Abstract

Overgrazing is a major driver of habitat degradation, especially in southern Africa. Although grasshoppers are adapted to and benefit from natural disturbances, such as grazing by indigenous game and burning, we do not know how they respond to heavy cattle grazing, and how this response interacts with different fire regimes. We also do not know whether grasshoppers respond principally to these disturbances, to changes in the vegetation layer, or to larger landscape attributes (e.g. elevation). We addressed these questions in the topographically heterogeneous Central Midlands of KwaZulu-Natal Province, South Africa. We compared grasshopper assemblages among sites differing in grazing intensity (light, moderate and heavy), fire regime, rocky outcrops and vegetation structure, and attributes of landscape heterogeneity. The local environment (rocky outcrops, bare ground cover, grass height and total vegetation cover) was more important than landscape attributes for all measures of diversity. Grasshopper species richness was best explained by grazing intensity, with the specific response determined by fire regime. Greatest species richness was consistently recorded in heavily-grazed grassland. Thus, we found no evidence in support of the Intermediate Disturbance Hypothesis. Grasshopper assemblage composition of areas with light grazing was different from those with heavy grazing, but areas with light grazing were similar to those with moderate grazing under all fire regimes. Different suites of grasshopper species were adapted to changes in the local environment, with greatest diversity (Shannon H') associated with elevated levels of bare ground and sparse vegetation cover. The greatest proportion of rare, endemic and sensitive grasshoppers (incl. Lentula minuta, Machaeridia conspersa and Chaschius fastigata) was associated with a greater proportion of vegetation cover. The sensitivity of grasshopper assemblages to fire-grazing interactions, and the habitat requirements of different suites of species necessitates consideration of different types (fire and grazing) as well as levels of disturbances when adjusting management practices. We recommend that conservation of rare, endemic and sensitive grasshoppers should be prioritized, as these are most vulnerable to local extirpation.

Key words

assemblage composition, burning regime, elevation, Grasshopper Conservation Index (GCI), grazing intensity, indicators, landscape heterogeneity, plants, Shannon diversity (H'), species richness, topographic position, vegetation structure

Introduction

Fire and grazing by indigenous large ruminant mammals are natural disturbances in Afromontane grassland, which is one of several consumer-controlled grasslands in the world (Bond et al. 2003, Bond and Keeley 2005). Natural disturbances maintain favorable conditions for species coexistence of stationary taxa, such as plants (Chesson 2000). The exclusion of fire causes grassland plant assemblages to change in composition and become species-poor (Paussas and Ribeiro 2017), especially in an African context (Kirkman et al. 2014). Grazing interacts with fire to change the richness and structure of the vegetation layer (Burkepile et al. 2016, Joubert et al. 2017), which then influences arthropod assemblages (Joern and Laws 2013). Superimposed upon these effects of disturbances and disturbance interactions on biodiversity are large-scale spatial and temporal phenomena, such as landscape fragmentation (Stoner and Joern 2004, Krauss et al. 2010), landscape heterogeneity (Batáry et al. 2007), seasonal changes (Fondeville et al. 2014) and weather cycles (Jonas and Joern 2007). It is necessary to identify drivers with large effects on biodiversity, and to understand how they relate with one another in natural landscapes in order to implement appropriate and effective conservation interventions.

Not all of biodiversity responds similarly to drivers of natural landscapes. Patterns in plant assemblages often show a lag in response to changes in the landscape, but respond quite rapidly to changes in the local environment (Krauss et al. 2010, Joubert et al. 2016a). Herbivorous arthropods respond more frequently and consistently to local changes in the vegetation layer than to changes in the landscape, while predatory arthropods respond more frequently to landscape than to local changes in vegetation structure (Collinge et al. 2003, Stoner and Joern 2004, Torma et al. 2014). Due to the taxonomic challenge and sheer numbers of insects (Cardoso et al. 2011), especially in sub-tropical grasslands, it is important to select indicators to represent biodiversity's response to ecosystem and environmental change (McGeoch 1998, Gerlach et al. 2013).

Grasshoppers are often used as indicators of grassland quality (Gerlach et al. 2013). This is because they are taxonomically well-
known and ecologically sensitive, they respond reliably to changes in their local environment (Bazelet and Samways 2011a) and they mimic the response of other invertebrate groups, e.g. butterflies (Marini et al. 2009, Bazelet and Samways 2012). As primary consumers, grasshoppers show greater response to local attributes than to changes in the landscape (Marini et al. 2007, Bazelet and Samways 2011b), but this may vary (Batáry et al. 2007). Grazing influences grasshoppers directly (e.g. mortality due to trampling or unintentional ingestion) and indirectly via the effect of cattle grazing on vegetation structure and specific plant assemblage (Joubert et al. 2005, Marini et al. 2009, Joubert et al. 2016b). In a global review of arthropod response to large grazing mammals, it was concluded that arthropod diversity only increases in grazed ecosystems if increased heterogeneity of the biotic and abiotic environment outweighs loss of resources and increased mortality (Van Klink et al. 2015).

Afromontane grassland is conserved in formally protected areas as well as Ecological Networks (ENs) among forestry plantations in South Africa (Samways and Pryke 2016). The conservation and management of the heterogeneity at the local and landscape spatial scale is central to the success of grassland ENs (Pryke et al. 2013). Design of ENs should incorporate the typical landscape heterogeneity found in the region (Pryke and Samways 2015), while management should avoid homogenization of grassland habitat by incorporating a patch mosaic burning regime (Bazelet and Samways 2011b, Joubert et al. 2016b) and encouraging grazing by indigenous game (Pryke et al. 2016). However, in ENs where domestic cattle replaced indigenous animals as dominant grazers, it is not clear how grasshoppers respond to different intensities of grazing. We also do not know whether grasshoppers respond primarily to these natural disturbances, to changes in the local biotic environment caused by these disturbances, or landscape heterogeneity.

The aim of this paper is to determine the main drivers of grasshopper assemblage composition, diversity and species richness in Afromontane grasslands. Are grasshoppers influenced mostly by grazing intensity, or phenomena at the local or landscape spatial scale? We hypothesize that grazing intensity and the local environment will have a larger effect than larger scale phenomena, because these small herbivores are sensitive to local changes in microclimatic niches, oviposition sites, and shelter from predators. Secondly, we hypothesize that grasshopper diversity will peak at intermediate levels of disturbance, as observed in the literature (Van Klink et al. 2015). Here, we also wish to identify indicator species of different grazing regimes. Thirdly, we expect different measures of grasshopper diversity to correlate with one another, as they correlate with other taxonomic groups (Bazelet and Samways 2012). Answering these questions will help us decide upon conservation action, specifically where it involves grasshoppers in ENs within transformed landscapes.

Methods

Description of study area.—The study took place in the mid-elevational grasslands (1168–1573 m a.s.l.) east of the Drakensberg mountain range in KwaZulu-Natal Province, South Africa. It is a summer rainfall area, with precipitation mostly in the form of thunderstorms and mist in summer, with mean annual precipitation of ~1120 mm. The topography is variable, and so are the vegetation patterns. Grasslands co-occur with natural wetlands in depressions and indigenous forest patches in steep valleys. Anthropic changes to the disturbance regime.—Fire and grazing are natural disturbances in these landscapes (Bond et al. 2003), but their frequency and intensity have changed greatly in response to change in anthropogenic land uses. Domestic livestock replaced indigenous game as dominant grazers, following the introduction of husbandry practices ~2000 years BP, and the influx of European settlers since the early 19th century (Deacon and Deacon 1999). Concurrently, the intensity of grazing increased (Rowe-Rowe and Scotcher 1986), impacting upon fuel load and spread of fire. Changes in land use from natural grasslands to agricultural crops (e.g. maize) and alien tree plantations further drove changes in the fire regime, as land users adapted fire as a tool for managing these novel landscapes. The current fire regime is more homogeneous than in the past due to legislative and organizational constraints that attempt to balance risks and benefits to commercial enterprises and remaining natural habitat.

Site selection and classifications.—Sites (n = 68) were in a large-scale EN in the Mt Shannon and Good Hope Forestry Estates, as well as in the adjacent Protected Area (PA), iMpendle Nature Reserve (Fig. 1). The variability in topography and disturbance regimes (found among sites) is representative of the variability found in the larger landscape. There were differences in abiotic landscape attributes (topographic position, elevation and aspect), or the local environment (rocky outcrops and vegetation structure).

Fire frequency was classified as either annual burning (AB) or longer fire rotations (LFR). Time since last fire at LFR sites were classified as recently-burned (RB) i.e. burned <12 months prior to sampling vs. unburned (UB) i.e. burned >12 months prior to sampling (Table 1). Grazing intensity at each site was categorized as light (reference sites in the PA), and moderate or heavy in the EN. Classification of sites was based on indicators of historical grazing (dominant grass composition and aerial cover by poisonous forbs - Senecio isatidius or S. retrorsus) and current grazing (grass height, bare ground due to trampling, and occurrence of cattle) (Joubert et al. 2017).

Sampling procedure.—Sites were >400 m apart to allow for independence of sampling. Except for annually-burned sites in narrow (<50 m) corridors, all sites were >30 m from forestry compartment edges in the interior of wider (>150 m) corridors. At each site, we sampled the grasshoppers three times: late spring (November 2012), mid-summer (January 2013), and early autumn (March 2013) with sweep nets. This involved sweeping a net (diameter: 400 mm; mesh size: 2 mm) back and forth in an 180° arc. There was one sweep with each step along four 100 m long transects that were spaced parallel to one another and 5 m apart; thus, 400 sweeps per sampling season and 1200 sweeps per site. Data from the three sampling seasons were pooled for analyses. Nets were emptied after every 25–30 sweeps to prevent escape of agile species. Grasshoppers were frozen, sorted and identified to the lowest possible taxonomic level (Disirh 1965, Johnsen 1984, Johnsen 1991, Cigliano et al. 2017).

For the local environment, we recorded vegetation attributes at each site. Plant assemblage composition outperforms vegetation structure at predicting response of different functional groups of arthropods (Schaffers et al. 2008), including grasshoppers (Kemp et al. 1990). However, vegetation structure and host plant diversity hingens upon the contribution of individual plant species (Joern and Laws 2013), especially in an African context (Gandar 1982). Therefore, using plant species richness and measures of vegetation structure as a proxy for change in the vegetation layer is justified.
Table 1. Description of the grazing and fire regime in each group of sites. Abbreviations for grazing intensity: light in iMpendle Nature Reserve (square symbols), and moderate or heavy in the ecological network. Abbreviations for fire regime: annual burning (AB, solid black symbols), grasslands with longer fire rotation that were recently-burned (RB, solid grey symbols) i.e. burned < 12 months prior to sampling and unburned (UB, open symbols) i.e. burned >12 months prior to sampling.

<table>
<thead>
<tr>
<th>Fire frequency</th>
<th>Time since last fire</th>
<th>Fire abbreviation</th>
<th>Grazing intensity</th>
<th>Sample size (n)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Annual burning</td>
<td>Recently-burned</td>
<td>AB</td>
<td>Light (PA)</td>
<td>8</td>
</tr>
<tr>
<td>Annual burning</td>
<td>Recently-burned</td>
<td>AB</td>
<td>Moderate</td>
<td>8</td>
</tr>
<tr>
<td>Annual burning</td>
<td>Recently-burned</td>
<td>AB</td>
<td>Heavy</td>
<td>8</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Recently-burned</td>
<td>RB</td>
<td>Light (PA)</td>
<td>8</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Recently-burned</td>
<td>RB</td>
<td>Moderate</td>
<td>7</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Recently-burned</td>
<td>RB</td>
<td>Heavy</td>
<td>7</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Unburned</td>
<td>UB</td>
<td>Light (PA)</td>
<td>8</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Unburned</td>
<td>UB</td>
<td>Moderate</td>
<td>7</td>
</tr>
<tr>
<td>Longer fire rotations</td>
<td>Unburned</td>
<td>UB</td>
<td>Heavy</td>
<td>7</td>
</tr>
</tbody>
</table>

At each site (~1000 m²), we recorded vegetation attributes in 24 discontinuous vegetation quadrats (1 m²) and six transects (i.e. six transects × 30 m = 180 measurements) (Joubert et al. 2017). In quadrats, we recorded vegetation cover of all plants (i.e. total vegetation cover), vegetation cover by only grasses (i.e. only grass cover), bare ground cover, rocky outcrop cover, and cumulative plant species richness in vegetation quadrats. The cumulative plant species richness of 24 discontinuous vegetation quadrats was used as a proxy for plant species richness of the whole site (1000 m²) (Güler et al. 2016). Vegetation quadrats were spaced evenly along vegetation transects. Along transects, we recorded vegetation height and basal distance at 1 m intervals. Basal distance serves as a proxy for trampling and erosion potential, especially on steep slopes, and measured as the distance from the bottom of a vertical rod (diameter: 15 mm) to where the nearest plant was rooted. Table 2 summarizes the differences in vegetation structure for each grazing intensity class. Transects were connected end-to-end, with orientation of each transect determined randomly. Averages were calculated for all attributes of the vegetation layer, except for cumulative plant species richness. Lastly, we recorded the following landscape parameters for each site: topographic position (footslope/valley bottom, midslope, and crest/ridge/escarpment), elevation and aspect.

Calculation of the Grasshopper Conservation Index.—The Grasshopper Conservation Index (GCI) estimates conservation value of a site based on occurrence of grasshopper species with specific traits related to extinction risk and sensitivity to habitat change. The standardized GCI site score (GCIn) is the sum of all GCI scores of spe-
Grasshopper species richness, Shannon H' diversity, and GCIn represented changes in plant species richness. Then, we tested for any correlations among different measures of grasshopper diversity using the same method, because we did not want to assume a linear relationship among variables (Hauke and Kossowski 2011). Finally, we used the indicator value (IndVal) method in the labdsv package of R (Dufrenee and Legendre 1997) to identify grasshopper indicators of grazing intensity.

Grasshopper assemblage composition was analyzed in PRIMER 6.0 software. Grasshopper data were standardized, and abundances were square root transformed to reduce the effect of dominant species. Then, a resemblance matrix was compiled based on the Bray-Curtis similarity index. We used canonical analysis of principal coordinates (CAP) to visualize patterns in grasshopper assemblage composition, i.e. how it responds to grazing intensity, vegetation structure and landscape attributes. This ordination method displays sites in a multivariate space based on the calculated similarity indices, i.e. sites grouped closely together are similar, whereas widely dispersed sites are different from one another (Anderson and Willis 2003). Then, we used two statistical tests – DistLM for continuous landscape and local variables, and permutational analyses of variance (PERMANOVA) for grazing intensity (i.e. categorical data) – to determine their effects on grasshopper assemblage composition. All continuous variables were imported as environmental data. Bare ground cover, basal distance and rock cover were log transformed. Continuous environmental variables that best describe grasshopper assemblage composition were identified using DistLM with a stepwise selection procedure and AICc selection criterion. We used PERMANOVA in the same software to test for the main effect of grazing intensity, and then to conduct pairwise comparisons among grazing intensity classes.

**Results**

**Grasshopper species richness, Shannon H' diversity and Grasshopper Conservation Index (GCIn).**—In the first model with all variables, grasshopper species richness was best explained by only grazing intensity (AICc = 342.44; Adjusted R² = 0.535, and LM, F = 10.15, P < 0.001). The greatest number of species was recorded in annually-burned areas with heavy cattle grazing, while the lowest number of species was recorded in unburned grassland with light grazing (Fig. 2). In annually-burned and unburned grassland, grasshopper species richness increased with increasing grazing intensity (light < moderate < heavy). For these fire regimes, we found significant differences between areas with light and heavy grazing (annual burning: light < heavy, t = -4.16, P = 0.003; unburned: light < heavy, t = -3.94, P = 0.006). In contrast, grasshopper species richness of recently-burned areas showed a unimodal response (light...
> moderate < heavy) to increasing grazing intensity. For recently-burned areas, species richness of moderate-grazed areas was significantly less than in heavily-grazed areas (t = -3.46, P = 0.026).

Out of all variables, Shannon H’ diversity was best explained by the local environment (Shannon’s diversity index, AICc = 53.07; Adjusted $R^2 = 0.175$, and LM, $F = 5.66, P = 0.001$), but not grazing intensity. There were significant increases in Shannon H’ diversity, as rocky outcrops ($F = 7.66, P = 0.007$) and bare ground cover ($F = 5.58, P = 0.02$) increased, and a near-significant increase as vegetation cover decreased ($F = 3.74, P = 0.058$).

The standardized GCI score per site (GCIn) was indicative of the proportion of rare, sensitive or range-restricted grasshopper species in the assemblage. Out of all variables, GCIn was best explained by total vegetation cover (AICc = 525.37, Adjusted $R^2 = 0.113$, and LM, $F = 9.57, P = 0.003$). The greatest GCIn score was 7 and recorded in an unburned site with light grazing in the PA. This site had only four grasshopper individuals representing three species (Lentula minuta, Machaeridia conspersa and Qachasia fastigiata), which each had a score of 7.

Relationships among diversity measures and environmental variables.— We found a significant positive correlation between grasshopper species richness and Shannon H’ diversity (Spearman, $Rho = 0.741, P < 0.001$). However, the standardized grasshopper conservation index (GCIn) was not significantly correlated with either grasshopper species richness (Spearman, $Rho = -0.031, P = 0.800$) or Shannon H’ diversity (Spearman, $Rho = -0.055, P = 0.658$). Also, plant species richness was not significantly correlated with grasshopper species richness (Spearman, $Rho = -0.154, P = 0.210$), Shannon H’ diversity (Spearman, $Rho = -0.045, P = 0.720$), or the GCIn (Spearman, $Rho = 0.012, P = 0.921$).

Environmental variables in this study were not independent of one another. There were significant correlations among several attributes of the local environment as well as larger landscape (Table 3). Elevation was significantly correlated with the local environment, i.e. rocky outcrops, bare ground cover and vegetation height (Table 3). Proportion of rocky outcrops was significantly correlated with most variables of the local environment:

Grasshopper assemblage composition.—Sites arranged along a continuum of disturbance intensity, with annually-burned and heavily-grazed sites to the left of the ordination space and unburned sites to the right (Fig. 3). Sites with heavy grazing grouped separately from sites with either light or moderate grazing. The bare ground: total vegetation cover gradient explained horizontal spread of sites along the first axis, while variation in rock and grass cover explained the vertical spread of sites along the second axis (Fig. 3). The two axes explained 15.5% and 11.5% of total variation in the dataset, respectively.

Grasshopper assemblage composition was best explained by the local environment (AICc = 531.02; Adjusted $R^2 = 0.157$; Table 4). Specific variables with a significant effect were total vegetation cover (Pseudo-$F = 2.59, P < 0.001$), grass height (Pseudo-$F = 3.13, P < 0.001$), bare ground cover (Pseudo-$F = 4.33, P < 0.001$), and rock cover (Pseudo-$F = 2.02, P = 0.015$).

Grazing intensity had a significant effect on grasshopper assemblage composition (Pseudo-$F = 2.19, P < 0.001$), with heavily grazed areas differing significantly from lightly grazed areas under...
Fig. 3. Canonical analysis of principal coordinates ordination (CAP) of grasshopper assemblage composition to display patterns in the data. Abbreviations for grazing intensity: light in the protected area (square symbols), and moderate (circular symbols) or heavy (triangular symbols) in the ecological network. Abbreviations for fire regime: annual burning (AB, solid black symbols), grasslands with longer fire rotation that were recently-burned (RB, solid grey symbols) i.e. burned < 12 months prior to sampling and unburned (UB, open symbols) i.e. burned >12 months prior to sampling. Significance values for pairwise comparisons are in Table 4.

### Table 5. Indicator species of grazing intensity, fire frequency, and time since last fire.

<table>
<thead>
<tr>
<th>Species</th>
<th>Disturbance</th>
<th>GCI</th>
<th>Ind Val</th>
<th>P-value</th>
</tr>
</thead>
<tbody>
<tr>
<td>Anablepia pilosa</td>
<td>RB-Light</td>
<td>6</td>
<td>0.74</td>
<td>0.001</td>
</tr>
<tr>
<td>Eyrepocnemis calceata</td>
<td>RB-Light</td>
<td>4</td>
<td>0.21</td>
<td>0.058</td>
</tr>
<tr>
<td>Pseudoaryctera cephalica</td>
<td>RB-Light</td>
<td>6</td>
<td>0.27</td>
<td>0.017</td>
</tr>
<tr>
<td>Dnophersula callosa</td>
<td>AB-Moderate</td>
<td>4</td>
<td>0.27</td>
<td>0.013</td>
</tr>
<tr>
<td>Tetrigrid sp. 3</td>
<td>AB-Moderate</td>
<td>7</td>
<td>0.26</td>
<td>0.031</td>
</tr>
<tr>
<td>Acorypha ferrifer</td>
<td>AB-Heavy</td>
<td>4</td>
<td>0.27</td>
<td>0.025</td>
</tr>
<tr>
<td>Catantops ochthephilus</td>
<td>AB-Heavy</td>
<td>5</td>
<td>0.38</td>
<td>0.003</td>
</tr>
<tr>
<td>Tetrigrid sp. 1</td>
<td>AB-Heavy</td>
<td>5</td>
<td>0.35</td>
<td>0.009</td>
</tr>
<tr>
<td>Coryphosima stenoptera</td>
<td>RB-Heavy</td>
<td>4</td>
<td>0.33</td>
<td>0.015</td>
</tr>
<tr>
<td>Lentula obtusifrons</td>
<td>RB-Heavy</td>
<td>7</td>
<td>0.30</td>
<td>0.065</td>
</tr>
<tr>
<td>Vitticatantops maculatus</td>
<td>RB-Heavy</td>
<td>4</td>
<td>0.28</td>
<td>0.013</td>
</tr>
<tr>
<td>Orthochtha sp. 2</td>
<td>UB-Heavy</td>
<td>3</td>
<td>0.33</td>
<td>0.078</td>
</tr>
<tr>
<td>Spathosternum nigrotaeniatum</td>
<td>UB-Heavy</td>
<td>6</td>
<td>0.49</td>
<td>0.001</td>
</tr>
</tbody>
</table>

We identified 13 species that were indicative of grazing intensity, of which eight species were associated with heavy grazing (Table 5). The GCI scores of two individual indicator species (Lentula obtusifrons and Spathosternum nigrotaeniatum) in heavily grazed areas were high (≥6).

### Discussion

Local versus landscape attributes.—Grasshopper assemblages responded primarily to changes in their local environment and not to larger landscape attributes. This was surprising, because earlier studies found large and significant effects of elevation and aspect on grasshopper assemblages in these mid-to-high elevational grasslands (Samways 1990, Gebeyehu and Samways 2006, Crous et al. 2013, 2014). In Afromontane grassland, grasshopper assemblage composition changed, and species richness increased with an increase in elevation (900–2200 m a.s.l.) (Crous et al. 2013). However, in Swaziland, grasshopper species richness showed the opposite response, as it declined with an increase in elevation (800–1400 m a.s.l.) (Wettstein and Schmid 1999). It is possible that the 400 m range in elevation in our study was not sufficient to detect this major ecological gradient. Alternatively, the effect of elevation might be explained by covariation among local and landscape attributes. There were significant correlations among landscape and local environmental attributes in our study. Our study is not unique. In the Succulent Karoo, there was sparser vegetation cover and greater grasshopper diversity on small hills (Gebeyehu and Samways 2006). Grasshopper assemblages in North America respond to large-scale and long-term environmental gradients (e.g. elevation and precipitation), but these variables are also known to correlate with changes in the local environment (Kemp et al. 1990, Jonas and Joern 2007). This is the case for calcareous and steppe grasslands in Germany (Fartmann et al. 2012, Weiss et al. 2013). Such relationships among environmental variables at the local and landscape spatial scale are a natural part of the landscape, and the reason...
The effect of grazing intensity.—Grazing intensity was the most important determinant of grasshopper species richness in our study. However, the specific response of grasshopper species richness to grazing intensity (light < or > moderate < heavy) depended on fire regime. This is because each fire regime exerts its own selection pressure on the grasshopper species assemblage (Joubert et al. 2016b), especially during the first year after fire (Little et al. 2013). The observed effect of grazing intensity is therefore on a subset of the complete species pool in these mesic grasslands. A case in point is recently-burned grassland where we found a significant response in assemblage composition and species richness when comparing moderately- and heavily-grazed areas. Such differences between moderately- and heavily-grazed areas did not exist in either annually-burned or unburned grassland. This scenario differs from a case where fire frequency and time since last fire had no such effect on grasshopper assemblage composition, causing grasshoppers to respond primarily to grazing and not to a fire-grazing interaction (Joern 2005). Because grasshoppers respond to a fire-grazing interaction in our study area, it is necessary to consider both types and different levels of these disturbances when making management adjustments.

Grasshopper assemblages in heavily-grazed areas were unique in composition and more species-rich than areas with light or moderate grazing. The shift towards a more species-rich grasshopper assemblage illustrates that grasshoppers are relatively tolerant of disturbance. This includes at least one flightless, narrow-range endemic species (Lentula obtusifrons) that was an indicator of heavily-grazed areas. The high degree of tolerance to heavy cattle grazing came as a surprise, although we knew beforehand that grasshoppers are adapted to and benefit from grazing by domestic livestock (Prendini et al. 1996) and indigenous game (Pryke et al. 2016). In North American tallgrass prairies, an increasing level of recent grazing by bison also increased grasshopper species richness (Joern 2005). In a global assessment of arthropod response to grazing, it was concluded that grazing can only increase the richness of grasshopper assemblages if it increases heterogeneity of the local environment, and if this increase in heterogeneity is large enough to make up for the loss of resources and increased mortality (Van Klink et al. 2015). This was expected to occur at moderate levels of grazing, and so lend support to the Intermediate Disturbance Hypothesis (Connell 1978). However, because greatest richness was documented in areas with heavy grazing, our findings did not meet these expectations. Grazing and its interaction with fire indeed increases the heterogeneity of vegetation layers across the landscape, with bare patches interspersed with patches of tall grass and grazing lawns (Archibald et al. 2005). These less disturbed vegetation patches are of great value for grasshoppers in an African savanna, especially in a disturbed mosaic (e.g. around a waterhole) where elevated levels of bare ground leaves insufficient cover for grasshoppers to escape predators and intense heat (Samways and Kreuzinger 2001, Gebyeeyu and Samways 2003). Management for heterogeneity should be prioritized to provide in the habitat requirements of different taxa – disturbance-tolerant species that benefit from heavy grazing, and less disturbed patches for other more sensitive species.

The effect of vegetation structure.—Full vegetation cover indicative of low levels of disturbance was most important for a suite of sensitive, rare and range-restricted grasshopper species that were of great conservation importance. This contrasts with the majority of grasshopper species that were more tolerant of disturbance, as indicated by the rich suite of species associated with elevated levels of bare ground, shorter grass, and sparser vegetation cover. Different grasshopper species are also associated with differences in bare ground cover and grass height in African subtropical grassland (Bazelet and Samways 2011a) and savanna (Prendini et al. 1996). Shorter grass benefitted grasshopper species richness in the Swiss Alps (Marini et al. 2009). Although a unique and rich suite of species were associated with greater levels of disturbance, the conservation of sensitive and range-restricted grasshopper species should be prioritized, as they are most vulnerable to local extirpation, especially when considering the large-scale occurrence of heavy grazing.

The effect of rocky outcrops.—Grasslands with more rocky outcrops supported a different and more diverse grasshopper assemblage than grasslands with less rocky outcrops. This concurs with an earlier study, which found surface rockiness to be a good abiotic indicator of grasshopper species richness in a nearby mesic grassland (Crous et al. 2013). Grasshoppers are very sensitive to changes in their local environment, and may use rocky crevices for shelter to escape large temperature fluctuations (Samways 1990). In the United Kingdom, sensitivity of grasshoppers to microclimate was illustrated by their movement away from prevailing winds relative to the direction of a footpath (Gardiner and Dover 2008). Grasshoppers also avoided excessively warm microhabitats devoid of sufficient vegetation cover to prevent them from overheating (Gardiner and Hassall 2009). In the temperate Cape Floristic Region in South Africa, behavior of small, endemic Betisoides species was influenced by wind intensity, temperature and vegetation height (Matenaar et al. 2014). We argue that rocky outcrops might add to the heterogeneity of microclimatic niches available in the landscape, directly by providing shelter and basking sites, and indirectly by altering the vegetation layer (Crous et al. 2014).

Surrogates of grassland diversity.—Apart from the significant, positive correlation between grasshopper species richness and Shannon H’ diversity, we found no meaningful relationships among measures of plant and grasshopper diversity. The proportion of rare, sensitive or range-restricted grasshoppers (GCln) was not correlated with either grasshopper species richness or Shannon H’ diversity. This contrasts with the findings in another study where small grasshopper species with localized distributions were good indicators of species richness in another arthropod group i.e. butterflies (Bazelet and Samways 2012). Also, we found no relationship among plant and grasshopper species richness. In fact, the greatest number of grasshopper species in this study was documented in annually-burned areas with heavy grazing, while this management practice was absolutely detrimental to indigenous plant conservation (Joubert et al. 2014). Non-congruence between plant and grasshopper species richness concurs with findings of a regional study conducted in Inner Mongolia (Hao et al. 2015). In contrast, there was a significant positive relationship between grasshopper and plant species richness in North American tallgrass prairie (Joern 2005) as well as in the Italian Alps (Marini et al. 2009). The absence of significant relationships among measures of plant and grasshopper diversity emphasizes the need to use multiple taxa and multiple measures of diversity to monitor change in grasslands, particularly since the same level of disturbance can cause gains in one taxonomic group and losses in another.

Management recommendations.—Afromontane grassland management should be cognizant of the individual and interactive effects of grazing and fire, as they each uniquely influence the richness and...
composition of grasshopper assemblages. The majority of species are adapted to high levels of disturbance causing them to persist well in grazed landscapes typical for large parts of the African continent. As such, they do not require special conservation measures to be put in place, provided these grasslands are grazed or burned. However, to also conserve the smaller, more sensitive suite of grasshopper species, patches of minimally disturbed grassland (i.e. areas where grazing or burning is difficult, such as rocky outcrops) should be left. Creating a mosaic of patches with different levels of disturbance is necessary to provide habitat for taxa with diverse requirements.

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