Synergistic effects of seasonal rainfall, parasites and demography on fluctuations in springbok body condition

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Summary

1. Seasonality of rainfall can exert a strong influence on animal condition and on host–parasite interactions. The body condition of ruminants fluctuates seasonally in response to changes in energy requirements, foraging patterns and resource availability, and seasonal variation in parasite infections may further alter ruminant body condition.

2. This study disentangles the effects of rainfall and gastrointestinal parasite infections on springbok (Antidorcas marsupialis) body condition and determines how these factors vary among demographic groups.

3. Using data from four years and three study areas, we investigated (i) the influence of rainfall variation, demographic factors and parasite interactions on parasite prevalence or infection intensity, (ii) whether parasitism or rainfall is a more important predictor of springbok body condition and (iii) how parasitism and condition vary among study areas along a rainfall gradient.

4. We found that increased parasite intensity is associated with reduced body condition only for adult females. For all other demographic groups, body condition was significantly related to prior rainfall and not to parasitism. Rainfall lagged by two months had a positive effect on body condition.

5. Adult females showed evidence of a ‘periparturient rise’ in parasite intensity and had higher parasite intensity and lower body condition than adult males after parturition and during early lactation. After juveniles were weaned, adult females had lower parasite intensity than adult males. Sex differences in parasitism and condition may be due to differences between adult females and males in the seasonal timing of reproductive effort and its effects on host immunity, as well as documented sex differences in vulnerability to predation.

6. Our results highlight that parasites and the environment can synergistically affect host populations, but that these interactions might be masked by their interwoven relationships, their differential impacts on demographic groups, and the different time-scales at which they operate.

Key-words: Bovidae, Eimeria, endoparasites, Etosha National Park, Namibia, Strongylida, Strongyloides

Introduction

Estimates of animal body condition are used to determine the influence of factors such as environmental degradation, life-history parameters and ecological interactions on animal health (Stevenson & Woods 2006). For ruminants, individual body condition varies cyclically with seasonal changes in energy and protein requirements (e.g. towards maintenance, growth, reproduction and lactation) and resource quality and availability (Parker, Barboza & Gillingham 2009). In balancing the body’s metabolic requirements against seasonal changes in the environment and the resource base, ruminants seasonally adjust their voluntary intake rate (Weber & Thompson 1998), digestive capacity, diet composition and foraging time (Owen-Smith 1994), all factors which contribute to the seasonal fluctuations in body condition.

Animal body condition can also be influenced by parasite infections; in wildlife populations, numerous examples exist...
of associations between gastrointestinal parasite infections and decreased host condition (e.g. Holmstad, Hudson & Skorping 2005; Lello, Buag & Hudson 2005; Newey et al. 2005; Hakkarainen et al. 2007; Craig et al. 2008). In hosts, gastrointestinal parasite infections can cause reduced voluntary intake rates, altered digestive function, altered protein metabolism (Fox 1997) and losses of endogenous protein (Van Houtert & Sykes 1996), all symptoms that would reduce the ability of infected individuals to maintain or increase body condition. Individuals may lose condition as a direct result of disease symptoms and in addition, animals in poorer condition may have a reduced ability to control parasite infections (Beldomenico et al. 2008) or may alter their foraging behaviour in ways that increases exposure to parasites (Hutchings et al. 1999). Whatever the ultimate cause, a parasite-associated reduction in host condition may result in decreased survival of parasitized individuals (Gulland 1992; Hudson, Newborn & Dobson 1992; Murray, Cary & Keith 1997).

For wildlife species with many natural predators, reductions in body condition as a result of parasite infection may increase vulnerability to predation (Murray, Cary & Keith 1997). Our study species springbok (Antidorcas marsupialis) is a medium-sized antelope with many natural predators including lion (Panthera leo) (Stander 1992), spotted hyena (Crocuta crocuta) (Trinkel 2010), black-backed jackal (Canis mesomelas) (Klare et al. 2010), cheetah (Acinonyx jubatus), leopard (P. pardus) (Hayward & Kerley 2008) and numerous other smaller mammalian and avian predators (Skinner & Louw 1996). Across springbok populations, the adult sex ratio is skewed towards females, a pattern which is more pronounced in populations with larger predators (Bednekoff & Ritter 1997). In our study area of Etosha National Park, Namibia, a full suite of medium- and large-bodied carnivores is extant, and estimates of the springbok adult sex ratio vary from 1:5:1 to 2:4:1 females to males, implying that predators are more likely to kill male than female springbok (Bednekoff & Ritter 1997). The goal of this study is to determine whether gastrointestinal parasites or rainfall has a larger influence on springbok body condition. However, distinguishing between environmental and parasite factors driving variation in body condition is complicated because there are many interrelated parasite and environmental factors that can affect body condition in opposing directions. Rainfall seasonality directly influences resource availability and parasitism by altering the development, survival and transmission of parasite life stages in the environment (Fayer 1980; Banks et al. 1990; O’Connor, Walkden-Brown & Kahn 2006). How individuals apportion their nutrient intake among bodily functions (e.g. maintenance, growth, reproduction, immunity) depends on resource availability and their age and reproductive status (Coop & Kyriazakis 1999). Furthermore, for a prey species, the condition observed in the population may be altered by predation, if predators selectively remove individuals in poorer condition or those of particular demographic groups.

In a prior study, we showed that gastrointestinal parasitism of springbok is strongly seasonal and that the prevalence or intensity of parasite infections can relate to host age (Turner & Getz 2010). Here, we build upon these patterns to disentangle the interrelated rainfall, parasite and demographic factors associated with changes in springbok physical condition. We test a series of models to investigate (i) the influence of rainfall, demographic factors and parasite interactions on parasite prevalence and infection intensity, (ii) whether parasitism or rainfall is a more important predictor of springbok body condition and how this varies among demographic groups, and (iii) how measures of parasitism and host condition vary across study areas situated along a rainfall gradient. We discuss how predation effects may or may not contribute to the patterns observed in this study.

Materials and methods

STUDY AREA

This research was conducted in Etosha National Park, a 22 915 km² reserve in northern Namibia, located between 18°30'–19°30'S and 14°15'–17°10'E. Etosha is a semi-arid savanna system with a seasonal rainfall pattern, mainly falling November–April, with the greatest monthly rainfall occurring January–February (Engert 1997). The main study area is located at Okaukuejo in the centre of Etosha (Fig. 1). There is a gradient of increasing rainfall from west to east across Etosha, and two additional study areas (Otjovasandu and Namutoni) were selected at either end of this gradient (Fig. 1). These three areas differ in annual rainfall, soil type (Bengler-Bell & Buch 1997) and vegetation communities (le Roux et al. 1988). The mean annual rainfall from 1971 to 2008 was 301 mm at Otjovasandu, 349 mm at Okaukuejo and 444 mm at Namutoni. During 2006–2007 when all three areas were sampled, the rainfall differences between areas were larger than apparent in the annual means (Fig. 1).

STUDY SPECIES

Springbok are an arid-adapted antelope (tribe Antilopini) occurring in drier regions of southern Africa (Nagy & Knight 1994) and a commercially important game species. Springbok segregate sexually and have a territorial mating system and a six-month gestation period. They are categorized as selective, mixed-feeding herbivores (Hoffmann, Knight & Skinner 1995). Springbok are the most abundant larger herbivore in Etosha, with a population estimate of 15 600 (13 200–17 900 95% CI) in 2005 (cf. Ministry of the Environment and Tourism aerial survey records). The parasites assessed in this study include nematodes in the superfamilies Trichostrongyloidea (order Strongylida, hereafter strongyles) and the genus Strongyloides (order Rhabditida) and two coccidian morphotypes in the genus Eimeria (hereafter Eimeria A and Eimeria B; Fig. 2). A morphtype may represent an undescribed species or a complex of multiple species; further research is underway to determine species units. We did not attempt to differentiate the strongyles based on egg morphology, and many strongyle genera parasitize springbok: Agriostomum, Cooperia, Cooperoides, Dictyocaulus, Haemonchus, Impalaiia, Longstrongylus, Nematodirus, Oesophagostomum, Ostertagia, Paracooperia and Trichostrongylus (Round 1968; Horak, Meltzer & de Vos 1982; De Villiers, Liversidge & Reinenecke 1985).
SAMPLE COLLECTION AND PARASITOLOGICAL ANALYSIS

Gastrointestinal parasitism was evaluated noninvasively using quantitative estimates of egg and oocyst shedding in springbok faeces from opportunistically selected individuals. Faecal specimens were collected from our main study area in Okaukuejo between July 2005 and April 2008, for a total of 718 samples. Collecting periods were July–August 2005, February–October 2006, February–June 2007 and January–April 2008. Monthly sample sizes ranged from 27 to 48 individuals, with a mean of 37 samples per month. Samples were collected from the other two study areas in April–May 2006 (wet season), July–August 2006 (dry season) and March–April 2007 (wet season). Thirty faecal samples were collected per area per month of sampling, collecting a total of 160 samples from each area.

In all study areas, samples were collected between 7:00 and 13:00, and areas were sampled in rotation: Okaukuejo was always sampled in the first week of the month, Namutoni in the second week and Otjovasandu in the third week. The laboratory for parasite analysis was based in Okaukuejo, so when working in other study areas, samples were collected over 2-5 days, stored in a refrigerator on the vehicle and then processed immediately on return to the laboratory. Samples were processed at most three days after collection and often at shorter time intervals.

In each sampling period (per month and area), we sampled at different waterholes and roads each day, to reduce the potential to resample individuals (elaborated in Turner & Getz 2010). On observing an individual defecate, its age, sex and body condition were recorded and the faeces collected within 10 min of deposition. Age and sex were determined via horn growth and morphology and genitalia (Raufenbach 1971). Age was assessed in three categories: juveniles <1 year, yearlings 1–2 years and adults 2+ years old. We adapted the condition scoring system developed by Berry & Louw (1982) for wildebeest to springbok. The condition categories (1 = very poor, 2 = poor, 3 = fair, 4 = good, 5 = excellent) were based on the shape of the hindquarters and the visibility of the ribs and pelvis.

Faecal samples were evaluated for gastrointestinal parasite eggs or oocysts using a modification of the McMaster method for faecal egg counts (Gibbons et al. 2005; detailed in Turner & Getz 2010). In rare cases where the number of oocysts on the slide was too high to count accurately, the sample was diluted and the counts adjusted in proportion to the dilution factor. The McMaster technique is generally used to quantify an estimate of the number of eggs per gram of faeces (the egg count seen in the two slide chambers multiplied by 50). We were...
interested in the relative differences in intensity, not estimates of parasites per gram of faeces, and multiplying the count data by 50 creates large discontinuities in the data set. Therefore, all intensity results were evaluated and reported based on slide counts rather than faecal egg counts.

DATA ANALYSIS

Two data sets were used to assess the objectives, the larger data set from the main study area with monthly level resolution (\(N = 718\) samples collected during 20 months over a 34 month period) and a seasonal-level data set with samples collected from the three study areas (in two wet and one dry seasons). For the analyses among areas (\(N = 524\) samples), only the subset of data from the main study area that matched the sampling period for the other study areas was used. The main data set was used to assess how parasitism and body condition were affected by host demography, rainfall and infection with other parasites. The study area data set was then used to determine whether parasitism and body condition differed in areas with higher and lower annual rainfall than the main study area. Statistical analyses were conducted in R 2.11.0 (R Development Core Team 2010).

Parasite prevalence and intensity

Statistical analyses of factors affecting parasite presence were conducted using logistic regressions, and analyses of factors affecting parasite intensity were performed using generalized linear models (GLMs) with a negative binomial error distribution and log link function. Parasite intensity is an estimate of parasite quantity shed by infected individuals; therefore, when parasite intensity was the dependent variable in an analysis, only samples for which that parasite was present (i.e. count > 0) were included. However, when intensity was an independent variable, all data for that parasite type, including zero values, were used in the analysis.

The prevalence or intensity of each parasite type was evaluated for relationships to host age and sex, monthly rainfall and associations with other parasites (as prevalence or intensity, to match the parasite measure of the dependent variable). The rainfall variables examined were monthly rainfall one and two months prior to sample collection; rainfall in the month of collection was not included because samples were collected in the first week of each month. Only proximate rainfall effects were examined as these are most relevant to the time-scale required to complete parasite life cycles.

Springbok body condition

Although condition scores were recorded using a five-category scale, 94% of the individuals sampled had condition scores of good or fair. As a result, the condition scores were collapsed into two classes – lower condition (poor or fair condition scores) and higher condition (good or excellent condition scores) – and analysed using logistic regressions.

As some parasite infections are more prevalent or intense in younger animals and others in adult animals (Turner & Getz 2010) and animals have different energetic requirements depending on their age and sex, the factors exerting a strong influence over an individual’s body condition will change over time. As a first level of analysis, we tested whether there were significant differences in body condition among age and sex classes. As a result of this analysis, the data were partitioned into juvenile, yearling, adult female and adult male data sets. This allowed for an assessment of the rainfall and parasite factors affecting springbok condition independent from demographic factors and also a comparison of how the factors affecting body condition differ for different ages or by sex. Each of these analyses contrasted the effects of rainfall (one and two months prior) and the intensity of each parasite type on body condition.

Study area comparisons

The study area data set was used to compare how environmental factors – including annual rainfall variation (study area), rainfall years (wet or dry sampling year) and season (wet or dry) – affected the measures of parasitism and body condition. Age was included as a factor in these analyses to control for age-related variation in parasitism or condition. Statistical tests for parasite prevalence, intensity and body condition were the same as used for the main data set.

Springbok movement patterns provide evidence that animals can migrate seasonally along the edge of the Etosha pan (Panagis & Stander 1989; W.C. Turner unpublished data); therefore, springbok populations in the intermediate-rainfall and wetter study areas may not be fully independent. There is no evidence supporting springbok movement corridors between the drier area and the other two study areas. Given the potential nonindependence of the wetter two study areas, we focus on the comparison between the drier and wetter area, which are separated by c. 250 km.

Results

PARASITE PREVALENCE AND INTENSITY

Strongyles, Strongyloides and Eimeria A all had positive associations between monthly rainfall and egg or oocyst shedding (Table 1; Fig. 3). Strongyle prevalence significantly increased with increased rainfall one and two months prior; however, strongyle intensity was only significantly related to rainfall one month prior. The prevalence and intensity of Eimeria A and Strongyloides were significantly related to rainfall one and two months prior. The prevalence of Eimeria B was not significantly related to either rainfall variable (Table 1).

There were significant positive associations in prevalence among strongyles, Strongyloides and Eimeria A and between the intensities of strongyles and Strongyloides (Table 1). There were no significant relationships between Eimeria B and the prevalence of other parasites.

The prevalence and intensity of strongyles significantly increased as host age increased (Table 1). Strongyles were also more prevalent in males than females but strongyle intensity in infected individuals was higher in females than males (Table 1). The prevalence estimates of strongyle nematodes were as follows: juvenile females 74%, juvenile males 66%, yearling females 83%, yearling males 91%, adult females 79% and adult males 87%. In contrast to strongyles, Eimeria spp. parasites had negative relationships with age. Eimeria B prevalence significantly decreased as age increased and although Eimeria A prevalence was not significantly related to host age, the intensity Eimeria A infections significantly decreased as age increased (Table 1). There were no statistically significant relationships between parasite prevalence or intensity and host sex or age for Strongyloides.
Prevalence and intensity of springbok parasites

Independent variables

<table>
<thead>
<tr>
<th>Parasite data types</th>
<th>Age</th>
<th>Sex</th>
<th>Rain 1</th>
<th>Rain 2</th>
<th>Strongyles</th>
<th>Strongyloides</th>
<th>Eimeria A</th>
<th>Eimeria B</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>β(SE)</td>
<td>P</td>
<td>β(SE)</td>
<td>P</td>
<td>β(SE)</td>
<td>β(SE)</td>
<td>β(SE)</td>
<td>β(SE)</td>
</tr>
<tr>
<td>Strongyles</td>
<td>0.338 (0.135)</td>
<td>* 0.503 (0.227)</td>
<td>* 0.013 (0.004)</td>
<td>** 0.015 (0.005)</td>
<td>** 1.092 (0.296)</td>
<td>*** 0.591 (0.281)</td>
<td>* 0.295 (0.332)</td>
<td>ns 718</td>
</tr>
<tr>
<td>Prevalence</td>
<td>0.287 (0.059)</td>
<td>*** 0.345 (0.086)</td>
<td>*** 0.006 (0.009)</td>
<td>*** 0.001 (0.001)</td>
<td>ns 0.007 (0.002)</td>
<td>*** 0.000005 (0.00009)</td>
<td>ns 0.00006 (0.0003)</td>
<td>584</td>
</tr>
<tr>
<td>Strongyloides</td>
<td>-0.066 (0.137)</td>
<td>ns 0.036 (0.214)</td>
<td>ns 0.014 (0.003)</td>
<td>*** 0.019 (0.003)</td>
<td>*** 1.67 (0.285)</td>
<td>*** 1.220 (0.235)</td>
<td>*** -0.259 (0.306)</td>
<td>ns 718</td>
</tr>
<tr>
<td>Prevalence</td>
<td>-0.105 (0.069)</td>
<td>ns 0.057 (0.101)</td>
<td>ns 0.008 (0.001)</td>
<td>*** 0.003 (0.001)</td>
<td>*** 0.006 (0.002)</td>
<td>** -0.002 (0.001)</td>
<td>* -0.0007 (0.0003)</td>
<td>* 480</td>
</tr>
<tr>
<td>Eimeria A</td>
<td>0.071 (0.133)</td>
<td>ns 0.063 (0.210)</td>
<td>ns 0.018 (0.003)</td>
<td>*** 0.015 (0.003)</td>
<td>*** 0.581 (0.280)</td>
<td>* 1.203 (0.239)</td>
<td>*** -0.463 (0.3)</td>
<td>ns 718</td>
</tr>
<tr>
<td>Intensity</td>
<td>-0.508 (0.097)</td>
<td>*** -0.699 (0.145)</td>
<td>ns 0.007 (0.001)</td>
<td>*** 0.010 (0.001)</td>
<td>*** 0.002 (0.003)</td>
<td>ns -0.002 (0.003)</td>
<td>ns -0.0006 (0.0005)</td>
<td>ns 438</td>
</tr>
<tr>
<td>Eimeria B</td>
<td>-0.745 (0.122)</td>
<td>*** -0.604 (0.210)</td>
<td>ns 0.002 (0.003)</td>
<td>ns 0.0003 (0.002)</td>
<td>ns 0.403 (0.324)</td>
<td>ns -0.090 (0.289)</td>
<td>ns -0.286 (0.283)</td>
<td>ns 718</td>
</tr>
</tbody>
</table>

Relationships are shown among parasite prevalence or intensity and monthly rainfall, host demographics and parasite interactions. Prevalence was tested with logistic regression; infection intensity with GLM. The variables rain 1 and rain 2 are respectively the rainfall one and two months prior to sample collection; samples were collected in the first week of each month. A positive coefficient for sex indicates the parasite measure was higher in males than females. Associations in Eimeria B intensity were excluded because of low prevalence (16.7%; 120 positives per 718 samples). Significance levels are *** P ≤ 0.001, ** P ≤ 0.01, * P ≤ 0.05, ^ P ≤ 0.1; ns = nonsignificant.

Table 2. Rainfall vs. parasite effects on springbok body condition

<table>
<thead>
<tr>
<th>Independent variables</th>
<th>Age/sex categories</th>
<th>Rain one month prior</th>
<th>Rain two months prior</th>
<th>Strongyles</th>
<th>Strongyloides</th>
<th>Eimeria A</th>
<th>Eimeria B</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>β(SE)</td>
<td>P</td>
<td>β(SE)</td>
<td>P</td>
<td>β(SE)</td>
<td>β(SE)</td>
<td>β(SE)</td>
</tr>
<tr>
<td>Juveniles</td>
<td>-0.018 (0.008)</td>
<td>*</td>
<td>0.015 (0.008)</td>
<td>*</td>
<td>0.003 (0.024)</td>
<td>ns</td>
<td>-0.018 (0.016)</td>
</tr>
<tr>
<td>Yearlings</td>
<td>-0.004 (0.004)</td>
<td>ns</td>
<td>0.011 (0.004)</td>
<td>**</td>
<td>0.003 (0.011)</td>
<td>ns</td>
<td>-0.018 (0.014)</td>
</tr>
<tr>
<td>Adult males</td>
<td>-0.0001 (0.032)</td>
<td>ns</td>
<td>0.007 (0.003)</td>
<td>*</td>
<td>-0.007 (0.010)</td>
<td>ns</td>
<td>-0.0007 (0.0080)</td>
</tr>
<tr>
<td>Adult females</td>
<td>-0.0007 (0.004)</td>
<td>ns</td>
<td>0.004 (0.003)</td>
<td>ns</td>
<td>-0.024 (0.008)</td>
<td>**</td>
<td>-0.002 (0.012)</td>
</tr>
</tbody>
</table>

Analyses were partitioned into demographic groups because of age and sex variation in body condition (Fig. 4). Significance levels are *** P ≤ 0.001, ** P ≤ 0.01, * P ≤ 0.05, ^ P ≤ 0.1; ns = nonsignificant.
Springbok were significantly more likely to be in the lower body condition class as age increased, and there was a near-significant interaction of age × sex on springbok condition (logistic regression: age, \( z = -6.64, P < 0.0001 \); sex, \( z = -1.49, P = 0.137 \); age × sex, \( z = 1.88, P = 0.061 \); Fig. 4). The percentage by age of animals in the lower-condition class was 14% juveniles, 42% yearlings and 56% adults (51% of adult males and 62% of adult females were in the lower condition class). As a result of these demographic patterns in springbok body condition, the effect of rainfall vs. parasite factors on condition were assessed separately for juveniles, yearlings, adult males and adult females.

The relative importance of rainfall vs. parasite factors on springbok body condition differed for the age and sex classes examined. The body condition of juveniles, yearlings and adult males was significantly related to rainfall whereas the condition of adult females was significantly related to strongyle intensity (Table 2). Although not always statistically significant, the directionality of relationships between condition and the rainfall variables was consistent among all groups examined: rainfall one month prior had a negative relationship with condition and rainfall two months prior had a positive relationship (Table 2). Juvenile springbok had a significant relationship between rainfall one month prior and body condition and a near-significant relationship between condition and rainfall two months prior. Yearlings and adult males had a significant relationship between condition and rainfall two months prior but not one month prior. Adult female condition was significantly lower for individuals with higher strongyle intensities (Fig. 5a; Table 2). For comparison, we repeated the model of adult female body condition excluding parasites from the analysis. When considering only rainfall variables, adult female condition was significantly and negatively related to rainfall one month prior (\( \beta = -0.009 \ SE = 0.003, z = 2.713, P = 0.007 \)). None of the other parasites examined had statistically significant relationships with springbok body condition.

**Fig. 3.** Monthly variation in rainfall and parasite prevalence in springbok.

**SPRINGBOK BODY CONDITION**

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**Fig. 4.** Springbok body condition by age and sex. Condition score is the average (1–5 scale) condition score recorded; means are presented ± standard errors.
The differences between adult males and females in strongyle parasitism and body condition are presented in Fig. 6, to explore why adult males and females differed in the factors affecting body condition. Data are presented monthly to show how parasitism and condition change seasonally and how the differences between the sexes change in relation to the generalized timing of important life-history events (i.e. parturition, lactation and breeding). Adult females have higher strongyle intensity and lower condition than adult males during the wet season, when females give birth and are lactating. In the dry season, after juveniles have been weaned, adult females have lower strongyle intensity than adult males and there are no detectable differences between the sexes in body condition. After the mating season, adult males experience a slight decrease in body condition compared to adult females.

**PARASITISM AND HOST CONDITION AMONG STUDY AREAS**

Parasitism increased from one study area to the next along the rainfall gradient only for *Strongyloides* prevalence and intensity (Table 3). For the other parasites, there was either no significant relationship between parasitism and the study areas (e.g. the prevalence of strongyles, *Eimeria* A and *Eimeria* B) or the significant relationships in parasitism among areas did not directly follow the rainfall gradient. Strongyle intensity was significantly lower in the intermediate-rainfall area than the wetter area but did not differ significantly between the wetter and drier areas. *Eimeria* A intensity was significantly higher in the wetter than the drier area, but the intermediate-rainfall area had higher *Eimeria* A intensity than the wetter area.

Despite the large interannual variation in rainfall between the two years of study, only *Eimeria* A (intensity) and *Strongyloides* (prevalence) showed significant differences between the two years, with parasitism higher in the wetter than drier year (Table 3). All parasite estimates were significantly higher in wet than dry seasons, with the exception of *Eimeria* B. *Eimeria* B was not statistically related to any of the environmental variables examined, although it had a near-significant negative relationship to year.

**Table 3. Environmental factors affecting springbok parasitism and body condition across study areas**

<table>
<thead>
<tr>
<th>Data type</th>
<th>Season (dry vs. wet)</th>
<th>Year (drier vs. wetter)</th>
<th>Area (drier vs. wetter)</th>
<th>Area (main vs. wetter)</th>
<th>Age</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>β (SE)</td>
<td>P</td>
<td>β (SE)</td>
<td>P</td>
<td>β (SE)</td>
</tr>
<tr>
<td>Strongyle prevalence</td>
<td>3.320 (0.481)</td>
<td>***</td>
<td>0.298 (0.356)</td>
<td>ns</td>
<td>0.003 (0.336)</td>
</tr>
<tr>
<td>Strongyle intensity</td>
<td>2.201 (0.132)</td>
<td>***</td>
<td>0.149 (0.095)</td>
<td>ns</td>
<td>0.267 (0.104)</td>
</tr>
<tr>
<td><em>Strongyloides</em> prevalence</td>
<td>4.075 (0.365)</td>
<td>***</td>
<td>0.628 (0.313)</td>
<td>*</td>
<td>0.729 (0.327)</td>
</tr>
<tr>
<td><em>Strongyloides</em> intensity</td>
<td>2.882 (0.255)</td>
<td>***</td>
<td>0.035 (0.115)</td>
<td>ns</td>
<td>0.672 (0.129)</td>
</tr>
<tr>
<td><em>Eimeria</em> A prevalence</td>
<td>4.142 (0.364)</td>
<td>***</td>
<td>0.232 (0.360)</td>
<td>ns</td>
<td>0.039 (0.336)</td>
</tr>
<tr>
<td><em>Eimeria</em> A intensity</td>
<td>3.171 (0.326)</td>
<td>***</td>
<td>0.609 (0.168)</td>
<td>***</td>
<td>0.047 (0.132)</td>
</tr>
<tr>
<td><em>Eimeria</em> B prevalence</td>
<td>0.398 (0.365)</td>
<td>ns</td>
<td>0.669 (0.364)</td>
<td>***</td>
<td>0.167 (0.247)</td>
</tr>
<tr>
<td>Body condition</td>
<td>0.610 (0.251)</td>
<td>*</td>
<td>1.449 (0.248)</td>
<td>***</td>
<td>0.016 (0.231)</td>
</tr>
</tbody>
</table>

Models included factors for the year (drier or wetter) and the season (dry or wet) and area (drier vs. wetter; intermediate (main) vs. wetter). A positive coefficient (β) for the categorical variables indicates the parasite or condition measure was higher in the wetter of the two categories considered. Significance levels are ***P ≤ 0.001, **P ≤ 0.01, *P ≤ 0.05, †P ≤ 0.1; ns = nonsignificant. Age is added when warranted (see Table 1) to account for age-related variation in parasitism or condition.
Springbok were in better condition in the wetter than drier years and in the wet than dry seasons (Table 3). Contrary to expectation, springbok in the drier area were significantly more often in the higher body condition class than were springbok in the other two areas (there was no significant difference in body condition between the intermediate and wetter areas). The sex differences for adults in the relationship between strongyloide intensity and body condition (Fig. 5a for the main study area) were consistent in each of the additional study areas (Fig. 5b).

Discussion

The goal of this study was to disentangle the effects of rainfall and parasitism on seasonal fluctuations in springbok body condition. Springbok body condition, and its relationship to rainfall or parasite factors, varied among demographic groups. The effects of rainfall on springbok body condition show both positive and negative relationships and direct and indirect influences on body condition, depending on the demographic group. Increased parasite intensity was associated with reduced body condition only for adult females. In all other demographic groups examined, body condition was related to rainfall and not parasite variables. Host resilience (i.e. productivity despite parasite infection) and resistance (i.e. ability to limit parasite establishment, growth, fecundity or persistence) to parasitism are affected by the balance between nutrition and the particular energy demands of the individual (Coop & Kyriazakis 1999) – factors that vary with age, sex and seasonality. This study demonstrates the importance of environmental variation and demographic influences on rates of parasitism and ultimately on animal body condition.

Rainfall vs Parasite Effects on Body Condition

The relative importance of rainfall or parasite factors in describing patterns of springbok body condition varied among demographic groups (Table 2). For yearlings and adult males, rainfall positively influenced body condition, and condition improved significantly with rainfall at a two-month lag. A lag between peak rainfall and improvement in body condition is expected and ruminants are generally in the month lag. A lag between peak rainfall and improvement in body condition improved significantly with rainfall at a two-month lag. A lag between peak rainfall and improvement in body condition, depending on the demographic group. Increased parasite intensity was associated with reduced body condition only for adult females. In all other demographic groups examined, body condition was related to rainfall and not parasite variables. Host resilience (i.e. productivity despite parasite infection) and resistance (i.e. ability to limit parasite establishment, growth, fecundity or persistence) to parasitism are affected by the balance between nutrition and the particular energy demands of the individual (Coop & Kyriazakis 1999) – factors that vary with age, sex and seasonality. This study demonstrates the importance of environmental variation and demographic influences on rates of parasitism and ultimately on animal body condition.

Parasitism and Condition Along the Rainfall Gradient

Against expectations, springbok were in significantly better body condition in the drier than the wetter study area (Table 3). There was a decreased presence and intensity of Strongyloides and a decreased intensity of Eimeria A in the drier area, but no significant differences were detected between the two areas in the prevalence or intensity of strongyles, the only parasite which affected body condition in the main study area (for adult females).

Beyond rainfall, these areas also differ in vegetation structure and composition and in the diversity and density of herbivorous mammals, all factors that could influence body condition and parasitism. Springbok are an arid-adapted, mixed-feeding species, and individuals graze when grasses are green and highly digestible (Bigalke & van Hensbergen 2007). A significant number of the grass species are pathogenic in livestock, whereas Strongyloides spp. infection is more often symptomatic (Bowman 2003). For Strongyloides spp. in springbok, this generalization is supported; there were no significant relationships between Strongyloides spp. and body condition, despite the significant associations between this parasite and infection with both strongyles and Eimeria A. Infections with the Eimeria morphotypes were more prevalent or intense in juvenile springbok, while infection with strongyles was more prevalent and intense in adults. Given the age-related patterns in parasite infections, the Eimeria spp. are the parasites most likely to affect juvenile springbok condition. The lack of a negative relationship between the Eimeria morphotypes and juvenile condition indicates that these parasites are not particularly pathogenic in free-ranging springbok.

area, 2.8 lions per 100 km² [Berry 1981; estimates for the wetter area are not available].

Rainfall seasonality was an important factor affecting parasitism, whether evaluated in terms of a lagged monthly rainfall effect (Table 1; Fig. 3) or as a categorical season variable in the study area comparison (Table 3). Of the environmental variables considered in the study area comparison, season had a larger effect on parasitism than did year or area effects. Despite the large interannual variation in rainfall between the two years of study, only *Eimeria* A (intensity) and *Strongyloides* (prevalence) showed significant differences between the two years, with parasitism higher in the wetter than drier year. *Eimeria* B showed no relationship with any of the environmental variables considered. This parasite was relatively rare, with a prevalence of 13.4–16.7% in the two data sets.

**SEX DIFFERENCES IN PARASITISM AND CONDITION**

Sex differences in parasitism are commonly observed and are attributed to ecological differences between males and females in parasite exposure or physiological differences in susceptibility (Zuk & McKean 1996). In this study, strongyle nematodes were more prevalent in males than females, and for infected springbok, strongyle intensity was higher in females than males (Table 1). The sex difference in prevalence implies a difference between males and females in exposure to strongyle larvae in the environment, a result that could be attributed to springbok social organization or mating system. Ruminants that sexually segregate often have differences in habitat selection, diet selection or activity patterns between the sexes (Bowyer 2004; Ruckstuhl 2007) – all factors that could lead to sex differences in parasitism.

Springbok males segregate from females, and springbok social groups include mixed-sex groups of females and young, male bachelor groups and lone territorial males (Skinner & Chimimba 2005). A territorial mating system may influence the exposure of males to parasites; Ezenwa (2004) found strongyle infection intensity was higher in territorial males than bachelor males or females for Grant’s gazelles (*Gazella granti*) but not for Thomson’s gazelles (*G. thomsoni*). Adult males were not apportioned into group types in this study, and it is possible that there are differences in parasite exposure for males on territories vs. in bachelor herds. However, only adult males may become territorial (Skinner & Chimimba 2005) and strongyle prevalence was higher for males than females in the yearling and adult age classes. Male springbok segregate from females into bachelor herds before adulthood (Skinner & Chimimba 2005); therefore, the observed sex differences in strongyle prevalence are more likely related to ecological differences resulting from sexual segregation than from territoriality.

The sex difference in strongyle intensity, where infected females had higher intensity than infected males (Table 1), may result from differences in immune responses to strongyle infection. Parasite egg intensity in faeces provides an estimate of a host’s ability to resist an infection given that the parasite is present and reproducing. Host immune responses against adult nematodes include constraints to worm body size and fecundity (Rowe *et al.* 2008) or worm expulsion (Balic, Bowles & Meeusen 2000). In many species, females show signs of a rise in faecal egg counts following parturition, a phenomenon called the ‘periparturient rise’ or ‘periparturient relaxation of immunity’. This increase in parasitism for reproductive females is linked to a decrease in immune function and is rapidly corrected when lactation ceases (Beasley, Kahn & Windon 2010). Late pregnancy and lactation are energetically costly for females: energy requirements for gestating females are 17–32% higher than non-gestation females, and lactation requires an additional 2–3 times more energy than gestation (Robbins 1993). The periparturient rise is thought to result from trade-offs in partitioning a finite nutrient pool between maintenance, reproductive activities and immune function (Coop & Kyriazakis 1999). Experimental reductions in energy demands (i.e. removing nursing young; Houdijk *et al.* 2006) or increases in the forage nutrient supply (i.e. dietary crude protein; Jones *et al.* 2011) can lead to improvements in the expression of immunity in females.

Adult female springbok had higher strongyle intensity and were on average in poorer condition than adult males from the peak in parturition and for the first two months of lactation (Fig. 6). After juveniles are weaned, adult females had lower strongyle intensity compared to adult males. We did not distinguish females by reproductive status, and the actual differences between lactating females and adult males in parasitism and condition may be more dramatic than those seen in our results. In addition, the seasonal timing of reproductive effort for males and females is different. The most energetically costly time for female ungulates often occurs just prior to peak rainfall, allowing females access to high quality or abundant forage when nutrient requirements are at a maximum (Sinclair, Mduma & Arcese 2000). However, this means females likely have increased reproductive stress and a dip in immunity in the season when potential contact with parasite larvae is high. These patterns may explain why changes in body condition were significantly related to parasites for adult females but not for the other demographic groups (Table 2). The springbok mating season occurs in the dry season, and any reproductive-related increases in stress or decreases in immunity for males therefore occurs when there should be little to no contact with parasite larvae in the environment. Our results support that adult females are more negatively influenced by strongyles than are other demographic groups, which may be due to increased energetic demands of gestation and lactation for females and the seasonal differences in reproductive effort between the sexes.

Adult males were more often in better condition than were adult females (Fig. 4). Adult females may be more negatively influenced by strongyles than are adult males, as discussed earlier, causing females on average to be in poorer condition than males. However, as springbok are an important prey species for many predators, the potential role of selective predation on the observed sex differences in condition and strongyle parasitism cannot be ignored. Springbok are the
primary food source for lions in Etosha, and adult males are killed significantly more often than adult females compared to the population sex ratio (Stander 1992). Springbok are also the primary food source for spotted hyena (Trinkel 2010), and hyenas may hunt adult males more than adult females (Martina Trinkel, pers. comm.). Sex differences in springbok vulnerability to predation may be due to their social organization, which is very similar to that of Thomson’s gazelles. Adult male Thomson’s gazelles are hunted more than adult females because males are more likely to be alone, in smaller groups or on the periphery of groups and are less vigilant than females (Fitzgibbon 1990).

A low overall hunting success rate implies that predators may be more successful hunting individuals in poorer condition (Temple 1987; Wirsing, Steury & Murray 2002; Packer et al. 2003). The success rate of lion hunts on springbok was low in comparison with other prey species, at 13% for springbok vs. 11–52% for other species (Stander 1992). Given that adult males are more vulnerable to predation and that hunting success on springbok is low, this may indicate that lions selectively remove males in poorer condition. If lions selectively remove adult males with low condition and high strongyle intensity from the population, this could create the infection/condition patterns observed in this study (Fig. 5). Females in poorer condition may have some degree of protection from predation by living in groups and if so, low condition-high parasite intensity females may be less likely to be removed from the population than similarly infected adult males. However, as territorial males likely have a higher predation risk than individuals in female or bachelor groups and must be in excellent condition to defend a territory, predator selection of more vulnerable adult males may not necessarily remove highly parasitized low-condition males from the population.

In reality, the two possible causes of the sex differences in strongyle intensity and body condition presented here are not mutually exclusive, and it is very likely that both sex differences in immunity and vulnerability to predation play a role in shaping patterns of parasitism and body condition observed in the springbok population. The sex differences in strongyle intensity and body condition were observed consistently across all the study areas (Fig. 5) providing evidence that these patterns are general to the ecology of springbok and are not caused by the peculiarities of any one study area. Additional research is required to distinguish the relative importance of immunity vs. predation on sex differences in parasitism and body condition, and how these factors may vary temporally or spatially.

Conclusion

Ecological systems are complex because there are typically many interacting factors affecting organisms at different scales. Environmental variability has been recognized as an important factor affecting host stress and immunocompetence (Altizer et al. 2006) and the behaviour and ecology of organisms (Owen-Smith 2002; Hopcraft, Oliff & Sinclair 2010). Parasites can have important effects on the life-history parameters of their hosts, reducing survival (Gulland 1992; Hudson, Newborn & Dobson 1992; Murray, Cary & Keith 1997), fecundity (Stien et al. 2002; Lello, Boag & Hudson 2005) or offspring condition (Hakkarakainen et al. 2007). Parasites and the environment can synergistically affect host populations, but the effects of parasites are often masked by the interwoven relationships among parasites, environmental variability and their differential effects on host demographic classes. This complexity is typified by the seasonal fluctuations in ruminant body condition, where animals must acquire the nutrients necessary for short-term survival but also for future reproductive events in dynamic systems alternating between resource abundance and scarcity. Host resilience and resistance to parasitism depend on the balance between nutritional quality of the resource base, the body’s energetic requirements and the magnitude of the detrimental parasite effects on the host. To understand the ecology of any single species, we must understand how environmental variation influences physiology, life history and the strength of species interactions in the web of consumers that surround the target organism, including both its predators and its parasites (Getz 2011). The time-scales at which species interactions can affect a target population vary substantially from multi-year cycles in the abundance of predators to seasonal cycles in resource abundance to weekly–monthly cycles in the abundance of parasites. Our results highlight the importance of considering the inertial effects of past influences on the current structure and health of a given population.

Acknowledgements

This manuscript benefited from comments from Steve Bellan, Carrie Cizauskas and Holly Ganz. We thank the Namibian Ministry of Environment and Tourism for permission to conduct this research and the staff in the Directorate of Scientific Services and Shayne Köttig at the Etosha Ecological Institute for logistical support and assistance. We thank Martina Küsters, Mathias Borsea, Aimee Boursaw, Emily Kalenius, Birgit Köttig, Gabriel Shatumbu, Nigel Berriman, Johannes Kapner and Seth Guim for assistance with sample collection. This research was supported by a Fulbright fellowship, Andrew and Mary Thompson Roeca Scholarships, Professor Earl Storie Memorial Scholarship, G. Fitzgerald Martin Scholarship, and a grant from the Department of Environmental Science, Policy and Management to WCT, and NIH Grant GM083863 to WMG.

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Received 11 January 2011; accepted 1 July 2011
Handling Editor: John Fryxell