

Population dynamics of *Thaumastocoris peregrinus* in *Eucalyptus* plantations of South Africa

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Abstract *Thaumastocoris peregrinus* is a sap-sucking insect that infests non-native *Eucalyptus* plantations in Africa, New Zealand, South America and parts of Southern Europe, in addition to street trees in parts of its native range of Australia. In South Africa, pronounced fluctuations in the population densities have been observed. To characterise spatiotemporal variability in *T. peregrinus* abundance and the factors that might influence it, we monitored adult population densities at six sites in the main eucalypt growing regions of South Africa. At each site, twenty yellow sticky traps were monitored weekly for 30 months,

together with climatic data. We also characterised the influence of temperature on growth and survival experimentally and used this to model how temperature may influence population dynamics. *T. peregrinus* was present throughout the year at all sites, with annual site-specific peaks in abundance. Peaks occurred during autumn (February–April) for the Pretoria site, summer (November–January) for the Zululand site and spring (August–October) for the Tzaneen, Sabie and Piet Retief monitoring sites. Temperature (both experimental and field-collected), humidity and rainfall were mostly weakly, or not at all, associated with population fluctuations. It is clear that a complex interaction of these and other factors (e.g. host quality) influence population fluctuations in an annual, site specific cycle. The results obtained not only provide insights into the biology of *T. peregrinus*, but will also be important for future planning of monitoring and control programs using semiochemicals, chemical insecticides or biological control agents.

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Introduction

Thaumastocoris peregrinus Carpintero and Dellapé (Hemiptera: Thaumastocoridae: Thaumastocorinae) is a widely distributed eucalypt pest of Australasian origin (Carpintero and Dellapé 2006; Noack et al. 2011; Laudonia and Sasso 2012; Nadel and Noack 2012). Prior to reaching significant infestation levels on planted eucalypt trees in Sydney, Australia in 2002, very little research had been undertaken on any species of *Thaumastocoris* (or any of the Thaumastocoridae) (Noack and Coviella 2006; Noack

2009; Noack et al. 2011; Nadel and Noack 2012). *T. peregrinus* has become a major pest on various species of *Eucalyptus* across climatic regions in Southern Africa, South America, Southern Europe and New Zealand (Jacobs and Naser 2005; Carpintero and Dellapé 2006; Noack and Coviella 2006; Martínez and Bianchi 2010; Nadel et al. 2010; Wilcken et al. 2010; Laudonia and Sasso 2012; Nadel and Noack 2012; Sopow et al. 2012). Knowledge regarding factors that may influence the population dynamics of this pest in several climatic regions is, however, lacking (Martins and Zarbin 2013; Oumar and Mutanga 2013).

Thaumastocoris peregrinus is a small (2–4.5 mm), sap-sucking and gregarious insect with both adults and nymphs feeding on the same leaf surface (Jacobs and Naser 2005; Noack and Rose 2007; Noack 2009). The life cycle lasts between 30 and 60 days including five nymphal instars (16–20 days) and an adult phase (\pm 14–42 days) facilitating several generations in a year (Noack and Rose 2007; Soliman et al. 2012). Lifetime fecundity is approximately 60 eggs which typically hatch within 6 days (at 17–22 and 26 °C) (Noack and Rose 2007; Soliman et al. 2012). The short life cycle and high reproductive potential of *T. peregrinus* females thus allow for the rapid build-up of populations in areas where optimal conditions for its growth and survival occur.

Thaumastocoris peregrinus is a major pest on non-native eucalypts in South Africa. Initially discovered in the Pretoria region of Gauteng in 2003, it was only recognised as a serious pest of commercially grown eucalypts in 2005 when it initially infested plantations in the Gauteng, Limpopo, Mpumalanga and North-West provinces (Jacobs and Naser 2005). The first symptoms of infestation were usually observed on the north to north-eastern edge of a compartment (Nadel et al. 2010). *T. peregrinus* infestations result in reduced photosynthetic ability and in some cases death of severely infested trees (Nadel et al. 2010).

Barcoding using COI mitochondrial DNA (mtDNA) revealed two separate introductions of *T. peregrinus* into South Africa, which very likely originated from urban centres in Australia (Nadel et al. 2010). The dominant mtDNA haplotype identified in the study by Nadel et al. (2010) was found to occur throughout South Africa, spanning all climatic regions where *Eucalyptus* is grown. Since its initial discovery in South Africa, *T. peregrinus* has spread throughout the country infesting numerous *Eucalyptus* species and commercially produced hybrid clones (Jacobs and Naser 2005; Nadel et al. 2010). The insect has also spread north in Africa, currently as far as Kenya (Nadel et al. 2010; Nadel and Noack 2012).

Initial observations and anecdotal reports have shown that *T. peregrinus* populations vary significantly during the year in South Africa, although population fluctuations in

different regions and on different *Eucalyptus* species do not exhibit a strong or apparent synchronicity. The first aim of this study was to determine critical temperature thresholds and growth responses of *T. peregrinus* to temperature under controlled laboratory conditions. The second objective was to study the timing and phenology of population fluctuations in distinct regions of South Africa, to assess evidence for synchrony and to evaluate potential climatic drivers of population dynamics.

Materials and methods

Growth and survival in relation to temperature for *T. peregrinus*

Critical thermal limits

Extreme temperatures can dramatically affect survival rates in insect populations and they can strongly influence population dynamics and persistence over time. Temperatures above and below thermal limits of an organism are defined as lethal temperatures, because they effectively result in death (Fry 1967; Block et al. 1990). An alternative method for measuring these thresholds, originally developed for experimentation on vertebrate poikilotherms, involves the measure of critical thermal activity limits, rather than lethal thresholds (Fry 1967). Critical thermal limits are defined as the maximum and minimum temperature just short of death, but where an organism loses its ability to move (Fry 1967). This method has been widely used in studies, including at least six invertebrate orders and 24 families, following the assumption that the identified thresholds closely track lethal minima and maxima (Kay and Whitford 1978; Mitchell et al. 1993; Klok and Chown 1997; Addo-Bediako et al. 2000).

To determine the critical thermal limits (Block et al. 1990) for *T. peregrinus*, 30 adults were placed into individual 1 ml sealed tubes. The tubes were submerged in a water bath, and the water temperature was either raised or lowered at a rate of 1 °C. min⁻¹, from an initial water temperature of 15 °C. All tubes were removed and inverted every minute, and the ability of the *T. peregrinus* adults to right themselves was assessed. The critical thermal limits (CT_{max} and CT_{min}) for each *T. peregrinus* adult were defined as the maximum and minimum temperature at which the adult could no longer right itself.

Egg developmental rate

To determine the effect of temperature on the rate of *T. peregrinus* egg development, eggs of known age were required. More than a hundred *T. peregrinus* adults were

collected in Pretoria and placed in a sealed container to which paper towelling had been added. Forty eggs were removed from the paper towelling within 2 h after the adult insects had been collected. Five eggs per treatment were placed onto moist filter paper and sealed in a Petri dish. Petri dishes were placed into one of eight incubators maintained at a constant temperature ranging from 0 to 35 °C at five degree intervals and were monitored every 12 h to record the number of hatched eggs. To determine the temperature at which egg development ceased (developmental zero), we examined the straight line linear relationship between rearing temperature and days to hatch (Campbell et al. 1974). The developmental zero of the eggs was estimated as being the point where the regression line intersected the temperature axis. To determine the accuracy of the developmental zero estimate and optimum developmental rate, eggs maintained 5 °C below the developmental temperature were monitored for thirty days to record any sign of hatching. They were later placed at the optimum developmental temperature to determine percent mortality and whether living eggs would develop normally.

Population modelling

DYMEX™ V3 (Maywald et al. 2007), a commercially available insect phenology modelling computer programme based on insect critical thresholds and developmental rates as a function of temperature and weather data, was used. We used DYMEX to develop a population model for *T. peregrinus* based on our laboratory-determined adult threshold temperatures (this study). The DYMEX model developed for this research used threshold temperatures reported in this study for both adult and egg development, in addition to using data from Noack and Rose (2007) to calculate the developmental rate of the five nymphal life stages of *T. peregrinus*. Modelling was undertaken using daily temperature and rainfall data from the Pretoria site only, because data from this site were the most complete.

Population monitoring

To determine an optimal sample protocol for monitoring *T. peregrinus* populations, a pilot trial was initiated to determine the optimum position of trap placement and the best colour traps to capture adults (data not shown). Yellow sticky board traps placed at mid-canopy were found to be most effective in capturing *T. peregrinus* adults, confirmed by results from similar studies (Martínez et al. 2010). Six sites ranging in elevation from 47 to 1,340 m above sea level (m a.s.l.) were placed in a stratified design across the main eucalypt growing areas of South Africa (Fig. 1; Table 1) and monitored between February 2007 and July 2009. The trial was established in collaboration with

several commercial forestry companies participating in the Tree Protection Co-operative Programme (TPCP) (www.fabinet.up.ac.za/tpcp). The trapping sites included one in the Gauteng province (Pretoria), one in the KwaZulu-Natal province (Zululand), two sites in the Limpopo province (Tzaneen) and two sites in the Mpumalanga province (Sabie and Piet Retief) (Fig. 1; Table 1).

At each of the six monitoring sites, we placed 20 plastic yellow board sticky traps (13 cm × 8 cm each; Insect Science (Pty) Ltd., Tzaneen, South Africa) on randomly selected trees at a height of 3 m, stratified by distance to the northern edge. For each site, ten traps were placed at the north-facing compartment edge, where initial infestation symptoms are usually first noticed (unpublished observation), and ten traps inside the compartment >10 m from any edge. Compartments had *Eucalyptus* trees of similar size and age (~5 m tall and 2–3 years old) across monitoring sites (Table 1).

We monitored temperature and humidity every 2 h using DS1923-F5 Thermochron *iButton*® (Dallas Semiconductor/Maxim, Dallas, TX, USA) data loggers placed in the middle of the canopy (~3 m), suspended from a tree located at the centre of each monitoring site. Data loggers and traps were exchanged weekly. Traps were returned to the laboratory, and total numbers of *T. peregrinus* adults were counted separately for each trap. Rainfall data were provided by the South African Weather Service from weather stations closest to the individual monitoring sites and aggregated to fortnightly averages. Insect abundance per trap and *iButton* data (temperature and relative humidity) were likewise averaged fortnightly. Data exploration and visualization (specifically Fig. 4) were performed in R (R Core Team 2012).

Results

Critical thermal limits

Thaumastocoris peregrinus adults were not able to right themselves at temperatures between 32 and 45 °C and between 1 and 6 °C. The mean CT_{max} for *T. peregrinus* adults was 38.6 ± 4.7 °C ($n = 30$), and a CT_{min} was with a mean value of 3.8 ± 1.9 °C ($n = 30$). Individual adults that were exposed to their critical thermal limits were subsequently able to recover normal movement when placed at temperatures ranging between 7 and 31 °C.

Egg developmental rate

Thaumastocoris peregrinus egg development was highest at a constant temperature of 30 °C (3 days with 15 % mortality) and lowest (15 days) at a constant temperature

Fig. 1 Map of South Africa showing placement of *T. peregrinus* monitoring sites within Pretoria and the major commercial *Eucalyptus* growing regions of Tzaneen, Sabie, Piet Retief and Zululand

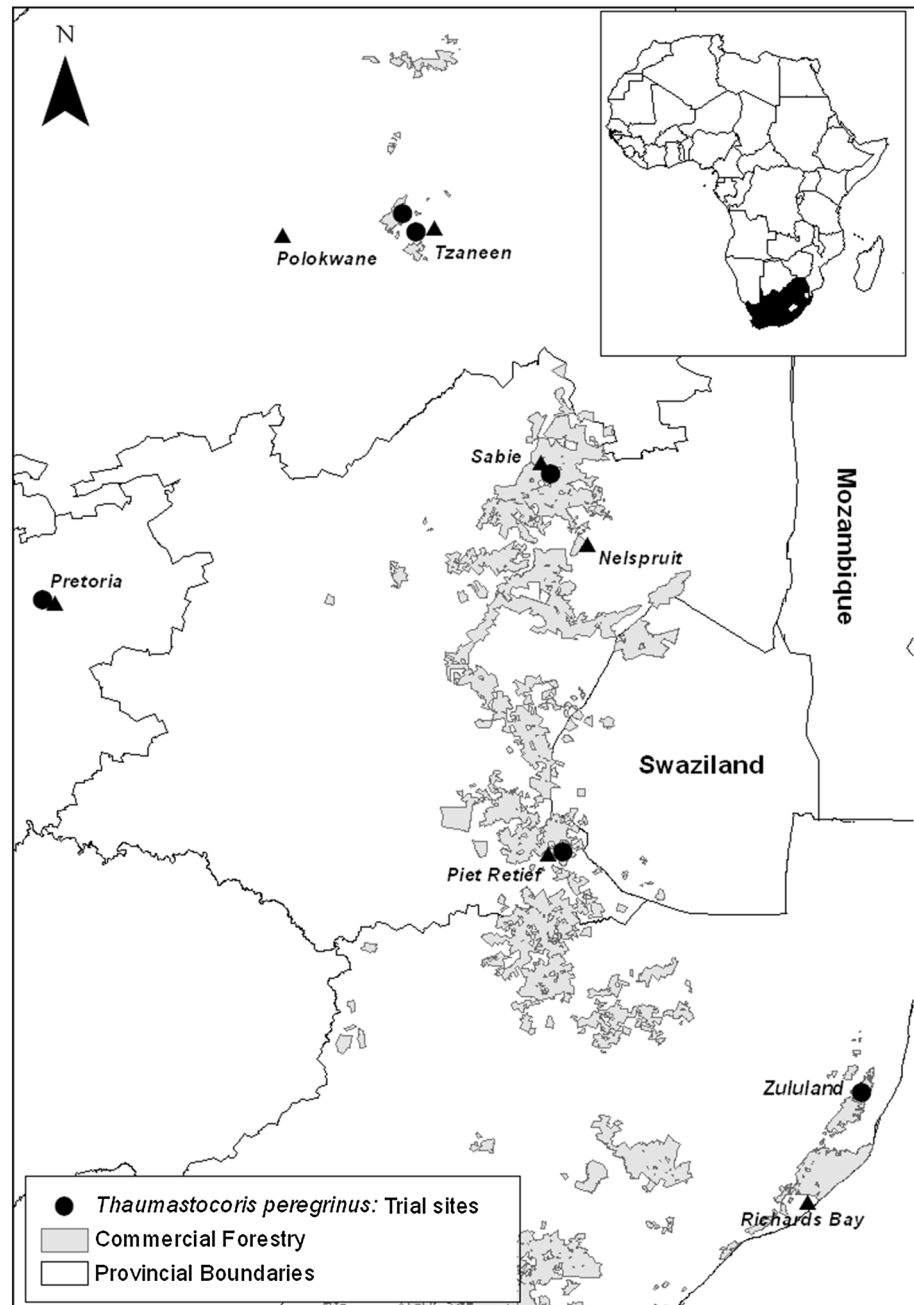


Table 1 Site characteristics for six *T. peregrinus* monitoring sites in South Africa

Site characteristics	Monitoring site					
	Piet Retief	Pretoria	Sabie	Tzaneen, Site 1	Tzaneen, Site 2	Zululand
Elevation (m a.s.l.)	1,219	1,340	1,280	1,276	870	47
Mean annual rainfall (mm)	841	711	1,142	1,253	1,235	917
Mean maximum temperature (°C)	23.2	24.9	22.1	23.6	25.5	27.1
Mean minimum temperature (°C)	10.1	11	10.8	13	12.5	16.6
<i>Eucalyptus</i> spp.	<i>E. macarthurii</i>	<i>E. camaldulensis</i>	<i>E. grandis</i>	<i>E. grandis</i>	<i>E. grandis</i> × <i>E. camaldulensis</i>	<i>E. grandis</i> × <i>E. camaldulensis</i>

of 15 °C. No egg development occurred at 0, 5, 10 and 35 °C (Fig. 2), and these data were excluded from the linear regression analysis, as undertaken by Campbell et al. (1974). The developmental zero of the eggs was calculated at 12.2 °C using the regression equation $y = 0.0178x - 0.2178$, $R^2 = 0.9731$, $p = 0.0135$.

DYMEX model

The DYMEX-simulated population dynamics of *T. peregrinus* predicted adult populations as peaking from October through to April at the Pretoria site and decreasing between the months May to July, with the opposite relationship occurring for the egg numbers at the site (Fig. 3). Predicted nymph numbers remained relatively constant throughout

the duration of the simulated model (Fig. 3). Compared to the actual adult population data obtained for this site (Fig. 4), discrepancies were revealed in the extent and timing of population build-ups and declines between the simulated and actual populations. At the Pretoria site, the adult population increased from December to January, peaking in March to April and decreasing during May (Fig. 4), whereas the deterministic model populations had relatively constant population levels during the same time period.

Monitoring trial

The monitoring trial revealed unique patterns of *T. peregrinus* population growth and decline for each of the populations monitored (Figs. 1, 4). Population abundance was strongly seasonal with a single peak per year at all sites. Trap catches varied dramatically between sampling time points, ranging from zero individuals to a maximum of approximately 60 individuals per trap at peak abundance at the Pretoria site. Despite long periods of low abundance, adults were present year round at all sites. Seasonality was consistent across the two years of sampling within sites (and between the two Tzaneen sites) but was highly divergent across sites. Populations peaked in Pretoria in autumn (February–April; Fig. 4), slightly earlier than at the warmer Zululand site (late summer, January–March). The three northernmost sites (Sabie and Tzaneen sites 1 & 2) peaked in spring (August–October), whereas Piet Retief had only one moderate peak in the spring of 2007. Surprisingly, there was little evidence of a consistent pattern with temperature or relative humidity between sites based on climate records (Fig. 4).

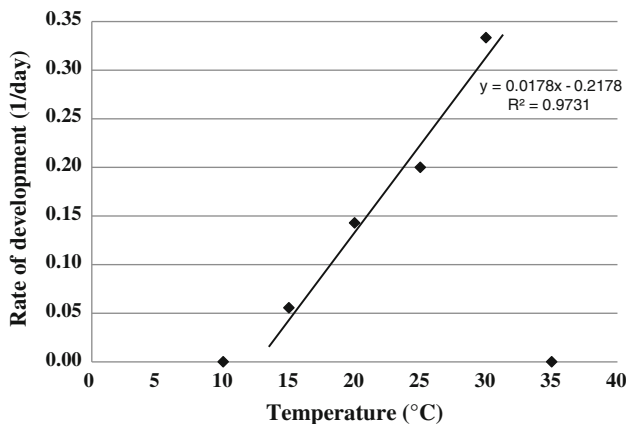


Fig. 2 The relationship between *T. peregrinus* egg development and temperature to determine the developmental rate equation ($y = 0.0178x - 0.2178$, $R^2 = 0.973$, $p = 0.01$) from hatching eggs

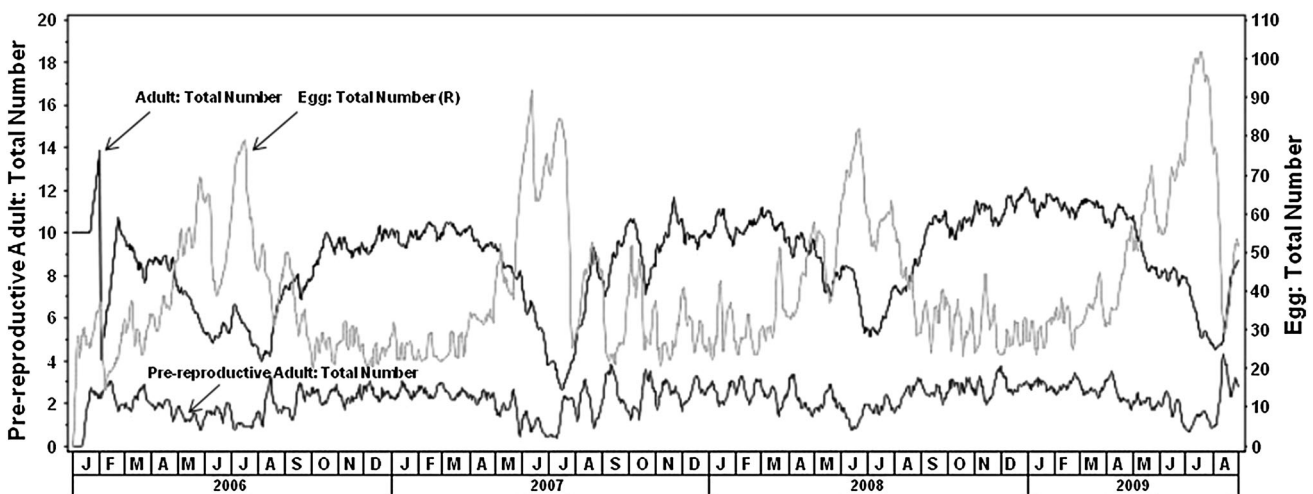


Fig. 3 An output of a DYMEX model simulation, predicting the population dynamics of the various life stages of *T. peregrinus* at the Pretoria monitoring over time

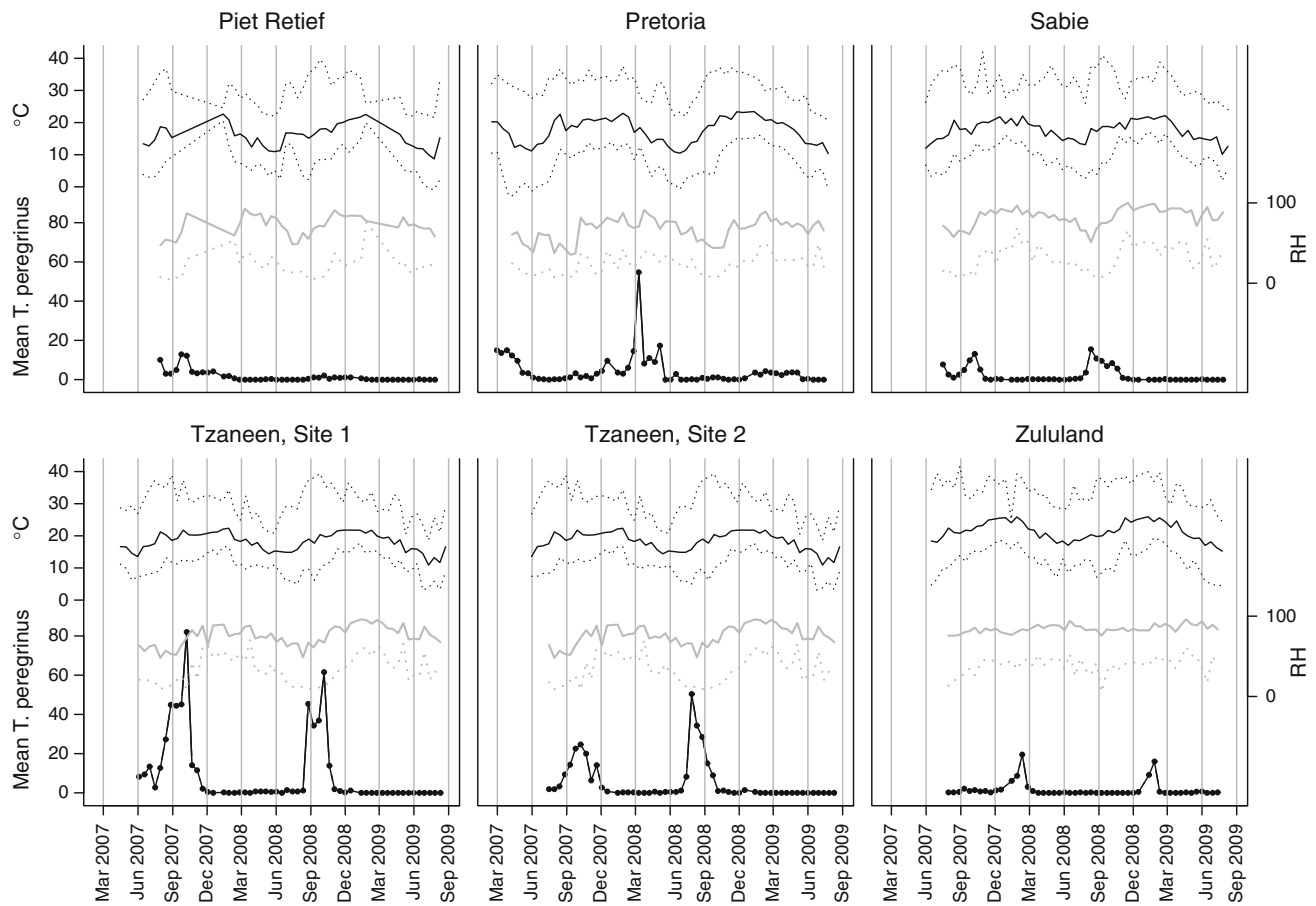


Fig. 4 Monthly temperature ($^{\circ}\text{C}$, left axes, top), relative humidity (RH, right axes) and the number of *T. peregrinus* adults collected per trap for each of the monitoring sites Piet Retief, Pretoria, Sabie, Tzaneen site 1, Tzaneen site 2 and Zululand

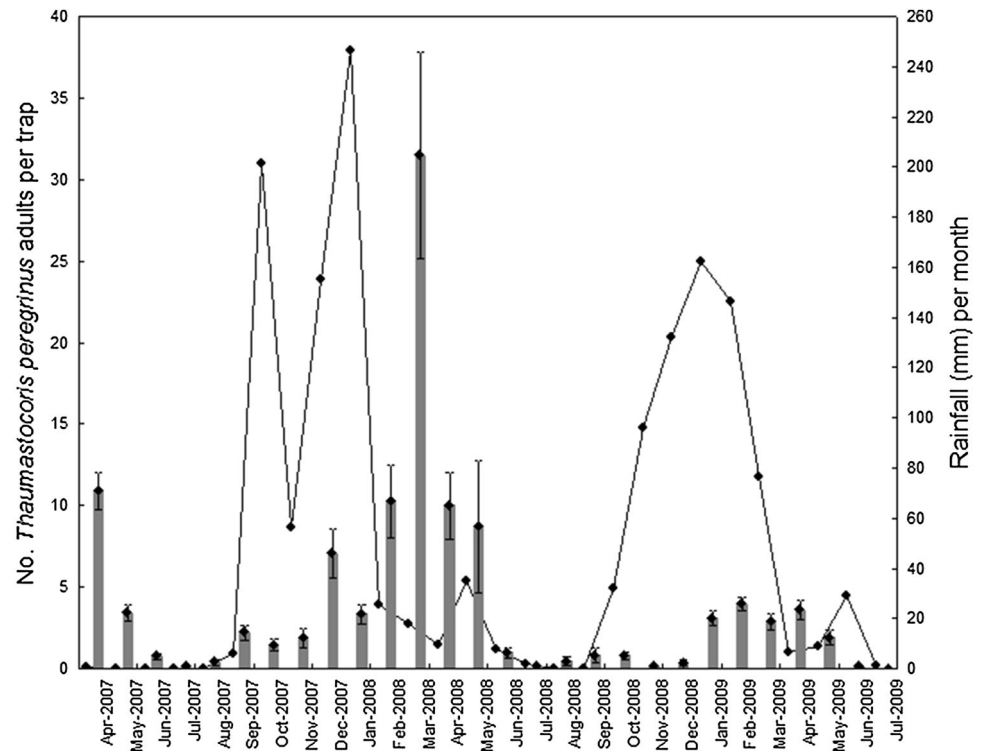
Rainfall data for the Pretoria monitoring site (the only site to have a weather station in reasonably close proximity) revealed that rainfall at this site peaked in January, when adult population levels were still relatively low (Fig. 5). Adult population levels were found to be highest during the 2007/2008 season (year with the higher rainfall) compared to 2008/2009 season (Fig. 5).

Discussion

Monitoring *T. peregrinus* populations for 30 months across the main *Eucalyptus* growing regions of South Africa revealed a regular pattern of population build-up and decline. The fluctuations in populations at each monitoring site were remarkably consistent between years or seasons, but were unique for the different regions. Contrary to expectation, there were no consistent correlations between the presence of *T. peregrinus* and temperature, relative humidity or rainfall across the sites. The relative synchronicity amongst sites in broadly similar regions suggests

that these patterns cannot simply be random and the scale of apparent co-variation would seem to indicate a driver acting at a broad spatial scale, such as weather (Moran 1953). Liebhold et al. (2012) recently reviewed the direct impact of weather on survival and reproduction (termed the “Moran effect” (Moran 1953)), reviewing dispersal, regional stochasticity and trophic interactions as factors affecting spatial synchronicity (Liebhold et al. 2012). One plausible hypothesis for this study is that weather and trophic interaction effects on tree phenology (e.g. leaf flush and senescence) drive *T. peregrinus* seasonality more strongly than a direct effect of temperature. It has been shown that mated *T. peregrinus* females demonstrated a preference to volatile organic compounds emitted by undamaged *Eucalyptus benthamii* plants compared to those damaged by *T. peregrinus* (Martins and Zarbin 2013). Thus, the combined effect of climate factors on the actual plant growth of undamaged trees may indirectly regulate the population dynamics of *T. peregrinus*, rather than a direct effect per se. Such an indirect effect might then limit the utility of evaluating the direct effects of weather

Fig. 5 Monthly rainfall (indicated by the *line plot*) and number of *T. peregrinus* adults collected per trap (indicated by the *bar plot*, vertical bars indicate 95 % confidence intervals) at the Pretoria monitoring site



(climatic factors), especially using relatively short time series. Our data were not sufficient to test for such indirect effects, but this may be a fruitful avenue for future research.

The phenological model for *T. peregrinus* employing laboratory-derived temperature parameters was roughly plausible for the Pretoria site on which it was based. Comparisons of actual adult *T. peregrinus* monitoring data with simulation data revealed faster population build-ups and longer periods of high population levels compared to that of actual populations. The peaks from the model data also did not correlate with the actual data in Pretoria, where the most weather data were available. Further refinement of the model using more accurate nymphal development thresholds and incorporating mortality factors may improve matching with field data. Important factors for future study and potential input to the model include the effect of temperature on adult fecundity, temperature thresholds for each nymphal life stage and bottom-up host effects, amongst others (Stiling 1988; Umbanhowar and Hastings 2002; Abbott and Dwyer 2007; Soliman et al. 2012).

Thaumastocoris peregrinus does not appear to undergo any diapause period and was active during all seasons at all monitoring sites, although sometimes in very low numbers. Diapause periods have likewise not been reported for any of the Thaumastocorinae. While *T. peregrinus* was present at all sites throughout the years, there were consistent

annual seasonal peaks in the numbers of individuals, albeit to different total numbers caught. The consistent and repetitive annual fluctuations of populations at each monitoring site further suggest a strong local environmental effect on the population dynamics of this pest. Whether these influences are direct or indirect is, however, not clear. The consistent pattern of population build-ups and decline also suggests that the population dynamics might be predicted at a particular site for a particular *Eucalyptus* sp. using a model parameterised by data of previous years. Such models would facilitate more efficient monitoring and control including the potential use of semiochemicals (e.g. use of male aggregation pheromones) (González et al. 2012; Martins and Zarbin 2013), insecticides (Noack et al. 2009) and deployment and/or augmentative release of potential biological control agents such as *Cleruchoidea noackae* Lin & Huber (Lin et al. 2007; Mascarin et al. 2012; Nadel et al. 2012) that rely on the presence of large numbers of eggs in field populations.

Observational data from countrywide surveys (Nadel et al. 2010) suggested that rainfall caused *T. peregrinus* populations to decrease at certain monitoring sites. The data from this study, however, did not support this hypothesis at the Pretoria site (the only site with reliable rainfall data that could be used for this purpose). Rainfall during spring usually results in a flush of new leaves and shoots that are high in nitrogen, often leading to increases in numbers and feeding of other insects (Wolda 1978;

Chilima and Leather 2001). *T. peregrinus* feeds on mature *Eucalyptus* leaves and possibly derives no benefit from this increased flush of shoots that usually occurs after first spring rainfalls. The apparent correlation between rainfall and a decrease in populations that has previously been observed (Nadel et al. 2010, authors unpublished) is most likely due to other factors (e.g. such as changes in plant tissue structure or chemistry, pathogen pressure, etc.) (Stiling 1988; Abbott and Dwyer 2007). Population fluctuations after rainfall periods could also be due to changes in other climatic factors, such as humidity and temperature that affect insects and plant-host growth (Specht and Brouwer 1975; Sutcliffe 1977; Jones 2001; Umbanhowar and Hastings 2002; Abbott and Dwyer 2007).

The positive linear relationship between *T. peregrinus* egg development and temperature from 12 to 30 °C is likely to be an important part in the explanation of the influence of average temperature and population size observed at all monitoring sites. The optimal developmental rate of 3 days with 15 % mortality for *T. peregrinus* eggs was at 30 °C. This rate of egg development was much higher in this study compared to other Thaumastocoridae such as *Baclozygum depressum* Bergroth, which had a developmental rate of 8 days for eggs at a temperature of 30 °C (Hill 1988). The rate of egg development in this study at 20 °C (7 days and 10 % mortality) was similar to the developmental rate of *T. safordi* Noack, Cassis & Rose eggs that were found to hatch within 8 days when kept at room temperature (Noack and Rose 2007; Noack 2009; Nadel and Noack 2012). However, these results differ from the developmental rates found for *T. peregrinus* (6 days) and *B. depressum* (15 days) at room temperature (Hill 1988; Noack and Rose 2007; Noack 2009). Egg developmental rates of insects have been found to decrease with further increases above optimum temperature (Kouskolekas and Decker 1966; Beck 1983; Son and Lewis 2005), as shown with *T. peregrinus* eggs that had 100 % mortality and appeared to dry out at 35 °C after 5 days.

Mean temperature provides an indication of the thermal environment to which insects are exposed and, conceivably, influence their developmental rate and behaviour (Beck 1983; Klueken et al. 2009). One would thus expect to encounter an increasing abundance of *T. peregrinus* with increasing mean temperatures due to the linear relationship with that of mean temperature and egg developmental rates from 12 to 30 °C. This relationship was not found at any of the monitoring sites, suggesting that in addition to mean temperature other environmental factors are likely to also be influencing *T. peregrinus* populations. Two possible other factors suggested by the data analysis were maximum temperature and humidity.

Maximum temperatures might have played a role in fluctuating populations at some of the sites. Observational

data from only the Tzaneen and Sabie monitoring sites, however, indicated that increases in mean temperatures above 20 °C did not necessarily result in increased numbers of *T. peregrinus*. This contradictory result was shown in the data to likely be due to large increases in the maximum (25–40 °C) and not that of mean (17–22 °C) temperatures experienced at these sites. This could result in a decrease in population size during hot summer months when thermal tolerance and thermal limits are met (Crozier 2004; Gaston 2009). For *T. peregrinus*, a maximum temperature of above 34 °C may result in egg development ceasing and in turn adult insects reaching their thermal tolerance. Maximum temperatures of over 34 °C that were consistently (23 days over a 122 day period) experienced at the Tzaneen and Sabie monitoring trial sites during the hot summer and early autumn months, sometimes for up to 6 h a day, could possibly explain the decrease in the number of *T. peregrinus* individuals encountered during the hot summer months.

High levels of relative humidity have been shown for other insects to positively affect the hatching of eggs and the survival of adults and immature life stages (Byrne et al. 2002; Duyck et al. 2006; Moriyama and Numata 2006; Simmons et al. 2008). However, high humidity at the Zululand, Tzaneen and Sabie monitoring sites was associated with a decrease in *T. peregrinus* population size. The reason for this is not clear. One possibility is that increases in humidity coupled with increasing temperature may allow for the growth of fungal pathogens on the eggs of *T. peregrinus* (Mascarin et al. 2012). This phenomenon has previously been observed on *T. peregrinus* eggs kept at high humidity levels on leaves under laboratory conditions (authors unpublished), but requires further testing. Fungal egg pathogens are not uncommon and have been effectively used to control numerous insect pest populations (Ferron 1978; Samuels et al. 2002).

Results of this study showed that *T. peregrinus* populations must be separately monitored and in the different regions of South Africa if the data are to be used to accurately predict seasonal abundance peaks. Seasonal synchronicity was quite consistent within sites across the two seasons monitored, potentially allowing for population models to be built that will allow for realistic population predictions and planning in future for specific *Eucalyptus* spp. in each region. While no clear or consistent link emerged between climate and population fluctuations, more work is needed to understand the strong seasonal patterns in abundance. There are also evidently other factors that affect population fluctuations, as would be expected. Another factor that appears to need specific attention includes the influence of variations in resource quantity, nutritional quality and volatile organic compounds of the various *Eucalyptus* spp. (Soliman et al. 2012;

Martins and Zarbin 2013) and how this impacts *T. peregrinus* population dynamics. Such data might be useful to improve our ability to predict future distribution and the severity of outbreaks internationally in regions that are not yet affected by this pest.

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