric savanna sites, activity and diversity peaked during the wet season rather than the transitional and is probably because waterlogged soils and flooding are not common features here, nor the rainfall enough to have a pronounced lag effect into the transitional season, suggesting that termites have an optimal rainfall range for activity, being inactive when conditions are excessively wet or dry. Unfortunately, transitional season data are unavailable for Hluhluwe.

Being soft-bodied and susceptible to desiccation, termites are also likely to move deeper into the soil during dry periods, becoming biologically inactive and less likely to be sampled with baiting or active searching. During the wet season, as the upper horizons of the soil become moister, termites move up in the profile and become more susceptible to sampling, thereby increasing recorded activity levels.

Fire effects
This study used long-term burning regimes, a broader range of fire treatments and a wider variety of sampling methods than previous studies (Chapter 2). This enabled us to assess fire effects on the entire termite assemblage. Termites have subterranean habits and are protected from the direct effects of fire, so the effects of repeated burning are thus likely
3. Termite responses and fire

to be primarily indirect, through changes to vegetation structure and resource (food) availability.

Overall, termite assemblages in this study showed striking resilience to fire: species richness and abundance showed few significant differences across burning regimes, although there were some differences between burning treatments at Pretoriuskop. This is in keeping with a previous study carried out on savanna ant assemblages using the same experimental burning plots (Parr et al. 2004), as well as studies elsewhere on savanna insect responses to burning (Andersen et al. 2003, Andersen et al. 2005) and for biodiversity more broadly (Parr and Andersen 2006). My results contrast, however, with studies on termites and fire from elsewhere (see Chapter 2 for a review). The contrast with other studies is likely to be because most previous studies focused solely on mound-builders and sampled only extremes of fire regimes (annual and unburnt) (e.g. Benzie 1986, Traoré and Lepage 2008).

While little response to burning was detected when examining species richness and abundance, at the assemblage level some patterns could be detected. This aspect of termite diversity has not been investigated previously within long-term burning regimes. Additionally, my study revealed that the sampling method used also had an effect on the termite responses recorded, indicating that different suites of termites (sampled by different methods) respond differently to fire; it may thus be difficult to get a full understanding of the effect of fire on termites unless a variety of sampling methods are employed.

When active searching (which samples the entire range of Feeding Groups, although still has a seasonal bias) was used, more pronounced differences in assemblage composition between fire treatments were recorded at the drier sites than the wettest (Pretoriuskop). Since soil feeding termites are more susceptible to desiccation and disturbance than other groups (Eggleton et al. 1995, 1996), at arid sites they may better be able to survive on unburnt sites which have more canopy and grass cover. Indeed *Promirotermes* spp., and *Astialotermes* spp., both FG III, were found almost exclusively on unburnt plots at Satara. However the presence of FG I termites (*Hodotermes mossambicus*) at the most arid site (Mopani) may have biased results here; those termites were only sampled at the December 3 plot and tend to occur in a patchy, sporadic fashion.
3. Termite responses and fire

(personal observation). These grass harvesters contributed 10% of the dissimilarity between the December 3 and August burns and did not occur on other fire plots. These results should also be treated with some caution as active searching has been little tested in arid savannas (Chapter 4, but see Zeidler et al. (2004) for a comparison with baiting) and its success is likely to be closely associated with the occurrence of rainfall events in those environments. If sampling had occurred on another day, *H. mossambicus* may well have not been sampled.

However, when the bait data are considered, responses to fire at the assemblage level differ more strikingly along the rainfall gradient. Baiting targets Feeding Group II, the most dominant, diverse and widespread feeding group in savannas and thus probably the most important from a functional perspective. My results suggest that for these termites, burning has differing levels of effect across the savanna habitats, and that effect is probably dependent on vegetational effects. At the arid Mopani site, there are negligible effects; here, FG III and IV are absent and vegetation is little affected by fire, resulting in few differences in the overall habitat and hence no measurable effect on termite composition.

In contrast, as rainfall increases, fire has a more pronounced effect on the vegetation (Parr et al. 2004), and the effect on termite diversity and activity becomes concomitantly more noticeable. At Pretoriuskop, the most striking differences are between unburnt and intense (August) fires, as well as between frequent, intense fires (August 1) and cool, less frequent fires (December 3). Since baiting does not generally attract soil feeders, these differences are likely to be attributed to differences in relative occurrences of termite species. Indeed, the differences between these plots are driven predominantly by the genus *Microtermes*, which is present on all plots but is most abundant on burnt (especially frequently burnt) plots. This would support the idea of FG II termites been excluded by soil feeders as burning regimes become less frequent and intense, allowing the humification of the feeding substrate to increase and the subsequent dominance of FG III and IV (Buxton 1981, Bignell and Eggleton 2000) where they can compete with fungi (Aanen and Eggleton 2005), indeed more termites from FG III and IV were present on unburnt plots (mean per plot) in the present study.
At the wettest site (Pretoriuskop), there is also a significant trend of more termites and activity with increasing fire intensity, and especially frequency. As discussed above, previous studies have found that as aridity increases (Eggleton et al. 1995, 1996) and the subsequent humification of the feeding substrate decreases (Buxton 1981, Bignell and Eggleton 2000), humivorous termites are excluded and replaced with fungus-growers and wood-feeders (which are attracted to baits). It would appear that fire has a similar effect in these mesic savannas, opening up the canopy for increased desiccation of the substrate, allowing fungus-growers and wood feeders of Feeding Group II to replace soil feeding termites, possibly as a result of fungus-growing termites being able to outcompete fungal decomposers in more desiccated environments where soil-feeders cannot survive (Aanen and Eggleton 2005). Also of importance here are the increased activity levels on annually burnt plots, which are possibly linked to vertebrate herbivore densities. Herbivores are likely to favour grazing on burnt plots due to higher forage potential and visibility (Wilsey 1996, Archibald and Bond 2004, Parrini and Owen-Smith 2009). This could result in a positive-feedback loop as more dung will be available here, both as food for termites (of FG II) and for increased soil nutrients which may influence termite distributions (Crist 1998).

Conclusions
Sampling termites from a variety of methods, although important in order adequately to sample all Feeding Groups, makes it difficult to draw broad conclusions because different data sets cannot always be combined. In addition, different termite feeding groups seem to react differently to fire, again making conclusions difficult.

However, there are distinct differences along the rainfall gradient, in terms of both termite diversity across savanna types and termite responses to fire. While diversity increases with rainfall so do responses to the long-term burning regimes. Although, termite communities were, overall, highly resilient to fire, the greatest effects were observed at Pretoriuskop (the most mesic site investigated) in terms of species richness, abundance, assemblage composition and activity levels (in terms of the dominant FG).

Re-visiting the concept suggested by Sankaran et al. (2005) of stable and unstable savannas dependent on rainfall, in this study Mopani and Satara would represent stable
3. Termite responses and fire

savannas and Pretoriuskop an unstable savanna. Sankaran et al. (2005) argue that fire has increased effect in unstable savannas in terms of vegetation structure and canopy cover, and my results indicate that faunal responses seem to follow this same pattern, at least with regard to termites, especially those attracted to baits.

References


3. Termite responses and fire


3. Termite responses and fire

**Table 1:** Analysis of similarity (ANOSIM) for termite assemblages between four savanna habitats, across three seasons. For the dry and transitional seasons, presence/absence data were used (data from active searching and baits combined), and for the wet season abundance data were used (data from the two sampling methods analysed separately). AS = active searching. Baits = cellulose baits. The R-statistic is a measure of similarity of assemblages, the closer this value is to 1, the more dissimilar assemblages are (Clarke & Warwick 2001). \( P \) is considered significant at the \( \alpha \)-level of 0.05. Highlighted values are those above or equal to 0.250 which are considered to be markedly dissimilar (Clark and Warwick 2001).

<table>
<thead>
<tr>
<th>Wet (AS)</th>
<th>Wet (baits) Dry</th>
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</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>R</td>
<td>P</td>
</tr>
<tr>
<td>Global R</td>
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<td>0.001</td>
</tr>
</tbody>
</table>

**Paired savanna comparisons**

- Mopani vs. Satara: 0.398 0.005 0.420 0.001 0.185 0.012 0.074 0.141
- Mopani vs. Pretoriuskop: 0.798 0.001 0.434 0.001 0.393 0.001 0.191 0.004
- Mopani vs. Hluhluwe: 0.376 0.002 0.287 0.002 0.203 0.006
- Satara vs. Pretoriuskop: 0.445 0.001 0.574 0.001 0.326 0.001 0.074 0.074
- Satara vs. Hluhluwe: 0.300 0.001 0.401 0.001 0.389 0.001
- Pretoriuskop vs. Hluhluwe: 0.339 0.001 0.242 0.011 0.262 0.004
3. Termite responses and fire

Table 2: Analysis of similarity (ANOSIM) for termite assemblages between four burning regimes at three savanna habitats and during three seasons. For the dry and transitional seasons, presence/absence data were used (data from active searching and baits combined), and for the wet season abundance data were used (data from the two sampling methods analysed separately). AS = active searching. Baits = cellulose baits. The R-statistic is a measure of similarity of assemblages, the closer this value is to 1, the more dissimilar assemblages are (Clarke and Warwick 2001). P is considered significant at the α-level of 0.05. Highlighted values are those above or equal to 0.250 which are considered to be markedly dissimilar (Clark and Warwick 2001).

<table>
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<th>Wet AS</th>
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<td>R</td>
<td>R</td>
<td>R</td>
<td>R</td>
</tr>
<tr>
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<td></td>
<td></td>
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<td>0.000</td>
<td>-0.111</td>
<td>0.241</td>
</tr>
<tr>
<td>Unburnt vs. August 2</td>
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<td><strong>0.296</strong></td>
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<tr>
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<td>-0.111</td>
<td>-0.093</td>
<td>0.241</td>
</tr>
<tr>
<td>December 3 vs. August 1</td>
<td><strong>0.750</strong></td>
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<td>-0.222</td>
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<tr>
<td><strong>Satara</strong></td>
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<td>-0.074</td>
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<td><strong>Pretoriuskop</strong></td>
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<td></td>
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<td>Global R</td>
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<td>-0.069</td>
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55
3. Termite responses and fire

<table>
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<th></th>
<th>Unburnt vs. August 1</th>
<th>Unburnt vs. August 3</th>
<th>Unburnt vs. December 3</th>
<th>December 3 vs. August 1</th>
<th>December 3 vs. August 3</th>
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</table>
Table 3: Mean percentage values for vegetation and overhead canopy cover as well as mean temperature readings recorded during the wet season at each savanna type and burning regime. G = grass, DGL = dead grass litter, DLL = dead leaf litter, BG = bare ground, F = forbs, WV = woody vegetation, DW = dead wood, R = rock and D = dung.

<table>
<thead>
<tr>
<th>Site</th>
<th>G</th>
<th>DGL</th>
<th>DLL</th>
<th>BG</th>
<th>F</th>
<th>WV</th>
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<th>R</th>
<th>D</th>
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<td>0.80</td>
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<tr>
<td>August 1</td>
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<td>1.55</td>
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<td>Hluhluwe</td>
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<td>14.45</td>
<td>23.45</td>
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</table>
3. Termite responses and fire

Figure 1: Mean (± 1 SE) observed termite species richness for each burning regime across three savanna types for all methods and seasons combined. Symbols denote significant differences. Letters denote sites which are the same statistically within a savanna habitat, differences are considered significant at the α-level of 0.05.
Figure 2: The relative contribution (measured as a percentage) each feeding group makes to the termite assemblage. Data are from all seasons and all sampling methods. Feeding group I comprises the lower termites which feed on dead wood and grass; group II is made up of Termitidae with a range of feeding habits which include dead wood, grass, leaf litter and micro-epiphytes; group III feed in the organic rich upper soil layers and are also Termitidae; and group IV contains the true soil-feeders which ingest mineral soil, again all Termitidae.
3. Termite responses and fire

![Graph showing mean termite species richness across seasons and savanna types]

**Figure 3:** Mean (± 1 SE) observed termite species richness recorded at paper baits (both buried and on the surface) across three seasons and four savanna types. Letters denote sites which are the same statistically within one savanna habitat; differences are considered significant at the α-level of 0.05.
3. Termite responses and fire

a.

![Graph showing termite responses and fire at different savanna sites.

- **Satara**
  - Mean number of encounters: b
  - Legend: ■ Cellulose baits  □ Active searching

b.

![Graph showing termite responses at different savanna sites following fire.

- **Satara**
  - Number of encounters: a
- **Pretoriuskop**
  - Number of encounters: a

Legend: ■ Unburnt  □ December 3  □ August 3  □ August 1

- **Mopani**
  - Number of encounters: a

Savanna site

Unburnt  December 3  August 3  August 1

Mean number of encounters

0  5  10  15  20  25  30

Mopani  Satara  Pretoriuskop  Hluhluwe

Digitised by the University of Pretoria, Library Services, 2012
c.

Figure 4: Overall termite abundance a) between savanna types for each wet season method separately and for b) active searching data and c) cellulose baits between burning regimes at each savanna type during the wet season. Letters denote sites which differ significantly within the respective savanna habitat, with differences being considered significant at the α-level of 0.05.
3. Termite responses and fire

**Figure 5**: Mean (± 1 SE) a) frequency and b) intensity of termite attack recorded at paper baits (both buried and on the surface) at four savanna sites and across three seasons. An asterisk denotes significant differences between savannas in a given season and letters denote significant seasonal differences within savanna sites, differences are considered significant at the α-level of 0.05.
3. Termite responses and fire

a. 

b. 

c. 

64
Figure 6: Mean (± 1 SE) intensity of termite attack recorded at paper baits (both buried and on the surface) across four burning regimes and three seasons at a) Mopani, b) Satara, and c) Pretoriuskop. Arrows distinguish sites which differ significantly, with differences being considered significant at the α-level of 0.05.
Appendix A

Termite species sampled in the four savanna habitats with their respective feeding groups. Codes in parenthesis indicate the season/s in which a particular species was sampled; w = wet, t = transitional, and d = dry season. Feeding group I comprises the lower termites which feed on dead wood and grass; group II is made up of Termitidae with a range of feeding habits which include dead wood, grass, leaf litter and micro-epiphytes; group III feed in the organic rich upper soil layers and are also Termitidae; and group IV contains the true soil-feeders which ingest mineral soil, again all Termitidae.

<table>
<thead>
<tr>
<th>Species</th>
<th>Mopani</th>
<th>Satara</th>
<th>Pretoriuskop</th>
<th>Hluhluwe</th>
<th>Feeding group</th>
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</thead>
<tbody>
<tr>
<td><strong>Hodotermitidae</strong></td>
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<td></td>
<td></td>
<td></td>
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</tr>
<tr>
<td><em>Hodotermitus mossambicus</em></td>
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<td>X(d)</td>
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<td>X(w,d)</td>
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<tr>
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<td>X(w,t,d)</td>
<td>X(w,t,d)</td>
<td>X(w,t,d)</td>
<td>X(w,d)</td>
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</tr>
<tr>
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<td>X(t)</td>
<td></td>
<td>X(w)</td>
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<td>X(w,t,d)</td>
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<td>X(w)</td>
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<td>X(w,d)</td>
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<td>X(w,t,d)</td>
<td>X(w,t,d)</td>
<td>X(w,d)</td>
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</tr>
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<td>X(w,d)</td>
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<td>X(w,d)</td>
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<td>X(w,t,d)</td>
<td>X(t,d)</td>
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<td>X(w)</td>
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<td><em>Anenteotermes</em> spp</td>
<td>X(w)</td>
<td></td>
<td></td>
<td></td>
<td>III</td>
</tr>
</tbody>
</table>
### Termite responses and fire

#### Astalotermes spp.
- X(w)
- X(w,t)
- X(w)
- **III**

#### Termitinae

- **Amietermes hastatus**
  - X(w)
  - X(d)
  - **II**

- **Amietermes messinae**
  - X(w)
  - X(d)
  - **II**

- **Angulitermes spp.**
  - X(w)
  - **III**

- **Cubitermes spp.**
  - X(w)
  - **IV**

- **Lepidotermes spp.**
  - X(d)
  - X(w)
  - X(w)
  - **IV**

- **Microcerotermes spp.**
  - X(w,t,d)
  - X(w,t)
  - X(w,t,d)
  - X(w,d)
  - **II**

- **Promirottermes spp.**
  - X(w)
  - X(w)
  - X(w)
  - **III**

#### Nasutitermitinae

- **Trinervitermes trinervoides**
  - X(w)
  - X(w)
  - X(w,d)
  - **II**

- **Rhadinotermes coarctatus**
  - X(w)
  - **II**
Supplementary Material

S1: Study site

The three replicates used in the Mopani section of KNP were Tsendse (23°41'S 31°31'E), Mooiplaas (23°34'S 31°27'E) and Dzombo (23°26'S 31°22'E); the replicates used in the Satara region were N'wanetsi (24°26'S 31°51'E), Marheya (24°32'S 31°46'E) and Satara (24°24'S 31°45'E); and the replicates in the Pretoriuskop area were Kambeni (25°15'S 31°26'E), Numbi (25°13'S 31°20'E) and Shabeni (25°12'S 31°23'E). In HiP, all sites were located in the northern Hluhluwe section of the reserve.

The Mopani plots were situated in the Mopani shrubveld, which is a mopani (Colophospermum mopane J. Kirk ex J. Léonard) dominated habitat with few other woody species present (Low & Rebelo 1996). The mean annual rainfall is 450 to 500 mm and the area ranges in altitude from 300 to 340 m a.s.l. (Gertenbach 1983). In the Satara region, the plots were situated in a mixed knobthorn Acacia nigrescens Oliv. and marula Sclerocarya birrea (A. Rich. Hochst.) savanna in the sweet lowveld bushveld (Low & Rebelo 1996). The area ranges in altitude from 240 to 320 m a.s.l. and has a mean annual rainfall of 550 mm (Gertenbach 1983). The Pretoriuskop region is sour lowveld bushveld, consisting of open tree savanna dominated by silver clusterleaf (Terminalia sericea Burch. ex DC), but bushwillow (Combretum colinum Fresen.) is also common (Low & Rebelo 1996). The region has a mean annual rainfall of 700 mm and the altitude varies from 560 to 640 m a.s.l. (Gertenbach 1983). The sites situated in HiP were located within Natal lowveld bushveld, dominated by various acacia species, notably A. tortilils and A. karoo., but with several other species also being common (Low & Rebelo 1996). The mean annual rainfall here is 800 to 900 mm and the altitude ranges from 500 to 750 m a.s.l. (Archibald et al. 2005). Both the Mopani and Satara regions have basalt-derived clay soils while the Pretoriuskop region has granite-derived sandy soils (Gertenbach 1983). At Hluhluwe the underlying geology is predominantly sandstone and shale resulting in sandy soils (Archibald et al. 2005).
S2: Modification of standardised transect method

The active searching transect method was modified from that of Jones & Eggleton in that less time was spent searching in each block (ten minutes in my study compared to the standard sixty minutes), this was primarily because the Jones & Eggleton method was designed for tropical forests which have a much higher termite diversity and density than semi-arid savannas and are also much easier to sample given the softer soils and more variable microhabitats. Additional logistical constraints prevented more time been spent searching each block. In this study, the transect was also divided into two 50 metre transects because of the small size of the burn plots and in order to limit any edge effects.
3. Termite responses and fire

a)

- S Obs
- ICE Mean
- Chao 2 Mean
- Jack 2 Mean
- MM Mean

b)
Figure S3: Species accumulation curves for all seasons data pooled, for a) Mopani, b) Satara, c) Pretoriuskop and d) Hluhluwe using estimators (ICE, Chao 2, Jack 2, Michaelis-Menten) from EstimateS (Colwell 2005).
3. Termite responses and fire

Table S4: SIMPER analysis for termite abundance per burning regime in each savanna type. Only those pair-wise tests which were significantly different in assemblage composition are included. Av. abund. = average abundance, Av. diss. = average dissimilarity, Cum. % = cumulative percentage of similarity.

<table>
<thead>
<tr>
<th>Burning regimes</th>
<th>Species</th>
<th>Av. abund.</th>
<th>Av. Diss.</th>
<th>Cum. %</th>
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<td>Mopani</td>
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<td><em>Microtermes</em> spp.</td>
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<td>1.00</td>
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<tr>
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<td>0.50</td>
<td>0.00</td>
<td>14.00</td>
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<tr>
<td></td>
<td><em>Allodontermes rhodesiensis</em></td>
<td>0.50</td>
<td>0.00</td>
<td>14.00</td>
</tr>
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<td>December 3 vs. August 1 (AS)</td>
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<td>3.50</td>
<td>22.81</td>
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<td>1.00</td>
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<td>1.00</td>
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<td>1.00</td>
<td>7.14</td>
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<tr>
<td>Satara</td>
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<td>Unburnt vs. August 1 (AS)</td>
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### 3. Termite responses and fire

<table>
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<tr>
<th>Termite Species</th>
<th>Unburnt vs. August 3 (AS)</th>
<th>December 3 vs. August 1 (baits)</th>
<th>Pretoriuskop August 3 vs. August 1 (AS)</th>
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<td>0.67</td>
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<td>6.78</td>
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</table>

Unburnt vs. August 3 (AS)

- Microcerotermes spp.: 3.33
- Astalofermes rhodesiensis: 2.67
- Ancistrofermes latinotus: 4.00
- Microfermes spp.: 5.33
- Odontotermes sp 3: 0.67
- Promirotermes spp.: 1.33
- Astalofermes spp.: 1.33
- Amitermes hastatus: 0.33
- Amitermes messinae: 0.33
- Trinervitermes trinervoides: 0.33

December 3 vs. August 1 (baits)

- Odontotermes 3 sp: 32.00
- Ancistrofermes latinotus: 1.00
- Macrofermes michaelseni: 4.33
- Microfermes spp.: 7.67
- Microcerotermes spp.: 0.33
- Allofermes rhodesiensis: 0.67

Pretoriuskop

- Astalofermes spp.: 5.00
- Allofermes spp.: 2.67
- Microfermes spp.: 1.67
- Allofermes rhodesiensis: 1.00
- Ancistrofermes latinotus: 1.00
- Alyscofermes spp.: 0.67
- Adaphriotermes spp.: 1.00
- Promirotermes spp.: 0.67
### 3. Termite responses and fire

<table>
<thead>
<tr>
<th>Termite Species</th>
<th>August 1</th>
<th>December 3</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Microtermes spp.</em></td>
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<td>10.67</td>
<td>5.00</td>
</tr>
<tr>
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<td>1.67</td>
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<tr>
<td><em>Odontotermes sp 1</em></td>
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<td>-0.33</td>
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<td>0.67</td>
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</table>

#### Unburnt vs. August 1 (baits)

<table>
<thead>
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<th>Termite Species</th>
<th>August 1</th>
<th>December 3</th>
<th>Change</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Microtermes spp.</em></td>
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<td>1.34</td>
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<td>-0.33</td>
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#### Unburnt vs. December 3 (baits)

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<th>December 3</th>
<th>Change</th>
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#### December 3 vs. August 1 (baits)

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<th>August 1</th>
<th>December 3</th>
<th>Change</th>
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<tr>
<td><em>Microtermes spp.</em></td>
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