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1	Dietary competition between the alien Asian Musk Shrew (Suncus murinus) and a reintroduced
2	population of Telfair's Skink (Leiolopisma telfairii)
3	
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18	sequencing, translocation
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24	
25	Running title: Niche overlap - alien vs. native predators

#### 26 Abstract

27 Reintroduction of rare species to parts of their historical range is becoming increasingly important as 28 a conservation strategy. Telfair's Skinks (Leiolopisma telfairii), once widespread on Mauritius, were 29 until recently found only on Round Island. There it is vulnerable to stochastic events, including the 30 introduction of alien predators that may either prey upon it or compete for food resources. 31 Consequently skinks have been introduced to Ile aux Aigrettes, another small Mauritian island that 32 has been cleared of rats. However, the island has been invaded by Asian Musk Shrews (Suncus 33 murinus), a commensal species spread by man well beyond its natural Asian range. Our aim was to 34 use next generation sequencing to analyse the diets of the shrews and skinks to look for niche 35 competition. DNA was extracted from skink faeces and from the stomach contents of shrews. 36 Application of shrew and skink-specific primers revealed no mutual predation. The DNA was then 37 amplified using general invertebrate primers with tags to identify individual predators, then 38 sequenced by 454 pyrosequencing. 119 prey MOTUs (molecular taxonomic units) were isolated, 39 though none could be identified to species. Seeding of cladograms with known sequences allowed 40 higher taxonomic assignments in some cases. Although most MOTUs were not shared by shrews and 41 skinks, Pianka's niche overlap test showed significant prey overlap, suggesting potentially strong 42 competition where food resources are limited. These results suggest that removal of the shrews from 43 the island should remain a priority.

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50 Introduction

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52 The introduction of locally extinct species to suitable habitats within their wider geographical range is 53 an increasingly important component of conservation strategies (Seddon et al. 2012). When the 54 distribution of a threatened native species has contracted to one or a few isolated sites it is highly 55 vulnerable to stochastic events, such as the introduction of alien species, which could rapidly destroy 56 a last remaining stronghold. Translocation of such a species to a new habitat becomes a conservation 57 priority. The habitat of such an alternative refuge should ideally be free of threats from alien species, 58 providing ecological conditions suitable for reintroductions. However, removal of alien species can 59 often be physically impossible (for example with many invertebrate species) or prohibitively 60 expensive. In some cases the effective techniques for removal of an alien need to be developed. 61 Under such conditions it may be necessary to attempt reintroductions under less than ideal 62 conditions and pragmatically determine whether a rare species can thrive in sympatry with 63 remaining aliens. Examples of successful translocations are birds such as the Kakapo (Strigops 64 habroptilus) between offshore islands in New Zealand (Elliott et al. 2001), and both pink pigeon 65 (Columba mayeri) and Mauritius Fody (Foudia rubra) to Ile aux Aigrettes (Seymour et al. 2005; 66 Cristinnace et al. 2009), and reptiles including whiptail lizards (Cnemidophorus vanzoi) to Praslin 67 Island, Saint Lucia (Dickinson & Fa 2000), Antiguan racers (Alsophis antiguae) to offshore islands of 68 Antigua (Daltry et al. 2001) and lizards to New Zealand islands (Towns & Ferreria 2001).

69

Asian Musk Shrews, *Suncus murinus* (Soricidae), are a highly invasive species spread by man to numerous locations outside its natural Asian range (Ruedi *et al.* 1996). It is a commensal species with man, often living in and around houses and spread by us between land masses. It was introduced to Mauritius in the 18<sup>th</sup> century and has been implicated in the loss of endemic vertebrate and invertebrate species there (Jones 1993; Cole *et al.* 2005; Cheke & Hulme 2008; Solow *et al.* 2008) as well as in other parts of the world, such as Guam (Fritts & Rodda 1998). Between 2009 and 2010, the

76 shrew invaded Flat Island to the north of Mauritius, leading to the localised loss of three endemic 77 reptile species within 18 months (N Cole unpublished data). It is thought to have been introduced to Ile aux Aigrettes (southeast of Mauritius) in the early 20<sup>th</sup> century where it spread rapidly (Cheke & 78 79 Hume 2008). Seymour et al. (2005) calculated that 20 females of S. murinus on Ile aux Aigrettes could 80 potentially generate a population of 550 individuals over a five month reproductive season. On Ile 81 aux Aigrettes, eradication programmes appeared to be successful for a while, but it soon became 82 clear that some individuals had survived and population recovery was rapid (Varnham et al. 2002; 83 Seymour et al. 2005; Solow et al. 2008). Cats (Felis catus) and brown rats (Rattus rattus) were 84 successfully eliminated from Ile aux Aigrettes by 1991 as part of a habitat restoration programme 85 (Jones & Hartley 1995), but this may have simply exacerbated the problem with the alien shrews, 86 releasing them from predation and competition with these equally alien predators.

87

88 Telfair's Skinks (Leiolopisma telfairii) are one of eight species of endemic Mauritian reptiles that 89 managed to survive on Round Island, where they are thriving in the absence of alien predators 90 (North et al. 1994; Pernetta et al. 2005). Historically these skinks lived on mainland Mauritius and on 91 a number of surrounding islands (Cheke & Hume 2008). As an insurance against loss of the Round 92 Island population, the skinks were introduced to Ile aux Aigrettes between 2006 and 2010 where the 93 adults are surviving well, but there is strong evidence that juveniles may be directly preyed on by 94 Asian Musk Shrews. There is also evidence that adult skinks prey upon shrews and annual population 95 surveys of terrestrial vertebrates along transect lines on Ile aux Aigrettes demonstrated a 68% 96 decline in the relative abundance of shrews since skinks were released (N. Cole unpublished data). 97 However, the skinks and shrews may also be limited by resource competition. Evidence from the 98 eradication programme, based upon live trapping, showed that as numbers of shrews declined, their 99 mean body mass increased considerably. This suggested that food resources were limiting and that 100 this increase in mass was the result of release from intraspecific competition (Seymour et al. 2005). It

101 follows that interspecific competition, between shrews and skinks, might also therefore have an 102 adverse effect upon the skinks if they share the same prey. Both shrews and skinks are omnivorous, 103 eating both plant and animal foods, which may buffer them against food shortages during the dry 104 season on Ile aux Aigrettes, when invertebrate prey are scarce (Cole & Harris 2011). Little is known 105 about the invertebrate prey species consumed by shrews and skinks, although morphological 106 identification of fragments of larger prey in faecal samples has provided some information but 107 mainly at higher taxonomic levels (Vinson & Vinson 1969; Pernetta et al. 2005; Richards 2007; Zuël 108 2009; Copsey et al. 2011). These studies using morphological examination of faecal samples from 109 skinks, revealed predation on Araneae, Blattaria, Chilopoda, Coleoptera, Collembola, Decapoda, 110 Dermaptera, Diptera, Embioptera, Hemiptera, Homoptera, Hymenoptera, Isopoda, Lepidoptera, 111 Opisthopora, Orthoptera, Pseudoscorpionida, Scorpionidae, Stylommatophora and Thysanura. Less 112 information appears to exist on invertebrates in the diets of Asian Musk Shrews, which are generally 113 considered to be highly omnivorous, incorporating significant quantities of arthropods in their diets 114 including Orthoptera, Hymenoptera, Blattaria and Chilopoda (Advani & Rana 1981; Prakash & Singh 115 1999; Lathiya et al. 2008). On Ile aux Aigrettes the African land snail Achatina fulica was consumed 116 when used as bait in traps (Varnham et al. 2002). Given their current wide geographical distribution 117 and adaptability, the shrews are likely to have very different diets within different regions and 118 ecosystems.

119

The problem with morphological identification of prey remains in the guts or faeces of vertebrates is that it is biased towards prey with hard parts that resist digestion (Symondson 2002). It requires a high level of taxonomic skill and the diagnostic features, essential for species-level identification, may not survive digestive processes (Ingerson-Mahar 2002; Sunderland *et al.* 2005). An alternative approach is to analyse gut and faecal samples using PCR (Symondson 2002; King *et al.* 2008), which can now be combined with next generation sequencing (NGS) (Pompanon *et al.* 2012). General

126 invertebrate primers can potentially amplify all invertebrates consumed, generating DNA 'barcodes' 127 (diagnostic sequences from a defined region of a gene) for each prey species (Pompanon et al. 2012). 128 In tropical ecosystems, such as on Ile aux Aigrettes, the invertebrate fauna has not been barcoded 129 and few, if any, taxa are likely to be found on databases such as GenBank or BOLD (Barcoding of Life 130 Database). However, the sequence output from NGS analyses can be clustered into MOTUs 131 (molecular operational taxonomic units) (Floyd et al. 2002) as a proxy for species and can be used to 132 analyse dietary overlap between predator species (Razgour *et al.* 2011). Two predator species may, 133 for example, be consuming the same families of invertebrates but completely different species, and 134 the MOTU approach will reveal this, even when the Linnaean identities of those species cannot be 135 determined. We therefore used next generation sequencing to analyse the invertebrate diets of the 136 shrews and skinks, then tested the hypothesis that there was significant niche overlap between the 137 alien and native species, potentially leading to competition. Tests such as Pianka's niche overlap test 138 (Pianka 1973) do not necessarily reveal where the most significant dietary overlaps lie. We therefore 139 further tested the hypothesis that many prey species are eaten occasionally, probably 140 opportunistically, while a smaller number of key prey species are shared and form a potentially 141 significant part of the diet. Only competition for these prey might be limiting for predator 142 populations. We also tested the hypothesis that shrews and skinks may be competing in a more 143 direct way, by preying on one another.

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146 Methods

147 *Predator sampling* 

Samples were collected over an eight week period from the 10<sup>th</sup> March to the 5<sup>th</sup> May 2011, on Ile aux Aigrettes, Mauritius. This 26 ha coralline island nature reserve is leased to, and managed by, the Mauritian Wildlife Foundation. Shrews were initially caught using Sherman traps. However, trapped

151 shrews had very little material in their guts by the time they were removed. Any remaining gut 152 contents often included bait, and shrews were observed to eat ants from the bait, creating false 153 trophic links. Shrews with full stomachs were subsequently caught more successfully by hand and 154 killed (using UK Home Office approved techniques, Animals (Scientific Procedures) Act 1986) during 155 surveys across the island, both in the early morning and late afternoon/early evening. They were 156 brought back immediately to the field station, dissected under sterile conditions to obtain stomach 157 samples, sexed and measured. Gender was confirmed by post mortem examination for the presence 158 or absence of testes. The length from nose to base of tail was measured to the nearest mm. The 159 presence or absence of foetuses was recorded for females. For males it was often possible to 160 determine adult or juvenile status based on the development of the testes. Females were classed as 161 juveniles if they were less than 12g. The stomach was stored in 94% ethanol at -20°C.

162

Telfair's Skinks were caught by hand and induced to defecate by gently massaging the belly. A sterile tube was placed below the cloaca to catch the faeces, which was topped up with 94% ethanol and kept at -20°C. Animals were sexed using morphological characteristics including hemipenal eversion of males. Each individual was identified from a unique subcutaneous PIT (Passive Integrated Transponder) tag number, which had been implanted during translocation from Round Island. Finally, measurements of snout-vent length (SVL) were taken. For a full list of both shrews and skinks caught and analysed, with measurements, refer to Table S4.

170

171 DNA extraction

DNA was extracted from faecal and gut samples using the QIAmp DNA Stool Mini Kit (QIAGEN), according to the manufacturer's instructions. Additionally, DNA was extracted from a range of invertebrate samples collected from Ile aux Aigrettes, along with tissue samples from shrews and

skinks, for primer testing, using the DNeasy tissue kit (QIAGEN), according to the manufacturer'sinstructions.

177

#### 178 Primer selection for pyrosequencing

179 Published universal PCR primers were tested in a number of different combinations for their ability 180 to amplify DNA from 29 different taxonomic groups of invertebrates (19 orders) collected from Ile 181 aux Aigrettes. Temperature gradient PCRs were performed for each primer pair to determine the 182 optimal annealing temperature at which the most taxa would amplify. PCRs were run on a Peltier 183 Thermal Cycler (Bio-Rad Laboratories, CA, USA) using Multiplex PCR kit (Qiagen) under the following 184 conditions: 1X Master Mix, 0.2  $\mu$ M each primer and 10ng /  $\mu$ L of DNA with an initial denaturation at 185 95°C for 15 min, 45 cycles of 94°C for 30 s, a gradient of 45-60°C for 90 s and 72°C for 90 s, and a 186 final extension at 72°C for 10 min. DNA of the shrews and skinks were also included so that primer 187 pairs which did not cross-amplify with the predators could be identified. Water was included in each 188 PCR in place of DNA as a negative control. From the large number of primers tested (some 189 unpublished) the best proved to be the forward primer LCO-1490 (Folmer et al. 1994) combined with 190 the reverse primer Uni-MiniBar-R (Meusnier et al. 2008), which produced a COI (cytochrome oxidase 191 I) amplicon of 177 bp. These primers were found to amplify 28 of the 29 local taxa at an annealing 192 temperature of 49°C and 42 cycles, with no cross-amplification of the predators (Table S1). A second 193 useful primer pair, combining LCO-1490 with ZBJ-ArtR2c (Zeale et al. 2011), produce a COI amplicon 194 of 225 bp, and was found to amplify 27 of the 29 taxa at an annealing temperature of 52°C and 40 195 cycles (Table S1), but in initial tests weakly cross-amplified the shrew. We therefore used the LCO-196 1490 / Uni-MiniBar-R for further analysis. All other primer combinations tested co-amplified the 197 shrew and/or skink DNA more strongly or amplified a lower range of invertebrate taxa.

198

199 Pyrosequencing

200 LCO-1490 and Uni-MiniBar-R, modified with fusion primers and MIDS (Multiplex Identifiers in the 201 form of unique DNA tags), were used to amplify faecal/gut DNA extracts from shrews and skinks 202 using PCR conditions described above. By using a unique combination of MIDS on both the forward 203 and reverse primers for each individual predator, MOTUs could be assigned to each predator later 204 bioinformatically. DNA from 41 shrew stomach samples and 29 skink faecal samples were 205 successfully amplified. PCR products were run through a 2% agarose gel stained with ethidium 206 bromide and quantified using UVP VisionWorks® LS Analysis software by comparing fluorescence 207 with known concentrations using MassRuler Low Range DNA ladder (Fermentas). Samples were then 208 pooled together in differing proportions to obtain an approximately equal amount of DNA in the final 209 mixed sample. The pooled sample was purified using the QIAquick PCR Purification Kit (QIAGEN) and 210 pooled DNA concentration quantified by Nanodrop ND-1000 Spectrophotometer.

211

212 The DNA was sent to the Genepool, Edinburgh, for NGS. This was performed using the Roche 454 GS-



214

215 Sequence Analysis

Sequences were analysed using the Galaxy platform (<u>https://main.g2.bx.psu.edu/root</u>, Giardine *et al.* 2005; Goecks *et al.* 2010; Blankenberg *et al.* 2010) and Bioedit (T. Hall, http://www. Mbio.ncsu.edu/bioedit/bioedit.html). Rare haplotypes (represented by <3 copies) were removed, plus sequences much longer or shorter than expected, and then aligned with the remaining haplotypes using clustal W in Bioedit. We then edited the alignment manually to remove indels and match reference sequences.

222

The sequences were clustered into MOTUs in the program jMOTU (Jones *et al.* 2011) and tested at thresholds from 1-10 bp. A graph of recovered MOTU vs threshold suggests that a 4 bp cut-off was

225 most appropriate in this data set (see Razgour *et al.* 2011). Representative sequences for each MOTU 226 were compared to the reference database in BOLD (www. barcodinglife.org) recording highest 227 sequence similarity. A phylogenetic tree was constructed of representative MOTUs and a series of 228 known reference sequences using maximum parsimony (MP) in MEGA 5 (Tamura *et al.* 2011) using 229 1000 bootstrap replications.

230

231 Ecological Analysis

Ecological analyses were performed in EcoSim V.7 (http://grayentsminger.com/ecosim.htm) and we compared extents of niche overlap using Pianka's (1973) measure of resource sharing (10000 simulated matrices) between shrews and skinks and between males and females in each predator species (equation 3 in Razgour *et al.* 2011). Null models were used to test whether niche overlap was greater than expected by chance. We then re-ran these analyses excluding prey that were only eaten by a single predator. Such occasional prey species are, individually, unlikely to have a significant effect on nutrition and hence on any prey overlap.

239

Dietary specialization and diversity were estimated using Levins' standardized measure of niche
breadth and Shannon's diversity index (equations 1 and 2 in Razgour *et al.* 2011).

242

243 Prey groups

Representative sequences from each MOTU were compared to sequences in the BOLD reference database and then included, with known references sequences, in a neighbour-joining reconstruction (Figure 1) in MEGA 5 (Tamura *et al.* 2011). The main prey groups were defined in the cladogram (Figure 1) into Lepidoptera, Dictyoptera, Diptera, Araneae and Gastropoda based on both similarity to known references (category 3 classification, Clare *et al.* in review) and clustering with known references sequences in the cladogram. Individual MOTUs which we found in more than 10% of

either shrews or skinks were also analyzed separately. The effects of predator species (shrew or skink), length, mass, age class (juvenile or adult), sex, and whether gravid, on consumption of prey groups, were explored within a Generalised Linear Model (GLM) (data in Table S4). Length was treated as a covariate and all other predictors as factors. The second order interaction predator:sex was included. A binomial error distribution was used with a logit link function. All analyses were conducted in the R statistical package version 2.9.2.

256

257 Species-specific shrew and skink primers

As the primers used for 454 sequencing did not, in practice, co-amplify either the shrew or skink DNA, species-specific primers were needed in order to determine whether there was intraguild predation between the two predators.

261

262 Cytochrome b sequences for the skinks (AF280133) and shrews (JF784171), along with sequences for 263 a broad range of vertebrates know to occur on the island (or their close relatives), were acquired 264 from GenBank and aligned in BioEdit in order to design species-specific primers. NetPrimer (Biosoft 265 International) was used to test primer sequences for potential primer-dimer and hairpins which 266 would reduce primer efficiency. LtF1 (5'-CCG TCC CCT ACA TTG GCA CTG-3') and LtR1 (5'-ACA GGA 267 GGT GAA GGA GAG ATA CC-3') were designed to amplify a 140 bp fragment of the skink while SmF1 (5'- TCG GAA TCT GCT TAA TTG CG-3') and SmR1 (5'- AAT AAC GAA TGA GTC AGC CAT AAT T-3') were 268 269 designed to amplify a 134 bp fragment of the shrew. Gradient PCRs were initially run to determine an 270 optimal annealing temperature for amplification of each target species.

271

272 Primers were tested for cross-amplification against DNA extracted from both shrews and skinks, from
273 a range of invertebrate taxa collected on Ile aux Aigrettes and identified to order (n=14) and

additionally from invertebrates (n=13) and vertebrates (n=10) collected in the UK (see
Supplementary Table S2).

Using the Multiplex PCR Kit (Qiagen) PCR conditions were: 1X Master Mix, 0.5  $\mu$ M each primer , 10% Q solution and 5ng /  $\mu$ L of DNA with an initial denaturation at 95°C for 15 min, 40 cycles of 94 °C for 30 s, 64.5 °C (for LtF/R) and 64 °C (for SmF/R) for 45 s and 72 °C for 30 s, and a final extension at 72 °C for 10 min. DNA samples were each tested twice, with water negatives included. Neither primer pair cross-amplified with any other taxa. Forty eight skink faecal DNA samples were subsequently screened with LtF/R primers and 49 shrew gut content DNA samples were screened with SmF/R primers, using the conditions described above.

284

#### 285 Results

286 Sequence Analysis

Prey DNA was successfully amplified from 42 shrews and 29 skinks, from which 237,402 sequences were recovered. After removal of rare haplotypes we also removed those that were <100bp and >220bp and, using the MID codes, the labelled sequences were assigned to individuals (female shrews n=14, male shrews n=27, one shrew gender unknown, female skinks n=19, male skinks n=10) and aligned using ClustalW in BioEdit. We edited this alignment to a reference sequence to remove indels. This combined screening of data yielded 3001 haplotypes. The primer, MID and adapter sequences were removed for further analysis.

294

The resulting Fasta files in jMOTU (Jones *et al.* 2011) were analysed following the same procedures employed by Razgour *et al.* (2011) resulting in the recovery of 119 MOTUs, using the 4bp threshold for assignment.

298

299 Ecological analyses.

Of the 119 recovered MOTUs, 53 were found in the diet of skinks and 76 from the diet of shrews with 14 shared between the two predators. Within the 53 MOTUs recovered for skinks, 44 were consumed by females, 17 by males and 8 were shared (one could not be assigned to an individual as sequencing did not recover the full MID). Within the 76 MOTUs recovered for shrews, 34 were consumed by females, 52 by males and 10 were shared.

305

306 Niche overlap was significantly greater than expected by chance between predator species (Pianka's 307 measure  $O_{ik}$ =0.55, p=0.012), between shrew males and females ( $O_{ik}$ =0.58, p=0.009) and between 308 skink males and females ( $O_{ik}$ =0.70, p<0.001) (but see Discussion). We then reanalysed the data, 309 excluding 95 MOTUs that were only recorded from the diets of one animal (rare prey), leaving 24 310 MOTUS (out of 119 or 20%) that were consumed at least twice. When prey species detected in only 311 one shrew or skink were excluded (Table S3), prey overlap was shown to be very strong (shrews vs. 312 skinks  $O_{ik}$ =0.80, p=0.002, male vs female shrews  $O_{ik}$ =0.80, p=0.003, male vs female skinks  $O_{ik}$ =0.91, p 313 < 0.0001). Overall, the niche breadth of both predator species was narrow (Levins' measure  $B_4$ =0.18 314 for skinks and  $B_A$  =0.20 for shrews) but high in diversity (H=3.54 for skinks and H=3.74 for shrews). 315 Niche breadth and diversity were similar in shrew females ( $B_A$ =0.26, H=3.27) and males ( $B_A$ =0.30, 316 H=3.53). Niche breadth was larger and higher in diversity in skink females ( $B_A=0.30$ , H=3.46) than in 317 skink males (*B*₄=0.16, *H*=2.69).

318

We could not reliably match any sequences to those in BOLD (<u>www.Barcodinglife.org</u>). A phylogenetic reconstruction of representative sequences for each MOTU was seeded with reference sequences (Figure 1) in order to give an indication of taxonomic groups. This showed a large portion of MOTUs clustering phylogenetically with the reference sequences, suggesting genetic relationships. Of these, 36 MOTUs were most similar to lepidopteran sequences in BOLD and were phylogenetically

324 placed in a clade with known lepidopteran sequences. Similarly, 34 MOTUs showed high sequence 325 similarity to representative Dictyoptera in BOLD (termites, cockroaches and mantids), clustered with 326 known Blattaria in the reconstruction, though a few also showed sequence similarity to reference 327 dipteran sequences.

328

#### 329 Analysis of consumption of prey groups

330 The following analyses were on the putative prey groups as defined in Figure 1. Consumption of 331 Diptera was significantly greater in skinks than in shrews ( $\chi^2 = 11.9$ , df = 1, P < 0.001) (Figure 2a), with 332 41% of skinks found to have consumed Diptera and only 7% of shrews. There was no significant 333 difference in consumption of Gastropoda between shrews and skinks, but male shrews were 334 significantly more likely to consume them than females ( $\chi^2$  = 4.3, df = 1, P = 0.038) (Figure 2b) with 335 44% of males having consumed them and only 14% of females. Consumption of Dictyoptera by shrews appeared higher than that of skinks but this was not quite significant ( $\chi^2$  = 3.3, df = 1, P = 336 337 0.068) (Figure 2c) with 63% of shrews having consumed them and 41% of skinks. Consumption of 338 individual MOTUs, numbers 8, 12 and 13 (all in the Dictyoptera group), were consumed by 20%, 24% 339 and 22% of shrews respectively, but not by any skinks. Conversely, consumption of MOTU number 10 340 (a dipteran) was found to be significantly higher in skinks than in shrews ( $\chi^2$ =10.1, df=1, P=0.001), 341 with 38% of skinks having consumed them compared to 7% of shrews. Length, age class, mass and 342 whether gravid had no significant effect on consumption of different prey groups.

343

344 Species-specific primers

No evidence was found for intraguild predation between the shrews and skinks; none of the shrew

346 gut samples contained skink DNA and none of the skink faecal samples contained shrew DNA.

347

348

349	Discussion
350	Overall our results demonstrate that prey overlap between the shrews and skinks is strong,
351	particularly so when rare prey, consumed only once (80% of prey species detected), were excluded
352	from the analysis. Both analyses may have been affected by sample size (42 shrews and 29 skinks)
353	but the effects are difficult to predict. Larger samples size would increase the probability that less
354	frequently eaten prey will be shared between predator species, but could also increase the number
355	of new rare MOTU's consumed. Rare species (weak links) in food webs may have little influence
356	individually but collectively can increase stability, and this pattern, of many weak links but a few
357	strong links, is commonly found in generalist predator food webs (e.g. McCann <i>et al.</i> 1998). All
358	measures of dietary overlap have been criticised (e.g. Wallace 1981) but when the levels of overlap
359	are so strong they are likely to accurately reflect what is happening in the field. We do not know,
360	however, the degree to which the overlap is driven by prey availability or whether at different times
361	of year prey choices by shrews and skinks change. The fact that so many prey were detected only
362	once implies that both shrews and skinks are adaptable and opportunistic, although more prey
363	species may be shown to be shared by the two predators with more sampling. Similarly, species-level
364	analyses of the diets of bats in previous studies showed rare species comprising approx. 50-90% of
365	recovered MOTUs (Clare et al. 2009, 2011; Bohmann et al. 2011). Strong niche overlap does not
366	necessarily imply significant competition if prey are numerous and not limiting. However, Seymour
367	et al. (2005) provided indirect evidence that prey availability can be limiting, by showing that the
368	mean biomass of shrews increased when their numbers were reduced. It is possible, however, that
369	shrew biomass increased for other reasons, such as reduced intensity of social interactions or
370	changes in abiotic conditions. Our field study coincided with when invertebrate resources are
371	considered to be relatively abundant in comparison to other times of year (Cole & Harris 2011).
372	

373 Although none of the prey could be conclusively identified to a specific taxon, the MOTU approach 374 provided an elegant means of testing for niche overlap between the two predators and between 375 sexes of each predator species, even without access to a reference collection. Data on precisely 376 which prey species are being exploited, particularly those consumed by both shrews and skinks, 377 would require a major barcoding exercise of taxa within the groups indicated on the tree (Figure 1). 378 This would need to be combined with a major effort by museum taxonomists to identify all the taxa 379 morphologically to species. This would not be difficult in, for example, Europe or North America, 380 where the fauna are less diverse and well-studied, but in tropical systems it would present a 381 significant challenge. Only if this were done could the MOTUs found amongst the diets of the shrews 382 and skinks be retrospectively assigned to species. However, analysis of our putative assignments 383 defined in Figure 1 did show some interesting differences. Although Lepidoptera were eaten by both 384 predators, skinks were approximately six times as likely to have consumed Diptera as shrews (Figure 385 2a). The near significantly greater consumption of Dictyoptera by shrews may relate to Blattaria 386 (Figure 2c), although these have been reported to be eaten by both skinks (Vinson & Vinson 1969; 387 Pernetta et al. 2005; Richards 2007; Zuël 2009; Copsey et al. 2011) and shrews (Advani & Rana 1981). 388 Dictyoptera are a superorder containing a large range of ecologically very different taxa (termites, 389 cockroaches and mantids), thus possibly masking dietary differences at the group level.

390

Shrews and skinks clearly have very different physiologies and it might be predicted that the homeothermic shrews would digest their prey more rapidly than poikilotheric skinks. However, we were able to access the shrew samples from an earlier stage of digestion (the stomach) while the skink diet was analysed from fresh faeces. What combined effects these may have had on prey detection, and the relative abundance of different MOTU consumed, could only be established through captive feeding trials.

397

398 Some differences were found between sexes, for example female skinks ate a greater diversity of 399 prey species than males, but the reasons for this, though intriguing, are not known. It may be that 400 the dietary needs of reproducing females are different to those of males. Sex differences in diet are 401 often related to sexual dimorphism, for example in birds and mammals (e.g. Rosalino et al. 2009; 402 Phillips et al. 2011) and arthropods (e.g. Symondson & Liddell 1993; Pekár et al. 2011), where the size 403 difference allows predators to access different prey, allowing intersexual partitioning of resources. 404 Adult male skinks and shrews are larger than females. Male shrews were more than three times as 405 likely to have eaten gastropods than females (Figure 2b). However, all of these results would have 406 been affected by the differences in sample sizes and they would require further work to verify.

407

408 Analysis with species-specific primers provided no evidence of direct intraguild predation by shrews 409 on skinks or skinks on shrews. However, this contrasts with observations on the island of juvenile 410 skink remains in the guts of shrews and shrew remains in the faeces of skinks (pelts and hair), plus 411 direct observations of mutual predation (N. Cole and D. Vencatasamy pers. obs.). Unavoidable delays 412 in conducting our work meant that shrews and skinks were sampled well after the peak period when 413 skinks hatch and are at their most vulnerable. The release of Telfair's skinks onto Ile aux Aigrettes 414 coincided with substantial declines in the abundance of shrews, possibly as a result of skinks preying 415 on shrews. However, at the current low shrew density dietary evidence of predation may not be 416 detected unless the number of skinks sampled was greatly increased. If prey are limiting then high 417 prey overlap between shrews and skinks may also have played a role in the decline of the shrews.

418

Any form of analysis of predation, whether morphological or utilising PCR, must always be qualified
by the fact that we cannot distinguish between predation, scavenging and secondary predation.
Scavenging of dead material by insect predators has been shown to be a likely source of error using
PCR (Foltan *et al.* 2005; Juen & Traugott 2005). Within invertebrate food webs, secondary predation,

where one predator eats another and the prey in the guts of the consumed predator can be detected, is probably a less important source of error (Sheppard *et al.* 2005). In all cases (predation, scavenging, secondary predation) the prey detected are contributing to the nutrition of the predator but the dynamics of the interactions are clearly very different.

427

The novel combination of existing primers proved to be highly effective at amplifying invertebrate DNA, covering a broad range of invertebrates but with no co-amplification of the predators. They proved to be a significant improvement on the Uni-MiniBar primers of Meusnier *et al.* (2008), UniMinibarF1 / UniMinibarR1, which have been criticised for their low taxonomic coverage (Ficetola *et al.* 2010). However, when is UniMinibarR1 combined with the general invertebrate forward primer LCO-1490 of Folmer *et al.* (1994) specificity and coverage were excellent.

434

435 As far as we are aware, this is only the second time that PCR has been used to analyse reptile diets 436 from faecal samples, the first being our previous study of predation on earthworms by slow worms, 437 the legless lizards Anguis fragilis (Brown et al. 2008). In that instance the primers used for NGS were 438 the earthworm group-specific primers developed by Harper et al. (2005). A further paper on the diets 439 of snakes in this special issue reports the vertebrate and invertebrate diet of the smooth snake 440 Coronella austriaca, analysed using prey-specific primers (Brown et al. submitted). The fact that PCR 441 and NGS can be used to analyse the diets of reptiles from faeces, despite the fact that many species 442 digest their prey to the extent of dissolving bones (Secor 2008), opens up a potentially rich field for 443 future research on reptile trophic ecology. A different molecular approach was taken recently by 444 Goiran et al. (2013), who demonstrated that fish eggs palpated from the stomachs of sea snakes 445 could be identified by sequencing their DNA.

446

447 Concerted trapping in 1999 to eradicate the shrews from Ile aux Aigrettes was only partially 448 effective. Some individuals are 'trap-shy' and can go on to generate a resurgent population within a 449 short time. It appears to be the case that shrews enter traps through curiosity, rather than 450 responding to baits (which are often left untouched) (Varnham et al. 2002; Seymour et al. 2005). 451 Thus analysis of their diets in the field provided an opportunity to identify favoured prey that, as bait 452 or food odours, could improve trap efficiency. The results of our analysis suggest that Lepidoptera 453 larvae or cockroaches may provide effective bait. Cockroach frass from laboratory cultures is highly 454 pungent and may be sufficient to attract shrews.

455

456 The ethics of killing vertebrates in order to obtain gut samples must be properly justified. Here we 457 caught and killed shrews in the field (using UK Home Office approved techniques), to obtain gut 458 samples. Once caught it was not considered ethically acceptable to release these pests back to the 459 wild, where they would continue to pose a threat to native wildlife. This allowed us to maximise the 460 information obtainable from these animals by analysing their stomach contents (rather than faeces) 461 where prey DNA was likely to be less degraded. A key aim of Mauritian conservationists has been to 462 eradicate shrews from offshore islands to permit further restoration processes. However, to date, 463 eradication attempts have only been successful using traps on topographically simple islands of a few 464 hectares or less (Varnham et al. 2002). The problem with the shrews is that traps do not catch them 465 efficiently and suitable poison baits have not been devised (Varnham et al. 2002; Seymour et al. 466 2005).

467

Our conclusion, therefore, is that shrews and skinks are feeding to a large extent on the same species of invertebrate prey, potentially leading to competition. If so then shrew control is likely to be beneficial to the fitness of the skinks. Mutual predation is known to occur, but our analysis failed to find evidence of this outside the period when juvenile skinks are particularly vulnerable. This is

472	probably because skinks grow too large to be attacked by shrews and similarly, at low densities,
473	shrews increase in biomass (Seymour et al. 2005) and may be too large for predation by skinks. Given
474	that the shrews pose a threat to island biodiversity, development of methods to eradicate them from
475	islands such as Ile aux Aigrettes should continue to be a priority.
476	
477	
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485	
486	
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641	Data accessibility
642	All sequences arising from NGS, fully processed, collapsed and aligned, plus allocated to individual
643	predators and ready for analysis, will be included as Online Supplementary Material after acceptance
644	of the paper. Three files will be included: all 'sequences pooled.fasta', 'all sequences shrew.fas' and
645	'all sequences skink.fas'.
646	
647	Author Contributions Box
648	Gut and faecal samples were collected from Mauritius by RB and DV, and DNA extracted by RB, who
649	designed and applied species-specific primers for analysing mutual predation between shrews and
650	skinks. Preparation of samples for NGS was conducted by DSB, along with analyses of predation on
651	key prey taxa. Bioinformatics and ecological analyses were conducted by ELC. Supervision of the
652	fieldwork in Mauritius was conducted by NC, who provided the expertise on Mauritian ecology. AM
653	conducted the 454 analysis. Overall supervision of the project and the writing of the paper were
654	primarily conducted by WOCS, with major contributions from co-authors.
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#### 665 Figure legends

666 Figure 1

667 Cladogram showing reconstructed relationships between all MOTUs retrieved from the guts or faecal

samples of Asian Musk Shrew and Telfair's Skinks, colour codes to denote prey consumed by shrews,

669 skinks or by both species.

670

671 Figure 2

672 Main significant or near significant differences in diet arising from analysis of putative higher-order

673 classifications, as defined in Figure 1. **a.** Predicted probability of consumption of Diptera (± s.e.) by

674 shrews and skinks, showing significantly higher consumption in skinks (*p* < 0.001). **b.** Predicted

675 probability of consumption of Gastropoda (± s.e.) by shrews, showing significantly higher

676 consumption in males than in females (p = 0.038). **c.** Predicted probability of consumption of

677 Dictyoptera (± s.e.) by shrews and skinks, showing a trend towards higher consumption by shrews (p

678 = 0.068).











2c

# **1 SUPPLEMENTARY MATERIAL**

- 2 Table S1
- 3 Invertebrates collected from Ile aux Aigrettes and tested for PCR amplification with the two primers
- 4 sets developed for 454 pyrosequencing, LCO-1490 / Uni-MiniBar-R and LCO-1490 / ZBJ-ArtR2c.
- 5

Potential prey	LCO-1490 / Uni-	LCO-1490 / ZBJ-
	MiniBar-R	ArtR2c
Coleoptera 1	V	v
Oligochaeta	v	v
Hemiptera 1	v	V
Isopoda	V	V
Dermaptera	V	V
Embioptera	V	V
Diplopoda	V	V
Hymenoptera (Vespa sp.)	v	V
Araneae 1	V	V
Gastropoda 1	v	V
Lepidoptera 1	v	V
Diptera	v	V
Blattaria 1	v	V
Odonata	v	v
Decapoda	v	V
Gastropoda 2	v	v

Hymenoptera - Formicoidea	v	v
Lepidoptera 2	٧	<b>√</b>
Scorpiones	V	v
Araneae 2	V	v
Coleoptera - Cerambycidae	V	v
Diptera - Culicidae	V	v
Collembola	V	√
Orthoptera - Gryllidae		V
Hemiptera 2	V	v
Hemiptera 3	V	
Chilopoda	V	√
Coleoptera 2	٧	
Blattaria 2	٧	√
Total	28/29	27/29

16 Table S2.

- 17 Non-target species tested for cross-amplification with skink-specific (LtF/R) and shrew-specific
- 18 (SmF/R) PCR primers. Neither primer set co-amplified any of these taxa.
- 19

Order	Species	Origin
Coleoptera	spp. x2	Ile aux Aigrettes
Lepidoptera	spp. x2	lle aux Aigrettes
Blattaria	spp. x2	Ile aux Aigrettes
Hymenoptera	spp. x2	lle aux Aigrettes
Diptera	spp. x2	lle aux Aigrettes
Isopoda	spp. x2	lle aux Aigrettes
Aranaea	spp. x2	Ile aux Aigrettes
Pulmonata	Arion intermedius	UK
	A. distinctus	UK
	Limax flavus	UK
Haplotaxida	Lumbricus terrestris	UK
	L. rubellus	UK
	Aporrectodea caliginosa	UK
	A. longa	UK
Coleoptera	Notiophilus biguttaus	UK
	Adalia bipunctata	UK
	Tachyporus obtusus	UK
Diptera	Tipulidae sp.	UK

Dermaptera	Forficula sp.	UK
Aranaea	Erigone ddentipalpis	UK
Squamata	Zootoca vivipara	UK
	Anguis fragilis	UK
	Coronella austriaca	UK
	Natrix natrix	UK
Rodentia	Myodes glareolus	UK
	Mus musculus	UK
	Apodemus flavicollis	UK
Soricomorpha	Neomys fodiens	UK
	Sorex araneus	UK
Caudata	Lissotriton helveticus	UK



31 Table S3

Numbers of shrews and skinks, of each sex, that contained each prey MOTU, excluding MOTUs that
were only found in one animal overall. Shrew N/R is an animal not sexed (see text). 'Total detections'
are the numbers of shrews+skinks testing positive for that MOTU. For a complete list, and to find
MOTU numbers, see Figure 1.

36

MOTU no.	Skinks	Skinks	Shrews	Shrew	Shrew N/R	Total
	male	female	male	female		detections
2	3	7	10	6	1	27
3	1	1	0	2	0	4
4	3	3	6	5	1	18
5	1	6	10	5	0	22
6	3	5	5	0	0	13
7	1	5	10	3	1	20
8	0	0	3	5	1	9
10	3	8	3	0	0	14
11	0	0	3	0	0	3
12	0	0	7	3	1	11
13	0	0	6	3	1	10
16	0	0	2	0	0	2
20	0	0	3	0	0	3
21	0	0	4	1	0	5

28	0	0	3	0	0	3
30	0	0	1	1	0	2
31	0	0	3	0	0	3
34	0	1	3	2	0	6
39	1	2	2	2	0	7
44	1	3	0	1	0	5
49	0	1	0	1	0	2
59	0	1	1	0	0	2
71	1	1	0	0	0	2
116	0	2	0	0	0	2

52 Table S4

File 'MOTUs consumed by shrews and skinks.xls'. Spreadsheet providing raw data on the shrews and
skinks from which we successfully amplified invertebrate DNA, including sex, mass, length,
adult/juvenile status and whether gravid.

56

57 Tables S5-S6

58 Spreadsheets including representative sequences for all haplotypes arising from NGS, fully 59 processed, collapsed and aligned, allocated to individual predators and ready for analysis. Divided 60 into 'All sequences shrew.fas' and 'All sequences skink.fas'.

	Skink		Shrew		
ΜΟΤυ	Males	Females	Males	Females	Unknown
2	11, 31, 46	20, 33, 36, 41, 45, 7, 9	11, 13, 21, 26, 33, 36, 44, 6, 8, 9	1, 25, 28, 29, 7, 37	2
3	42	20		7, 32	
4	10, 31, 42	3, 7, 9	12, 22, 26, 40, 41, 44	19, 20, 38, 39, 32	2
5	11	20, 29, 2, 36, 41, 44	13, 15, 21, 26, 33, 35, 44, 6, 8, 9	25, 29, 38, 7, 50	
6	11, 15, 42	12, 18, 20, 36, 41	12, 21, 22, 26, 33		
7	39	18, 20, 33, 41, 48	14, 17, 22, 30, 34, 3, 41, 43, 44, 9	45, 46, 49	2
8			21, 28, 8	1, 25, 7, 32, 37	2
10	11, 43, 6	14, 2, 33, 3, 40, 41, 44, 45	15, 40, 44		
11			41, 44, 9		
12			15, 16, 17, 26, 33, 41, 48	24, 25, 49	2
13			21, 33, 36, 44, 8, 9	25, 28, 37	2
16			40, 41		
20			26, 33, 48		
21			22, 26, 48, 46	49	
28			26, 33, 48		
30			41	29	
31			26, 33, 48		
34		9	40, 44, 9	37, 50	
39	42	4, 9	40, 44	4, 33	
44	37	33, 41, 9		49	
49		18		24	
59		7	6		
71	11	41			
116		12, 41			