Diets of impala from Kruger National Park: evidence from stable carbon isotopes

M. Sponheimer, C.C. Grant, D.J. de Ruiter, J.A. Lee-Thorp, D.M. Codron and J. Codron


Impala are known to exhibit dietary flexibility, relying primarily on browse in some areas and graze in others. In this study we use stable isotope analysis of faeces and hair to examine the diets of impala in Kruger National Park. As expected, the data show that impala are mixed-feeders and highly distinct from grazing buffalo and browsing kudu. Moreover, impala, buffalo, and kudu faeces contain 2.1%, 1.4%, and 2.9% nitrogen respectively, suggesting that impala diets are of intermediate quality. There are also marked differences between impala populations in the northern and southern regions of the park. The northern impala graze less than their southern counterparts. This difference probably reflects decreased availability of herbaceous forage in the mopane-dominated north. Males and females also have different diets, with males grazing more than females.

Key words: impala, carbon isotopes, faeces, hair, diet.

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Introduction

Impala (*Aepyceros melampus*) are known to exhibit great dietary flexibility. They can make large changes in the percentages of browse and graze consumed seasonally (Monro 1980; Dunham 1982; Smithers 1983; Meissner *et al.* 1996; Wronski 2002). One study suggested that grasses make up 90% of impala diets during the rainy season, but only 33% during the dry season (Meissner *et al.* 1996). Impala diets are also believed to differ markedly between localities reflecting changes in local vegetation. For instance, it has been reported that some Tanzanian impala populations eat over 90% grass (Lamprey 1963), whereas some Zimbabwean populations have diets dominated by browse (Smithers 1983). Here, we investigate the diets of impala from Kruger National Park, South Africa, using stable carbon isotope analysis of faeces and hair. Stable carbon isotopes of hair or faeces have been used to investigate the diets of wildlife (Tieszen & Imbaba 1980; Van der Merwe *et al.* 1988; Witt & Beeton 1997; Wit *et al.* 1998; Schoeninger *et al.* 1999), but such studies are few in number. Furthermore, these materials have never been analysed in tandem.

Stable carbon isotope analysis is particularly useful for determining the percentages of browse and graze in the diets of African herbivores (Vogel 1978; Ambrose & DeNiro 1986). Grasses (graze) in tropical environments use the *C*$_4$ photosynthetic pathway, while trees, shrubs, and forbs (browse) use the *C*$_3$ pathway. This results in these food
types having highly divergent, non-overlapping ratios of $^{13}$C/$^{12}$C (Smith & Epstein 1971). Because the carbon from these food sources is incorporated into faeces and hair in well-understood ways (DeNiro & Epstein 1978; Jones et al. 1981; O’Connell & Hedges 1999), we can then use the $^{13}$C/$^{12}$C ratios of these materials to determine the percentages of graze and browse in an animal’s diet.

Methods

The Kruger National Park is broadly divided into granite-derived soils in the west and basalt-derived soils in the east. The rainfall decreases from south to north and to a lesser extent from west to east (Joubert 1986). Tail hair samples were obtained from darted animals in December, 2001 and faeces were collected in June, 2002. Our collections included animals from the northern and southern regions of the park, to see if the different rainfall regimes led to divergent trophic behavior. In order to provide a comparative baseline, we also gathered tail hair and faeces samples from the grazing cape buffalo (*Syncerus caffer*) and browsing kudu (*Tragelaphus strepsiceros*). The darted animals were from healthy populations that were undergoing routine veterinary study by Kruger National Park staff. Thus, our sampling strategy was perforce opportunistic.

About 6 cm of tail hair was homogenised so that analysis would provide a long-term dietary average. Hair grows at the rate of about 1 cm per month (Saitoh et al. 1969), so the samples we analysed represented about 6 months’ growth. Faecal samples were dried, homogenised and ground. Hair and faecal samples were combusted in an automated Carlo-Erba device (Carlo-Erba, Milan) and stable carbon isotopes were analysed using a flow-through inlet system on a continuous flow isotope ratio mass spectrometer (Finnigan, Bremen). This provided us with the $^{13}$C/$^{12}$C ratios and percentage nitrogen of each sample. The $^{13}$C/$^{12}$C ratios are expressed here as δ values in parts per thousand (‰) relative to the PDB standard. The standard deviation for replicate measurements of a yeast standard was < 0.1 ‰. Data from grazing buffalo and browsing kudu are included to provide a comparative baseline. We used one-way analysis of variance to look for species, region, and sex differences.

The δ$^{13}$C values were converted to % C$_4$ consumed using –12.5 ‰ and –27.0 ‰ as pure grazing and browsing dietary endpoints (Vogel et al. 1978), and assuming isotopic fractionations of –0.9 ‰ and +3.0 ‰ for faeces and hair respectively (Jones et al. 1981; Cerling & Harris 1999; Sponheimer pers. obs.). For example, an animal with hair $\delta^{13}$C of –24.0 ‰ would have a dietary $\delta^{13}$C of –27.0 ‰, and thus be

<table>
<thead>
<tr>
<th>Species</th>
<th>N</th>
<th>$\delta^{13}$C</th>
<th>% N</th>
<th>% C$_4$</th>
</tr>
</thead>
<tbody>
<tr>
<td>Faeces</td>
<td></td>
<td></td>
<td></td>
<td></td>
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<tr>
<td>Impala Combined</td>
<td>39</td>
<td>-20.5±2.8</td>
<td>2.1±0.5</td>
<td>53±20</td>
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<tr>
<td>Impala North</td>
<td>18</td>
<td>-22.2±2.6</td>
<td>2.1±0.7</td>
<td>41±19</td>
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<tr>
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<td>21</td>
<td>-19.2±2.2</td>
<td>2.1±0.4</td>
<td>63±16</td>
</tr>
<tr>
<td>Buffalo</td>
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<td>-14.7±0.5</td>
<td>1.4±0.3</td>
<td>95±3</td>
</tr>
<tr>
<td>Kudu</td>
<td>19</td>
<td>-26.8±0.4</td>
<td>2.9±0.5</td>
<td>8±3</td>
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<tr>
<td>Hair</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Impala Combined</td>
<td>36</td>
<td>-15.2±3.4</td>
<td>na</td>
<td>63±24</td>
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<td>na</td>
<td>44±20</td>
</tr>
<tr>
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<td>-12.6±1.3</td>
<td>na</td>
<td>82±10</td>
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<tr>
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<td>-16.6±3.8</td>
<td>na</td>
<td>53±27</td>
</tr>
<tr>
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<td>na</td>
<td>67±21</td>
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<tr>
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<td>na</td>
<td>98±2</td>
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<tr>
<td>Kudu</td>
<td>19</td>
<td>-23.5±0.5</td>
<td>na</td>
<td>4±3</td>
</tr>
</tbody>
</table>
Results and Discussion

Both faeces and hair data show impala to be mixed-feeders and significantly different from buffalo and kudu (\(P < 0.0001\)) (Table 1; Fig. 1). The faeces data suggest impala have a diet of 53 % \(C_4\) vegetation (grass), whereas the hair data suggest a diet of 63 % grass. These results are consistent with previous studies of impala in Kruger National Park and elsewhere (Van Zyl 1965; Hofmann & Stewart 1973; Monro 1980; Smithers 1983; Van Rooyen 1992; Meissner et al. 1996; Vogel 1978; Wronski 2002). The faecal nitrogen contents of impala, buffalo, and kudu are also significantly different (\(P < 0.0001\)) (Table 1). Befitting a mixed-feeder, impala faecal nitrogen concentrations are higher than those of buffalo, but lower than those of kudu. These findings are also consistent with a multiple year study of ungulate faecal nitrogen concentrations in Kruger National Park (Grant et al. 2000).

Both faeces and hair data also show strong dietary differences between impala from the northern and southern regions of the park (\(P < 0.0001\)) (Table 1; Fig. 2), with southern impala grazing more than their northern counterparts. The faeces data suggest a diet of 41 % grass in the north and 63 % grass in the south. Similarly, the hair data suggest a 44 % grass diet in the north and a 82 % grass diet in the south. The difference between the faeces and hair data in the south may reflect the fact that the hair data represent a long-term dietary average, whereas the faeces data largely reflect the season when they were collected (the beginning of the dry season). Nonetheless, both data sets concur that impala have different diets in the mopane-dominated north than they do in the marula-knobthorn and bushwillow woodlands of the south (Gertenbach 1983). Intriguingly, there is no difference in the faecal nitrogen concentrations of impala in the northern and southern regions of Kruger National Park. Note that both hair and faeces data agree that impala graze more in the south.
Because the hair samples were collected from darted individuals, we were also able to look for sex differences in diet. Throughout the park males grazed 14% more than females (Table 1; Fig. 3); however, this difference was only statistically significant in the southern portion of the park ($P = 0.03$). This provides further evidence for sex based dietary differences as observed by other researchers (Wronski 2002). In future, it might also prove fruitful to explore possible dietary differences associated with disparate social structures. For instance, do males in bachelor herds and those with harems have divergent diets?

**Conclusion**

In this study we were able to show regional and sex based dietary differences of impala in Kruger National Park. Further stable isotope analysis should prove a powerful tool for supplementing traditional studies of wildlife diets. Observational studies are very time-consuming, which necessarily limits the number of populations that can be observed. Furthermore, although these studies provide rich detail about the variety of plants consumed and the time spent feeding on each diurnally, they cannot easily quantify the actual amounts of browse and graze consumed both day and night.

In contrast, stable isotope analysis is very fast and can provide precise information on the amount of browse and graze digested. Once samples have been obtained, preparation and analysis of 100 samples can be completed in a single day. Thus, although stable isotope studies cannot be used to identify individual plant species consumed, they are ideal for quickly comparing the percentages of browse and graze consumed by different groups, whether they are intra- or inter-populational.

Stable isotope analysis also has the advantage that it can be used to study extinct populations. The hair of mounted trophies and even fossilised teeth can be used to glean something of the diets of animals long since dead (e.g., Lee-Thorp & van der Merwe 1987; Macko et al. 1999; Sponheimer et al. 1999). Thus, we can use this tool not only to study the nutritional ecology of modern wildlife, but to study the development of modern dietary adaptations through time.

Moreover, emerging techniques will make it possible to derive even more ecological information from such analyses. For instance, sampling tail hair in small increments (< 1 cm), makes it possible to trace dietary change through time (O’Connell & Hedges 1999). This should prove particularly useful for documenting seasonal changes in browse and graze consumption. It has also been suggested that nitrogen isotopes in hair can be used as indicators of nutritional stress (Hobson et al. 1993). Given present capabilities and the increasing potential offered by constant technical advances in the field, stable isotope analysis should prove a valuable supplement to traditional studies in mammalian dietary ecology in the future.
Acknowledgments

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