1 Ecological flexibility as measured by the use of pioneer and exotic plants in two lemurids:

2 Eulemur collaris and Hapalemur meridionalis

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Short title: Do lemurid diets shape ecological flexibilities?

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Keywords: pioneer species; exotic plants; edge habitat; collared brown lemur; southern bamboo
lemur; Madagascar

8

9 Abstract

Primate responses to habitat alteration vary depending on the species' dietary guild and 10 forest type. Leaves from secondary vegetation can provide nutritious resources to folivorous 11 12 primates, whereas frugivores, burdened with a scattered spatial and temporal distribution of fruiting resources, require larger home ranges, potentially limiting their ability to cope with 13 altered landscapes. Within coastal south-eastern Madagascar, we sought to determine whether 14 15 two lemur species occupying contrasting ecological niches (i.e., dietary guilds) respond differently to the changing features of their degraded and fragmented habitat. We conducted 16 behavioural observations between 2011 and 2013 on frugivorous collared brown lemurs 17 (Eulemur collaris) and folivorous southern bamboo lemurs (Hapalemur meridionalis). In order 18 to estimate the ability of lemurs to use pioneer species, we categorised all plants used for feeding 19 and resting as either 'fast-growing', 'mid-growing', or 'slow-growing'. We fitted linear mixed-20 effects models, one for each plant growth category with monthly proportional use rates as the 21 dependent variable, and included species (E. collaris and H. meridionalis), activity (feeding and 22 23 resting), and season (dry and wet) as fixed effects. Our results show that E. collaris used both

slow- and mid-growing plant species most often, while *H. meridionalis* were more likely to use
fast-growing plants, which indicated an ability to utilise secondary/disturbed vegetation.

26 Frugivorous E. collaris appear more limited by climax plants, while folivorous H. meridionalis

appear to be slightly more adaptable, a finding that is consistent with other primate folivores.

28

29 Introduction

Tropical deforestation is one of the primary threats to global biodiversity (Achard et al. 30 2002; Asner et al. 2009; Dirzo and Raven 2003; Gibson et al. 2011; Sala et al. 2000). The 31 destruction, fragmentation, and degradation of remaining habitats threaten many species' ability 32 to survive (Oates 2013). While forest fragments typically persist after deforestation, they 33 effectively become islands within an anthropogenic landscape, most of which are unsuitable 34 habitat for the majority of forest species (Broadbent et al. 2008; Laurance et al. 2009, 2011). 35 Ecological flexibility is loosely defined as the ability of an organism to adjust to changes, 36 e.g., anthropogenic, gradual, and stochastic, within its environment (Isaac and Cowlishaw 2004; 37 38 Nowak and Lee 2013; Wieczkowski 2003). In more specific terms, *flexibility* may encapsulate various behavioural modifications including the diet, i.e., exploitation of alternative food 39 sources, as well as altering activity, ranging pattern and vertical strata use in response to new 40 dietary opportunities. This ability to expand niche breadth is key to withstanding the risks of 41 anthropogenic and/or stochastic habitat modification (Lee 2003). 42

It is important to understand behavioural responses of forest dwelling primates to habitat degradation and fragmentation due to the increasing rate of habitat alteration and limited ability of most species to move between forest fragments (Marsh 2003). How a primate responds to habitat degradation, however, seems to vary depending on species and type of forest (Chapman

et al. 2000; Cowlishaw and Dunbar 2000; Onderdonk and Chapman 2000). Secondary growth 47 may produce foods of higher dietary quality compared to foods available in mature forests, thus 48 making folivorous (i.e., leaf-eating) primates less affected by habitat degradation (Chapman et al. 49 50 2002; Ganzhorn 1995; Ganzhorn et al. 1999b; Plumptre and Reynolds 1994). For example, populations of folivorous black howler monkeys (*Alouatta carava* and *A. pigra*), have been 51 documented to use and rely heavily on fast-growing, exotic plant species (e.g., *Eucalyptus* and 52 shaded cocoa plantations) for both occasional food and resting/sleeping within fragmented, 53 anthropogenic landscapes (Bicca-Margues and Calegaro-Margues 1994; Bonilla-Sánchez et al. 54 2012; Zárate et al. 2014). Similarly, black-and-white colobus (Colobus guereza) appear to do 55 well in some disturbed (i.e., previously logged) habitats (Chapman et al. 2000; Tutin et al. 56 1997b). Frugivorous (i.e., fruit-eating) primates, however, have to cope with the scattered spatial 57 and temporal distribution of fruiting resources, thus often requiring larger home ranges (Estrada 58 59 and Coates-Estrada 1996; Rode et al. 2006; but see Tutin et al. 1997a). Many frugivorous primates avoid forest fragments, e.g., grey-cheeked mangabeys (Lophocebus albigena) and 60 61 Mexican spider monkeys (Ateles geoffroyi vellerosus), and appear to be restricted to continuous forests (Estrada and Coates-Estrada 1996; Tutin et al. 1997b). Despite these potential limitations, 62 some frugivorous primates, such as chimpanzees (*Pan troglodytes*) and Sumatran orangutans 63 (Pongo abelii), demonstrate an ability to survive within degraded, anthropogenic landscapes, 64 foraging on a mixture of crops and wild fruits (Campbell-Smith et al. 2011; Hockings and 65 McLennan 2012; McLennan and Hockings 2014). As frugivorous primates are important seed 66 dispersers, their ability to cope within anthropogenic landscapes has major implications for the 67 maintenance of forest diversity: they are fundamental in the regeneration of degraded habitats 68 (Chapman 1995; Ganzhorn 1995; Razafindratsima and Dunham 2014). 69

70	On a global scale, frugivorous strepsirrhines from Madagascar contribute on a larger
71	scale to their respective ecosystems, e.g., seed dispersal, compared to primates in the Neotropics
72	or mainland Africa (Jernvall and Wright 1998). As frugivorous lemurs are essential to
73	maintaining the unique forests of Madagascar, their demise would likely trigger extinction
74	cascades (Federman et al. 2016; Ganzhorn et al. 1999a; Jernvall and Wright 1998;
75	Razafindratsima and Dunham 2014). Within Madagascar, for example, greater than 80% of
76	forest area exists less than 1 kilometre from an edge (Harper et al. 2007), and thus fragmentation
77	is of great concern for the survival of forest fauna and flora species (Hannah et al. 2008; Waeber
78	et al. 2015). This can be further complicated by introduced exotic and invasive species that
79	threaten the preservation of endemic biodiversity as well as ecosystem restoration efforts
80	(Braithwaite et al. 1989; D'Antonio and Vitousek 1992). While the limits of lemurs' tolerance,
81	i.e., coping strategies, to fragmented, secondary, and degraded habitats are poorly understood
82	(Campera et al. 2014; Donati et al. 2011; Eppley et al. 2015a; Gardner 2009; Irwin et al. 2010;
83	Lehman et al. 2006), it is imperative to understand the relationship between species and these
84	altered habitats if we are to properly conserve primates and other species (Cristóbal-Azkarate
85	and Arroyo-Rodríguez 2007; Isabirye-Basuta and Lwanga 2008; Onderdonk and Chapman
86	2000).

The Anosy region along the southeast coast of Madagascar provides a complex mosaic of heavily fragmented upland and swamp forest habitats, mono-dominant exotic species, old and new timber plantations, and a large-scale ilmenite ore mine and separation plant facility (Barthlott et al. 1996; Ramanamanjato et al. 2002; Ganzhorn et al. 2007b). This area provides an excellent model with which to explore the behavioural and feeding ecological flexibilities amongst the lemurs which inhabit it (Bollen and Donati 2006; Eppley et al. 2015a;

Rabenantoandro et al. 2007), and how they cope with habitat disturbance. Within southeast 93 Madagascar, sympatric collared brown lemur (Eulemur collaris) and southern bamboo lemur 94 (Hapalemur meridionalis) occupy different ecological niches, the frugivorous and folivorous 95 96 dietary guild, respectively. Previous research has shown that E. collaris is tolerant to habitat degradation and strong seasonal resource availability by flexibly modifying many aspects of its 97 behavioural ecology, such as feeding strategies and home range use (Campera et al. 2014; Donati 98 et al. 2011). Similarly, *H. meridionalis* display a flexible ecology, utilising three distinct habitats 99 (littoral forest, littoral swamp, and Melaleuca-dominated swamp) for both resting and feeding 100 purposes (Eppley et al. 2015a). 101 While the use of disturbed habitats by these two lemurids has been increasingly 102 documented (Campera et al. 2014; Eppley et al. 2015a), our study sought to contrast the two taxa 103 104 and specifically determine to what extent their ecological flexibility played a role in their ability to use altered and degraded habitats. Habitat edges often contain a higher abundance of pioneer, 105 i.e., fast-growing, species as compared to climax habitat (Laurance et al. 2006, 2007). 106 107 Furthermore, climax, i.e., slow-growing, plants struggle to regenerate in open habitats as is very often the case in edge areas and/or plantations (Benitez-Malvido 1998). The general observation 108 that folivores are able to cope better within degraded environments led us to predict that H. 109 meridionalis will use more forest edge habitat compared to E. collaris. In terms of utilising plant 110 species as feeding and/or resting resources, we sought to determine whether the dietary guilds of 111 our two taxa are predictive of an ability to use fast growing plant species, typically represented 112 by pioneer species, and how this is a potential signal of ecological flexibility to altered habitats. 113 As fast-growing tropical plant species often provide a continuous (i.e., non-seasonal) and 114 relatively large biomass presence of young leaves (Coley et al. 1985; Poorter 1999), which 115

consequently produce more protein (Wasserman and Chapman 2003), we predict that H. 116 *meridionalis* will use fast-growing tree species more often than sympatric frugivorous E. 117 *collaris.* Due to the contrasting dietary guilds of our two study species, we predict that there will 118 119 be little overlap in feeding resources. As these are the two largest lemur species within the southeast coastal landscape, however, we predict that they will use similar tree species for 120 resting, as this degraded littoral environment has a limited number of mature trees. We expect 121 122 the largest differences in feeding tree use to be evident for *E. collaris* during the dry season, i.e., when there are fewer available food resources (Bollen and Donati 2005; Campera et al. 2014). 123 Lastly, exotic plant species (e.g., introduced, non-endemic), are shown to incur lower levels of 124 leaf herbivory compared to endemic plant species (Lake and Leishman 2004). Thus, we predict 125 that the folivorous *H. meridionalis* would avoid exotic plant species. 126

127

128 Methods

129 Study site

We conducted our study in the Mandena Conservation Zone (24°95'S 46°99'E; hereafter 130 Mandena), along the southeast coast of Madagascar, approximately 10 km north of Fort-Dauphin 131 (Tolagnaro). Located within three kilometres of the coast and characterized by a low canopy 132 growing on sandy substrate (Dumetz 1999), this protected area consists of approximately 82 ha 133 of seasonally inundated swamp among 148 ha of degraded littoral forest fragments (Ganzhorn et 134 al. 2007a). This littoral zone experiences less seasonality than the humid eastern forests (Bollen 135 and Donati 2005), with a mean temperature of 22.5°C (range: 9.5 – 35.0°C) and total annual 136 precipitation of 2,808mm, typically generating a wet season between November and April 137 (Eppley et al. 2015a, 2016b). Compared to the less degraded littoral forests further north (Bollen 138

and Donati, 2006), the degree of anthropogenic degradation in Mandena resulted from the 139 historical extraction of utilitarian timber species and charcoal production due to the close 140 proximity of the Anosy region capital (Ingram and Dawson 2006; Vincelette et al. 2007b). The 141 142 area immediately surrounding these fragmented forests is composed of mono-dominant timber plantations, an exposed sand-scrub matrix, and the large-scale ilmenite mining concession and 143 associated administration and extraction/separation facilities (Ganzhorn et al. 2007b). In addition 144 to the two cathemeral lemurids, i.e., E. collaris and H. meridionalis, this littoral area is inhabited 145 by four nocturnal strepsirrhines: Ganzhorn's mouse lemur (Microcebus ganzhorni), eastern fat-146 tailed dwarf lemur (*Cheirogaleus medius*), greater dwarf lemur (*C. major*), and the southern 147 woolly lemur (Avahi meridionalis). 148

149

150 Study species

Our study focuses on two sympatric lemurs inhabiting Mandena: E. collaris and H. 151 meridionalis. Both are medium-sized lemurs, although E. collaris is considerably larger, with a 152 153 mean body mass of 2.2 kg (Donati et al. 2011), compared to the mean body mass of H. meridionalis which is 1.1 kg (Eppley et al. 2015b). Both of these lemurid species exhibit a 154 cathemeral activity pattern (Donati et al. 2007; Eppley et al. 2015c). Species are classified 155 according to dietary guild based on diets comprising \geq 50% of a specific food category 156 (Ganzhorn 1997). As the annual diet of *E. collaris* consists of \geq 70% fruits, it is classified as 157 frugivorous (Donati et al. 2007, 2011). The annual diet of *H. meridionalis* consists of \geq 70% 158 foliose matter, thus this species is classified as folivorous (Eppley et al. 2011, 2016a). 159 We captured lemur subjects via Telinject[®] blow darts (administered by an experienced 160 Malagasy technician) containing a hypnotic anaesthesia (4-5 mg/kg of ketamine hydrochloride or 161

162	tiletamine hydrochloride. Four individuals (one for each group) of E. collaris were captured and
163	equipped with radio-collars (TW-3, Biotrack, 29 g). We captured ten individuals of H.
164	meridionalis from four social groups, and radio-collared with data-logging tags (ARC400,
165	Advanced Telemetry Systems, Inc.; Isanti, MN, USA). We utilised radio-collars to expedite the
166	amount of time it took to locate lemur groups each day; however, not all adult focal individuals
167	were radio-collared. All subjects recovered from anaesthesia within 1.5 h and were not moved
168	from the capture area. Furthermore, we followed lemurs until they regained full mobility in trees.
169	There were no injuries as a consequence of the captures. The collars were below the 5%
170	threshold of the subjects' weight. For more specific information on the capturing/collaring
171	processes of <i>E. collaris</i> , see Campera et al. (2014), and for <i>H. meridionalis</i> , see Eppley et al.
172	(2015c, 2016c).

174 Data collection

We collected data for each species during different years. For E. collaris, MB and MC 175 176 observed group AB from March 2011 to January 2012, and group C from June 2011 to January 2012. Data collection was conducted on a focal individual from 06:00 to 18:00 h. We collected 177 behavioural data in 5 min intervals via instantaneous sampling (Altmann 1974), specifically 178 179 noting the tree species used for feeding and resting. Furthermore, we recorded the position of the focal E. collaris individual in 30 min intervals via a handheld GPS. For H. meridionalis, TME 180 conducted full-day focal observations (from sunrise to sunset) with groups 1, 2, and 4 (we used 181 group 3 exclusively for home range data collection) between January and December 2013. We 182 identified all observed plant food items consumed by the focal individual, noting the plant 183 species' scientific name, and recorded feeding duration via continuous sampling (Altmann 184

185	1974). Furthermore, we recorded all instances ≥ 15 min for continuous resting. Lastly, we
186	recorded <i>H. meridionalis</i> focal waypoint locations via GPS in 15 min intervals.
187	JR and FR identified all plant species used for feeding and resting by both lemur species,
188	and we categorized these into three successive growth rates as they occur under natural
189	conditions. As such, <i>fast-growing</i> plant species reached maturity < 2 years, <i>mid-growing</i> plant
190	species reached maturity between $2-5$ years, while <i>slow-growing</i> species reached maturity in >
191	5 years, with categories based on previous botanical assessments (cf. Vincelette et al. 2007a).
192	Furthermore, JR identified exotic plant species (i.e., non-endemic), which we validated with an
193	index of exotic and invasive species in Madagascar (Gérard et al. 2015).
194	
195	Ethical Note
196	Our research protocols were approved and permits authorized by the Commission
197	Tripartite of the Direction des Eaux et Forêts de Madagascar (Autorisation de Recherché
198	n.29/11/MEF/ SG/DGF/DCB.SAP/SCB du 20/01/11 and
199	n.240/12/MEF/SG/DGF/DCB.SAP/SCB du 17/09/12), adhering to the legal requirements of
200	Madagascar. We conducted research under the collaboration agreement between the Department
201	of Animal Biology of the University of Antananarivo and the Department of Animal Ecology
202	and Conservation of the University of Hamburg, and QIT Madagascar Minerals (QMM).
203	
204	Data analyses
205	We entered all ranging data into ArcGIS 10.2 (ESRI) using the Geospatial Modelling
206	Environment (GME) spatial ecology interface (Beyer 2012). Ranging and statistical analyses
207	were conducted using R statistical software version 3.2.3 (R Development Core Team 2015). We

determined each group home range with a 95% kernel density estimate, while core areas were
determined as a 50% kernel density estimate (Worton 1989). We then created a forest edge
polygon buffer 100 m inside the littoral forest boundary which allowed us to calculate the total
amount of forest edge and non-edge habitat within each lemur species' home range and core
areas within Mandena (Laurance et al. 2007; Lehman et al. 2006).

From our behavioural sampling of *E. collaris* and *H. meridionalis*, we calculated monthly 213 proportional utilisation rates for all feeding and resting trees. For each of the three plant species 214 growth categorizations, we fitted General Linear Mixed-effects Models (LMM) using the Imer 215 function of the lme4 package developed for R (Bates et al. 2012). For each LMM, our dependent 216 response variable was the monthly proportion of plant species used (i.e., fast-growing plants, 217 mid-growing plants, and slow-growing plants), while our fixed effects were the lemur species (E. 218 219 *collaris* and *H. meridionalis*), activity (feeding and resting), and season (dry and wet). We included lemur social group as random effect to control for repeated sampling. We then used the 220 ANOVA function to calculate likelihood ratio tests for model comparison, allowing us to 221 222 determine which model had the best explanatory power by comparing Akaike's Information Criterion (AIC) values for all possible models. P-values were obtained with a likelihood ratio 223 test using the afex package (Singmann 2014) developed for R, with significance considered at P 224 225 < 0.05. Residuals from the analyses did not deviate from normality according to the Kolmogorov-Smirnov test. 226

To determine which factors are linked to the utilisation of exotic plants within Mandena, we fitted Generalized Linear Mixed-effects Models (GLMM) using the glmer function of the lme4 package developed for R (Bates et al. 2012), with the monthly use of an exotic plant as a binomial dependent variable, as opposed to endemic plants. As with the LMMs, our fixed effects

were lemur species, activity, and season, with group included as random effect to control for
repeated sampling. We then used the ANOVA function to calculate likelihood ratio tests for
model comparison and determined which model had the most explanatory power by comparing
the AIC values for all possible models.

- 235
- 236 **Results**
- 237 Ranging

We observed E. collaris for 962 h, while H. meridionalis were observed for 1,762 h. Both 238 lemurid species' home ranges were within the central to northern portions of Mandena, and were 239 not limited to only littoral forest areas, but rather encompassed a mixture of both littoral forest 240 and swamp (Fig. 1). Considering species' home ranges, E. collaris used considerably larger areas 241 than *H. meridionalis* (Table 1). The proportion of edge habitat used by both species within their 242 home range were similar, with forest edge comprising a mean of 37.4% of *E. collaris* home 243 ranges (N = 2), and 45.6% of *H. meridionalis* home ranges (N = 4) (Table 1). Considering only 244 245 the core areas, forest edge comprised similar mean percentages of E. collaris (50.6%) and H. meridionalis (42.6%) habitat. 246

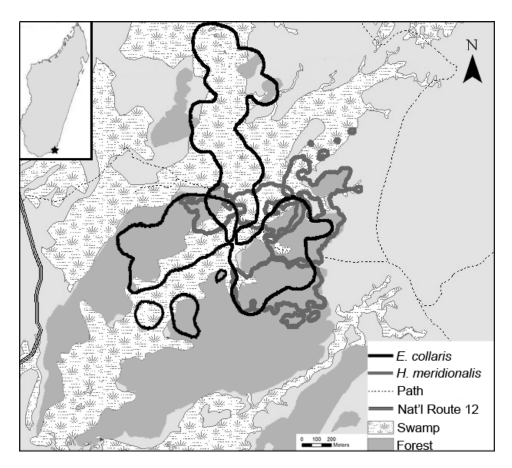


Fig. 1. Location of *E. collaris* and *H. meridionalis* group home ranges (95% kernel density
estimates) within the Mandena littoral forest and swamp. Portions of the swamp are composed of
mono-dominant strands of exotic *Melaleuca*, while lighter grey areas to the east are a sand-scrub
matrix and those to the west are a matrix of sand-scrub and Eucalyptus plantations. Data on *E. collaris* were collected between March 2011 and January 2012, and *H. meridionalis* between
January and December 2013.

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Table 1. Area (in hectares) of both home range (95% kernel density estimate) and core area
 (50% kernel density estimate) for *E. collaris* and *H. meridionalis* groups in Mandena. Edge

habitat was calculated as the area (ha) within 100 m buffer from the forest edge. Data were

collected on *E. collaris* between March 2011 and January 2012, and on *H. meridionalis* between

			Home range (ha)	_		Core area (ha)	_	
		Months of		-			_	
Species	Group	observation		Edge (ha)	Edge (%)		Edge (ha)	Edge (%)
E. collaris								
	AB	11	41.16	19.07	46.33	5.61	4.31	76.83
	С	8	83.32	23.68	28.42	17.98	4.39	24.42
H. meridionalis								
	1	12	18.39	8.40	45.68	7.76	1.20	15.46
	2	12	17.66	9.75	55.21	3.74	2.68	71.66
	3	12	6.60	3.65	55.30	N/A	N/A	N/A
	4	12	10.43	2.75	26.37	2.09	0.85	40.67

260 January and December 2013.

262 Diet

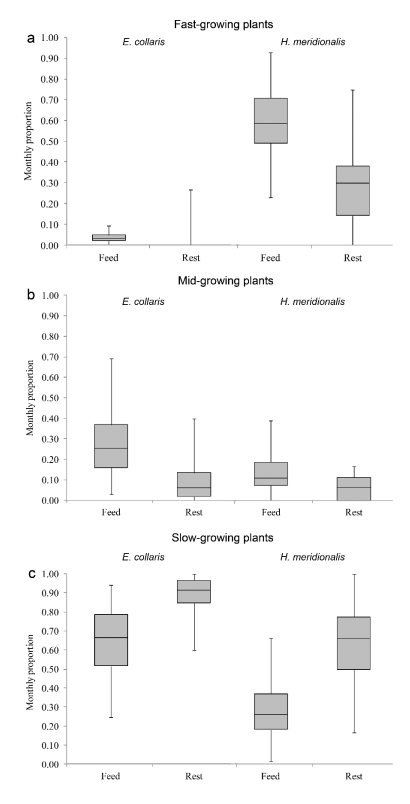
263	We identified 105 different plant species used by <i>E. collaris</i> and 112 species used by <i>H</i> .
264	meridionalis for feeding and resting (Table 2). Twenty-four plant species were eaten by both
265	lemurs. More specifically, E. collaris food resources comprised 16.9% fast-growing, 28.6% mid-
266	growing, and 54.6% slow-growing plants. For <i>H. meridionalis</i> , food resources comprised 38.0%
267	fast-growing, 16.9% mid-growing, and 45.1% slow-growing plants. Twenty-seven plant species
268	were used for resting by both lemur species. For <i>E. collaris</i> , we categorized 14.3% of all resting
269	plants as fast-growing, 24.8% as mid-growing, and 61.0% as slow-growing, while for <i>H</i> .
270	meridionalis, we categorized 27.7% of their used plants as fast-growing, 14.3% as mid-growing,
271	and 58.0% as slow-growing. Both E. collaris and H. meridionalis displayed large differences in
272	their utilisation of these plant growth categories between activity (Fig. 2) and season (Fig. 3).
273	

Table 2. Number of species within feeding and resting plant species' growth categorization for
 E. collaris (March 2011 to January 2012) and *H. meridionalis* (January to December 2013) in
 Mandena.

	<u>E. collaris</u>			<u>H. meridionalis</u>		
Plant categories	Feed	Rest	Total	Feed	Rest	Total
Fast-growing	13	2	15	27	5	31
Mid-growing	22	11	26	12	8	16
Slow-growing	42	43	64	33	57	65

Note: *E. collaris* also relied on six unidentified plant species that were not included in the
analyses

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Fig. 2. Comparison of monthly proportional medians (including interquartiles and ranges)

between *E. collaris* and *H. meridionalis* on their selection of (a) fast-growing, (b) mid-growing,

- and (c) slow-growing plants for feeding and resting. Data were collected between March 2011 and January 2012 on E collaring and between January and December 2012 on H maridian discussions
- and January 2012 on *E. collaris*, and between January and December 2013 on *H. meridionalis*.

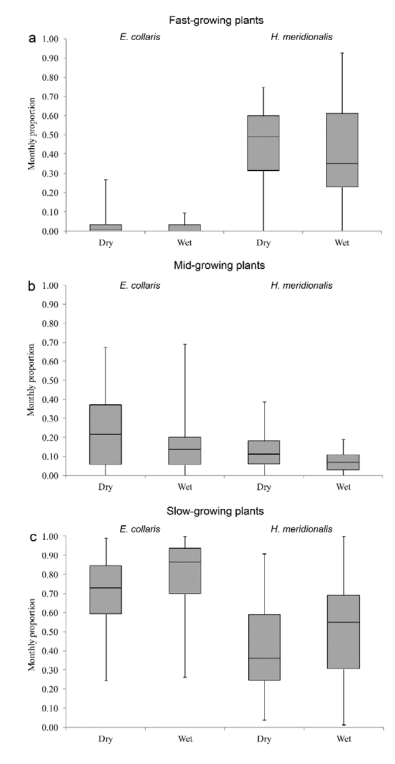




Fig. 3. Comparison of monthly proportional medians (including interquartiles and ranges)

between *E. collaris* and *H. meridionalis* on their selection of (a) fast-growing, (b) mid-growing,

and (c) slow-growing plants during the dry and wet seasons. Data were collected between March

291 2011 and January 2012 on *E. collaris*, and between January and December 2013 on *H*.

292 *meridionalis*.

294	The model with the best predictive value for fast-growing plants (AIC = -93.06, χ^2 =
295	21.59, $df = 1$, $P < 0.001$) showed that both lemur species and activity were likely to influence
296	their use (Table 3) while season had no effect. Specifically, fast-growing plants were most likely
297	to be used by <i>H. meridionalis</i> , and most often for feeding (Fig. 2a). Season was not significantly
298	predictive (Fig. 3a). The model with the best predictive value for mid-growing plants (AIC = $-$
299	163.11, $\chi^2 = 9.29$, $df = 1$, $P < 0.01$) showed that all fixed-effects, i.e., species, activity, and
300	season, influenced use of these plants (Table 3). Specifically, E. collaris was most likely to use
301	mid-growing plants. Furthermore, these plants were more likely to be used for feeding (Fig. 2b),
302	and to be used in the dry season (Fig. 3b). The model with the best predictive value for slow-
303	growing plants (AIC = -63.90, χ^2 = 21.87, df = 1, P < 0.001) showed again that all fixed-effects,
304	i.e., species, activity, and season, influenced use of these plants (Table 3). E. collaris was most
305	likely to use slow-growing plants, with these plants most often used for resting (Fig. 2c),
306	specifically during the wet season (Fig. 3c).
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323	Table 3. Linear mixed models	predicting increased	d monthly proportion o	f using fast-growing
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plants, mid-growing plants, and slow-growing plants by *E. collaris* and *H. meridionalis* in

Mandena, Madagascar. Data were collected between March 2011 and January 2012 on *E*.

<i>collaris</i> , and between January and December 2013 on <i>H. meridionalis</i> .

Growth class	Variable	β	SE	95% CI	t	Р
Fast-growing						
	Fixed effects					
	Intercept	0.13	0.04	0.06, 0.21	3.42	
	Species	0.41	0.42	0.33, 0.49	9.78	<0.0001
	Activity	-0.19	0.29	-0.25, -0.14	-6.66	<0.0001
	Season	-0.02	0.03	-0.08, 0.03	-0.85	0.39
	Random effect					
	Group	Variance	0.03			
	Residual	Variance	0.15			
Mid-growing						
	Fixed effects					
	Intercept	0.29	0.02	0.24, 0.34	12.24	
	Species	-0.10	0.02	-0.15, -0.06	-4.67	<0.001
	Activity	-0.12	0.02	-0.16, -0.08	-5.47	<0.0001
	Season	-0.06	0.02	-0.10, -0.02	-2.74	<0.01
	Random effect					
	Group	Variance	0.00			
	Residual	Variance	0.11			
Slow-growing						
	Fixed effects					
	Intercept	0.55	0.04	0.48, 0.63	14.86	
	Species	-0.29	0.04	-0.36, -0.22	-8.29	<0.0001
	Activity	0.32	0.03	0.25, 0.38	9.52	<0.0001
	Season	0.08	0.03	0.02, 0.15	2.45	0.01
	Random effect					
	Group	Variance	0.00			
	Residual	Variance	0.17			

Bold indicates factors significant at P < 0.05, with values obtained using likelihood-ratio test.

328 β standardised regression coefficient, SE standard error, CI confidence interval, t t-value

329

330 Exotic species in Mandena

There were five plant species in Mandena classified as exotics, likely the consequence of

human activities and then dispersed in various ways (e.g., wind). These were broad-leaved

paperbark tree (*Melaleuca quinquenervia*), guava (*Psidium* spp.), Pemba grass (*Stenotaphrum*

dimidiatum), Polynesian arrowroot (*Tacca leontopetaloides*) and soapbush (*Clidemia hirta*). We

observed *H. meridionalis* feeding on *M. quinquenervia* flowers and resting in this species, while 335 they fed on the leaves (grass blades) of S. dimidiatum. We observed E. collaris using four exotic 336 species, feeding on the ripe fruits of C. hirta, Psidium spp., and T. leontopetaloides, and resting 337 in *M. quinquenervia*. *H. meridionalis* used exotics in 33 of 36 total months $(6.6 \pm 1.5\%)$ of plants 338 used monthly), whereas E. collaris only used exotic plant species in four of 19 total months (0.3 339 \pm 0.2% of plants used monthly). The model with the best predictive value (AIC = 105.91, γ^2 = 340 1.33, df = 1, P < 0.001) showed that exotic plants were most likely to be used by *H. meridionalis*, 341 and most often for feeding (Table 4). Season was not included in the best-fit model. 342

343

Table 4. Generalized linear mixed model predicting monthly utilisation of exotic plants. Data
 were collected between March 2011 and January 2012 on *E. collaris*, and between January and

346	December 2013 on <i>H. meridionalis</i> .	

	ionanis.				
Variable	B	SE	95% CI	Ζ	Р
Fixed effects					
Intercept	-1.52	0.74	-3.24, 0.07	-2.07	
Lemur species	3.39	0.94	1.47, 5.68	3.63	<0.01
Activity	-2.45	0.61	-3.78, -1.35	-4.05	<0.0001
Random effect					
Group	Variance	0.66			
Bold indicates factors signifi	cant at $P < 0.05$				

Bold indicates factors significant at P < 0.05.

350 **Discussion**

351 Similar to Lehman et al. (2006), we found that both *E. collaris* and *H. meridionalis* used

similar proportions of forest edge habitat within their home ranges and core areas, thus our

353 prediction that *H. meridionalis* would use greater edge habitat was not supported. As predicted,

the frugivorous *E. collaris* was more likely to use both slow- and mid-growing plant species,

355 while the folivorous *H. meridionalis* was more likely to use fast-growing plants in Mandena. In

terms of activity, slow-growing trees were particularly important for *E. collaris* resting, in line

³⁴⁸ *SE* standard error, *CI* confidence interval

³⁴⁹

with our prediction, while *H. meridionalis* used a similarly large amount of slow-growing trees 357 for resting. As expected, fast-growing plants (comprising mostly herbs and scrubs) seem to be 358 preferred by *H. meridionalis* which exhibited greater ability to include pioneer species in its diet, 359 360 a finding that is consistent with other studies of folivorous primates (Bicca-Margues and Calegaro-Marques 1994; Bonilla-Sánchez et al. 2012; Ganzhorn et al. 1999b). However, the use 361 of exotic (non-endemic) plant species for feeding by *H. meridionalis* did not support our 362 prediction, as these small-bodied folivores consumed items from these non-native plants nearly 363 every month. 364

Although bamboo lemurs are folivores, they are often considered to be dietary specialists 365 due to the large proportion of their feeding focused on bamboos (Ballhorn et al. 2016; Tan 1999). 366 However, when there are alternative habitats adjacent to a degraded habitat (e.g., mangrove 367 swamp, mono-dominant plantation), even dietary specialists can adapt and exploit them (Galat-368 Luong and Galat 2005; Grimes and Paterson 2000; Nowak 2008). Such is the case with bamboo 369 lemurs which have been observed to use alternative and/or degraded habitats (Grassi 2006; 370 371 Martinez 2008; Wright et al. 2008; Eppley et al. 2015a). Furthermore, the occasional use of wetland habitat by primates may become obligate if preferred upland habitat becomes 372 increasingly disturbed (Nowak 2008, 2013; Quinten et al. 2010); however, when species are 373 374 highly selective within their habitat, the loss of key resources may result in their ultimate demise (Lee and Hauser 1998). In contrast, low selectivity may enhance a species' chances for survival, 375 even in heavily disturbed habitats (Guo et al. 2008). 376

In general, bamboo lemurs (*Hapalemur* spp. / *Prolemur simus*) appear less susceptible to habitat degradation than more frugivorous species, i.e., *Propithecus* spp., *Eulemur* spp., *Varecia* spp. (Arrigo-Nelson 2006; Dehgan 2003; Irwin et al. 2010; Schwitzer et al. 2007). Despite this,

there appears to be some variation in bamboo lemur responses to degraded habitats. For example, 380 H. occidentalis have been observed to feed on invasive C. hirta and crop forage on rice (Orvza 381 sativa) in agricultural fields adjacent to Masoala National Park (Martinez 2008), while H. griseus 382 383 have been observed shift their diet to exotic guava (P. cattleianum) during fruiting periods in a previously selectively logged area of Ranomafana National Park (Grassi 2006). Furthermore, the 384 greater bamboo lemur (*P. simus*) is known to inhabit shaded coffee plantations (Wright et al. 385 2008). Similar to these fragment-tolerant bamboo lemurs, H. meridionalis displayed an ability to 386 adjust across various habitats (i.e., littoral forest, littoral swamp, and an invasive Melaleuca-387 dominated swamp), and though this was slightly seasonal, they were able to feed and rest for 388 large portions of time in each habitat in all seasons (Eppley et al. 2015a). Additionally, they 389 exhibited the highest dietary diversity recorded for a bamboo lemur species (Eppley et al. 390 2016a). In addition to the flexible activity pattern exhibited by *H. meridionalis* in Mandena, 391 these lemurs are also able to adjust flexibly to contrasting floristic and structural habitats, 392 exploiting resources that are specific to each environment (Eppley et al. 2015a, 2016a). 393 394 Two previous studies on *E. collaris* in Mandena indicate that these lemurs in the fragmented littoral forest tend to remain highly frugivorous but they expand their home range 395 when compared to less disturbed forests (Campera et al. 2014; Donati et al. 2011). This flexible 396 strategy differs from other brown lemur populations that seem to be able to shift seasonally to a 397 more folivorous diet (e.g., E. macaco macaco; Colquhoun 1997, E. mongoz; Curtis 2004, E. 398 rufifrons; Sussman 1977), and for a detailed meta-analysis, see Sato et al. (2016). The feeding 399 preference of *E. collaris* for mid- and slow-growing species, that tend to represent large trees 400 rather than herbs/scrubs and thus are rarer in highly fragmented areas than in pristine forest, is in 401 line with an expansion of the threshold of area requirement. Our results show a preference of E. 402

collaris for mid-growing species in the dry seasons while slow-growing, usually climax trees, are 403 selected more often in the wet season. This is an indication that E. collaris may tend to use 404 pioneer species more frequently during periods of low resource abundance (e.g., the dry season 405 406 in Mandena) when climax trees show phenological bottlenecks. This hypothesis is worth exploring in future studies matching fine-grained phenological data with lemur seasonal feeding. 407 The preference for fruiting trees does not mean that *E. collaris* is not capable of using 408 pioneer or exotic species growing in edge areas both for feeding and for resting, as indicated by 409 the similar values of edge use and their use of four exotic plant species. In Mandena, E. collaris 410 have been seen to move in the periphery of forest fragments in order to feed on fruits of the 411 exotic *Psidium* spp. (Campera et al. 2014; Donati et al. 2011) and domestic lychee (*Litchi* 412 chinensis; Donati pers. observ.). In Ste. Luce (20 km north of Mandena), E. collaris have also 413 414 been observed to move to the forest edge, or even outside of it, to feed on the fruits of exotic and/or pioneer species, e.g., the fruits of the pioneer meramaintso (Sarcolaena multiflora; 415 Campera et al. 2014). This pattern does not seem to be unusual for brown lemurs even in less 416 417 disturbed forests as migrations from familiar areas to feed on exotic *Psidium* spp. have also been recorded in E. rufifrons in Ranomafana (Overdorff 1993; Wright 1999). 418 In areas more heavily affected by habitat alteration, the genus *Eulemur* may rely heavily 419 420 on exotic trees, in most cases for fruits or for resting/sleeping. In the gallery forest fragment of

421 Berenty, during specific periods of the year the hybrids *E. rufifrons* x *E. collaris* base the

422 majority of their diet on fruits of the exotic Manilla tamarind (*Pithecellobium dulce*; Donati,

423 unpublished data). In Ampasikely, a 50-ha coastal private landholding located in northwestern

424 Madagascar, *E. macaco* feed on 23 exotic plant species that were introduced as cash crops, such

425 as coffee (Coffea spp.), papaya (Carica papaya), mango (Mangifera indica), and lebbeck or

woman's tongue (*Albizia lebbeck*; Simmen et al. 2007). Thus, the low level of reliance on exotic
species by *E. collaris* recorded in our study seems to be more the consequence of the low
frequency of suitable exotic species than the lack of flexibility of these collared brown lemurs to
include unusual food species in their diet.

Habitat disturbance may benefit folivorous lemurs in several ways. It can increase the 430 heterogeneity of a forest and therefore increase the amount or density of food resources (Oates 431 1996). Disturbance can increase the relative abundance of certain plant species that may be 432 preferred food sources, such as pioneer and light-gap species, and terrestrial herbaceous 433 vegetation (Oates 1996). Light gaps created by tree falls and/or selective felling may help to 434 maintain floristic diversity by harbouring a higher density of tree stems (Brokaw and Busing 435 2000). These gaps can also increase the number of early successional specialists, which tend to 436 have leaves with increased protein, less fibre, and lower phenolic content, as well as increasing 437 the quantity of young leaves and improving the quality of mature leaves (Chapman et al. 2002; 438 Ganzhorn 1992, 1995; Oates, 1996). Our finding that *H. meridionalis* exhibit a flexible 439 440 behavioural and feeding ecology is not all that surprising. Bamboo lemur congeners exploit bamboo, which is highly prevalent in their habitat and thrives particularly well in slightly 441 disturbed areas. The increased sunlight reaching both the canopy and forest floor further 442 increases the quantity and quality of staple foods (bamboo and leaves) and provides higher 443 quality supplemental foods (light-gap species and introduced species). Furthermore, similar to 444 our H. meridionalis results, H. griseus in Ranomafana National Park exhibit a tolerance to forest 445 edge (Lehman et al. 2006). Ultimately, the ability to use forest edge may have future benefits, in 446 that altered landscapes with habitat matrices could provide potential conservation value as vital 447 448 refuges (Chapman and Lambert 2000; Riley 2007).

449	Various folivorous primates, i.e., Alouatta spp., are able to inhabit
450	anthropogenically-disturbed habitats, likely due to a broad range of behavioural adaptations
451	(Bonilla-Sánchez et al. 2012; Zárate et al. 2014). Notably, within these habitats howler
452	monkeys are able to flexibly increase their dietary breadth (Bicca-Marques 2003), similar to
453	observations of <i>H. meridionalis</i> in Mandena (Eppley et al. 2016a). By comparison, arboreal
454	frugivores such as brown spider monkeys (Ateles hybridus) are not as flexible, and have
455	been shown to be adversely affected by the constraints of living in an anthropogenic,
456	degraded forest (Marsh et al. 2016). This is not always the case, however, as even
457	frugivorous primates, e.g., red-bellied lemurs (E. rubriventer), display an ability to utilise
458	and be tolerant of forest edge (Lehman et al. 2006). Although E. collaris and H.
459	meridionalis displayed differences in the degree of pioneer exotic plant species they used,
460	the both used similar proportions of forest edge within their home ranges and core areas.
461	The further fragmentation of remaining forests is of great concern if forest species of
462	Madagascar are to persist (Ganzhorn et al. 2014). Although the fate of all lemur species should
463	be considered precarious due to increasing habitat destruction, the knowledge that some lemurs
464	are able to cope with this degradation (to a certain degree) should be seen as positive. Some
465	primate species adapted to narrow ecological specializations may be sensitive to natural or
466	anthropogenic habitat perturbations (Harcourt et al. 2005; Kamilar and Paciulli 2008), whereas
467	others have been shown to adjust to changing environments (Anderson et al. 2007; Nowak and
468	Lee 2013). Not surprisingly, our study on two lemurids living in the highly disturbed littoral
469	forest fragments shows that the lemurs are able to use both pioneer and exotic species for feeding
470	and resting. However, while frugivorous E. collaris appear more limited by climax plants,

471	folivorous <i>H. meridionalis</i> show a wider range of adaptability, probably favoured by its diet and
472	smaller body size.
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