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1 SPECIAL ISSUE ARTICLE

- 3 Living with hyraxes: Biogeography and comparative ecology of West African
- 4 Fornasinius beetles
- 5

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- 36 Hyrax and beetles

37 ABSTRACT

38 Among the flower beetles (Scarabaeidae, Cetoniinae), the Goliathini comprise several genera of 39 medium and large-sized beetles widely distributed in sub-Saharan Africa. In this tribe, the genus Fornasinius 40 Bertoloni, 1853 includes two species found in West Africa: F. higginsi (Westwood) and F. klingbeili Zöller, 41 Fiebig, & Schulze. In this study, we present new data on the comparative ecology of these two species, 42 including sex ratio, population structure by size, monthly activity patterns, and habitat features at two 43 different spatial scales. These observations were conducted over a twenty-year period in Cote d'Ivoire, 44 Ghana, and Togo. Both species exhibited similar overall population structure and morphometrics. They 45 were characterized by a male-skewed adult sex ratio (with possible bias), male-larger sexual size 46 dimorphism, and consistent population structure by size; the two species are seemingly parapatric (with 47 the potential contact zone being separated in eastern Ghana by the Volta river and lake). Both are primarily 48 found in the vicinity of trees that contained middens of hyraxes, such as Dendrohyrax dorsalis (F. higginsi), 49 D. interfluvialis (F. klingbeili), and Procavia capensis (F. klingbeili). Fornasinius higginsi has been primarily 50 observed in forest habitat, while F. klingbeili was found in both forests and Guinea savannahs. Occurrence 51 sites of K. higginsi were predominantly characterized by higher tree cover, less bare areas and built-up 52 zones. Although these beetles may be locally abundant within hyrax middens, they typically occur in only a 53 relatively small number of sites within their preferred habitat. Monthly activity patterns differed between 54 the two species. Adults of both species were observed throughout the year, sightings of *F. higginsi* were 55 more common during the wet season, whereas F. klingbeili sightings were more frequent in the dry season. 56 The observed differences in monthly activity patterns between these ecologically and morphologically 57 similar species may help minimize interspecific competition in potential – yet unknown – sites of syntopy. 58 Although some Fornasinius populations might be locally threatened by deforestation or overhunting of 59 hyraxes, the careful management of hyrax populations should adequately ensure the conservation of these 60 two beetle species.

61 Keywords: Africa; Ecology; Goliathiinae; Habitat; Interspecific competition; Population structure;
62 Seasonality

63

64 **1** | **INTRODUCTION**

65 The Goliath beetles in the genus Fornasinius Bertoloni, 1853 are a group of about seven relatively large-66 sized species (up to 68 mm long, including the clypeal apophysis) with distinctive morphology and large 67 cephalic processes in the males. While previously included by some authors in genus Goliathus Lamarck, 68 1801 as a subgenus (Weibes 1968; Croizat, 1994; Maquart & Malec, 2017), Fornasinius is currently treated 69 as a valid genus (De Palma, 2018). All Fornasinius species are found in sub-Saharan Africa (De Palma & Di 70 Gennaro, 2017; De Palma & Takano, 2018; De Palma, 2018) and are heavily targeted in the international 71 entomological trade. Despite this, these beetles have received little ecological study, with only scattered 72 natural history data available (e.g., Maguart & Malec, 2017).

In this study, we utilize data gathered opportunistically over 20 years in Cote d'Ivoire, Ghana, and
Togo, to provide information on the ecology of two apparently rare *Fornasinius* species: *F. higginsi* and *F. klingbeili*. We focus on monthly activity patterns, sex-ratio, population structure by size, and habitat
characteristics/partitioning. These two species are morphologically similar (De Palma, 2018), and, despite
having received attention from entomological collectors, have been extremely rare in collections (e.g.,
Lachaume, 1983) until the recent discovery of their association with hyrax middens.

The two West African *Fornasinius* species are entirely black, although sparse yellow cretaceous spots are often present on the elytra (Figure 1). *Fornasinius higginsi* males are readily distinguished from those of *F. klingbeili* by the very distinctive shape of the clypeal plate and clypeal horn; females display only minor differences in the anterior margin of the clypeus. Both species differ significantly from other members of the genus and arguably could be included in the now defunct genus *Sphyrorrhina* Nickerl, 1890 as a subgenus of *Fornasinius*.

In West Africa, the two *Fornasinius* species have a nearly continuous and parapatric distribution; *Fornasinius higginsi* recorded from western Cote d'Ivoire (Fediere et al., 1987) eastwards to the Volta
region in Ghana, while *F. klingbeili* is found in Togo westwards to the Dahomey gap in Ghana (Maquart &

Malec, 2017). *Fornasinius aureosparsus* (van de Pool, 1890), a distantly related species, is found in southeastern Nigeria, Cameroon and the Congo basin (De Palma, 2018). Given their parapatric distribution and similar habits, *F. higginsi* and *F. klingbeili* may serve as an interesting case study to examine patterns of coexistence and competition among Goliathine beetles in West African forest and savannah habitats.

92 In this paper, we aim to address the following key questions: (1) Given that the two West African 93 Fornasinius species exhibit remarkable morphological similarity and are genetically related (M. De Palma, 94 unpublished data), what ecological and/or biogeographic mechanisms have led to the separation of these 95 species in otherwise analogous ecological niches? (2) Do potential differences in their ecological niches 96 involve variations in annual phenology and/or habitat preferences? If so, are these differences observed at 97 the site scale and/or the landscape scale? (3) Are the two species linked to middens of different hyrax 98 species or do they occur in middens of a same hyrax species? (4) What are the conservation implications of 99 our findings?

100

101 2 | MATERIALS AND METHODS

102 2.1 | Field protocol

All numerical data were obtained opportunistically between 2012 and 2022, during other fieldwork in Togo, Ghana, and Cote d'Ivoire (Figure 2). Qualitative and distribution records were gathered from 2000 to 2024. Beetles were recorded during surveys investigating aspects of the conservation and ecology of other animals, primarily reptiles.

107 The geographic coordinates of all observed beetles were recorded; however, these are not provided 108 here for conservation reasons. This precaution is necessary because these beetles are heavily sought after 109 by collectors for the entomological market trade.

We spent a total of 68 field days in sites in Ghana and Cote d'Ivoire (range of *F. higginsi*) and 609 field days in Togo (range of *F. klingbeili*). The number of field days per month in each country is given in Appendix Table A1.

We surveyed beetles across a wide range of potential habitats, including dry savannahs, wet rainforests, and mangroves. At each study site, we conducted random surveys from 06:00 to 24:00 local time, primarily targeting reptiles and amphibians but also searching for beetles. We ensured that similar amounts of time were spent in different habitats and under various weather conditions, from full sun to slight rain. Transects were walked concurrently by at least three individuals moving independently of each other.

In addition to recording live beetles, we documented individuals found dead on the ground. When an individual was found, we assessed the characteristics of the microhabitat and noted the presence of nearby hyrax groups. Hyrax middens in tree holes are often located at heights over 5 meters (our unpublished observations), making them difficult to examine directly. Therefore, we focused on checking the surrounding area for *Fornasinius* specimens or their remains on the ground.

124 The following habitat categories were considered for each observed individual:

i) DRS = dry (= Sudanese) savannah, characterized by a large dominance of drier grasslands
 interspersed with *Acacia* bushlands, and the Combretaceae and Caesalpinioideae trees. The
 herbaceous belt consists of mostly species of the genera *Andropogon* and *Hyparrhenia*, but on
 shallow soils also of *Loudetia* and *Aristida*.

WES = wet (Guinean) savannah, that consists mainly of grasslands crossed with abundant trees
 and gallery forest strips alongside streams and on hillsides. *Lophira lanceolata* is among the
 dominant tree species found in this habitat type.

132 iii) DRF = semi-deciduous forest, that is characterized by some tree species exhibiting a partial
 133 shedding of leaves in the dry season. These forests are very diverse in terms of plant species

- and include the coexistence between humid forest species transgressing towards North and dry
 forest species transgressing towards South.
- iv) PLT = plantation, mainly of cacao, coffee, banana, plantain, pineapple, cassava, yam, etc. These
 plantations are normally interspersed with few large trees and human settlements and are very
 disturbed and altered forest-derived habitats.
- v) FTP = forest-plantation mosaic, is a mixed landscape with portions of territory similar to DRF
 and others to PLT.
- vi) MAF = mature (and pristine) rainforest, consisting of patches of forests that have not been
 altered by humans in the last thirty years, with many trees more than 30 m height. These
 patches are typically sacred grooves, inner parts of community forests or legally protected
 areas.
- vii) SEF = secondary or altered moist forest. This is typically the same vegetation type as MAF, but
 with a stronger/more recent human intervention. The number/density of very large trees
 (taller than 30 m height) is much smaller than in MAF, the average tree height is about 12-15
 meters, and the surface is often crossed by paths and roads.

149 2.2 Sample sizes

150 None of the live specimens encountered were collected or sacrificed. Only those specimens that 151 were found dead in the field were retrieved and saved for further analyses. In the following text, we define an "individual record" as the event observed during quantitative transects in which Fornasinius specimens 152 153 were observed at a single point and at a single time (date and hour-of-the-day). Several individual records 154 consisted of more than one specimen, including both live and remains of dead animals. However, all 155 specimens found in the same locality and at the same time were considered as single records to avoid 156 pseudoreplication in the analyses of activity patterns and habitat (at both spatial scales). In quantitative 157 transects, we collected 77 individual records of F. higginsi and 39 individual records of F. klingbeili.

158 The 77 individual records of F. higginsi involved 105 measurable and sexually identifiable 159 specimens, while the 39 individual records of F. klingbeili involved 44 measurable and sexually identifiable 160 specimens. In the seasonality analyses, we included all individual records. For habitat analyses of F. higginsi, 161 we excluded 14 out of 77 individual records because these involved individuals found squashed on roads or 162 dead in ponds, making it impossible to define a clear habitat. The sample size for habitat analyses of F. 163 klingbeili included a total of 50, since 11 additional individual records with habitat information were obtained outside the quantitative transect surveys. In addition to transect data, the biogeography analysis 164 165 also incorporated data collected by one of the authors (M. De Palma) over more than two decades of 166 research into African Goliath beetles.

167 The total body length of each beetle collected was measured from the tip of the clypeus to the 168 apex of elytra. It is important to note that the total body length includes the cephalic horn present in male 169 specimens. Therefore, a greater body length does not imply a greater body mass.

When analysing the monthly activity patterns, we only considered specimens encountered during
random walks. We excluded individuals found dead and in small fragments, as it was not possible to
determine the month(s) of their activity and death. However, we included some specimens from collections
with reliable tags indicating the precise date of collection. All voucher specimens are deposited in the
Entomological Collections of the Université de Lomé (Togo), Luiselli's collection in Rome (Italy), and De
Palma's collection in Lausanne (Switzerland).

176 2.3 | Statistical analyses

177 Monthly activity patterns of adult beetles was analysed relative to monthly field effort. We first 178 calculated the monthly relative sampling effort (MRSE) for each of the two species separately. We used a 179 null hypothesis of equal encounter frequency among months. The MRSE was calculated using the following 180 formula:

[1] MRSE = $n \times 100 / N$

181

where *n* represents the number of field days spent in that month in the study area, and *N* represents the total number of field days in that area. We then generated the expected number of beetles in each month by multiplying the total number of beetles of each species found during the entire research period by MRSE. Finally, we compared the observed number of beetles per month against the expected number using an observed-versus-expected χ^2 test. The same statistical design was applied in other field studies of beetles (see Dendi et al., 2023).

188 To evaluate the correlation between the activity of adult beetles and rainfall patterns, we used the 189 number of rainy days per month as a proxy for rainfall. Rainfall data was obtained from Accra, Ghana 190 (downloaded from <https://weatherspark.com/y/42322/Average-Weather-in-Accra-Ghana-Year-191 Round#Sections-Precipitation>, last accessed on 20 May 2022) for F. higginsi, and from Kara, Togo 192 (<https://weatherspark.com/y/45806/Average-Weather-in-Kara-Togo-Year-Round#Sections-Precipitation>, last accessed on 20 May 2022) for F. klingbeili. We conducted a Spearman's rank correlation (r_s) test for 193 194 each month, with the number of rainy days as the independent variable and percentage monthly observed 195 adults as the dependent variable.

To evaluate habitat preferences of both species, we used two different approaches across two spatial scales: (i) the sighting site scale, and (ii) the landscape scale. For habitat preference at the sighting site scale, each beetle record was assigned to the dominant (or the most representative) of the seven habitat type categories described above, within a 200 m buffer around the capture site of each specimen. We assessed differences in sighting frequencies between habitats using observed-versus-expected χ^2 tests. Additionally, we calculated the habitat niche breadth for numerical data using Simpson's (1949) measure of niche breadth (B_s).

To analyze habitat preferences at the landscape scale, we utilized a GIS-based methodology with QGIS software version 3.32. For this analysis, we employed the LC Map of Africa 2016 raster layer, released by the European Space Agency (ESA) between October 2, 2017, and February 6, 2018, with a pixel size of 20 meters. The raster classifies land use into the following categories: 1 - Tree cover areas (TRC); 2 - Shrub

207 cover areas (SHR); 3 - Grassland (GRS); 4 - Cropland (CRP); 5 - Aquatic or regularly flooded vegetation
208 (VGA); 6 - Sparse vegetation (SPA); 7 - Bare areas (BAR); 8 - Built-up areas (BLT).

For each occurrence point, we created a square buffer area of 2.2 km per side. We used the "saga_histogram" algorithm to perform zonal statistics, which calculates the counts of each unique value from the raster layer within defined zones. This allowed us to count the number of pixels for each land use category at each beetle occurrence point and determine the percentage of occupancy for each land use category. To assess interspecific differences in the mean percentage occupancy of various land use categories, we employed a non-parametric Kruskal-Wallis ANOVA.

The sex ratio was evaluated using an observed-versus-expected χ^2 test. Intersexual differences in mean body sizes were assessed using a one-way ANOVA followed by Tukey HSD post-hoc tests for pairwise comparisons. We tested all variables for normality and homoscedasticity using a Shapiro-Wilk test, and if necessary, we log-transformed them before conducting any parametric analysis. All analyses were performed using PAST version 4.0 software, with all tests being two-tailed and alpha set at 5%.

220

221 **3 | RESULTS**

222 3.1 | Sex ratio and body length

Out of the total of 105 individuals of *F. higginsi* for which sex and body size were recorded, the observed sex ratio (males: females) was 1.92: 1 ($\chi^2 = 11.11$, df = 1, P < 0.001). Data on body length and sex were also collected on 44 individuals of *F. klingbeili*, with a sex ratio of 1.44: 1, which was not significantly uneven (χ^2 = 1.45, df = 1, P = 0.228). The distribution of body lengths in the observed sample of the two species is shown in Appendix Figure A1.

The distribution of body length in the observed sample of the two species is given in Appendix
Figure A1. In both species, there was a significant male-larger sexual size dimorphism, driven by the
presence of the cephalic horn in the males (Figure A1; *F. higginsi* – males: 53.1 ± 4.7 mm, median = 54 mm,

n = 69; females: 45.9 ± 3.3 mm, median = 46 mm, n = 36; *F. klingbeili* – males: 52.3 ± 6.1 mm, median =
51.5, n = 25; females: 48.1 ± 5.7 mm, median = 47 mm, n = 19. A) (after log-transformation, t = 8.33, P <
0.0001 for *F. higginsi* and t = 2.33, P < 0.05 for *F. klingbeili*). A one-way ANOVA (on log-transformed body
size measurements) followed by Tukey HSD post-hoc test revealed that male size did not differ significantly
between species (Tukey = 1.18, P = 0.836), and the same was true for the female size (Tukey = 2.233, P =
0.394).

3.2 | Apparent abundance and **Seasonality**

238 Considering only the individuals encountered during our transect walks, we recorded 63 individuals 239 of *F. higginsi* and 50 individuals of *K. klingbeili*. The number of individuals observed relative to field effort 240 was higher in *F. higginsi* (0.93 individuals per field day in appropriate habitat) than in *F. klingbeili* (0.07 241 individuals per field day.

Adults of both species were observed throughout most of the year. However, *F. higginsi* was not sighted in September-October, and *F. klingbeili* was not sighted in September (Figure A2). The frequency of monthly sightings was significantly uneven in *F. higginsi* ($\chi^2 = 62.52$, df = 11, P < 0.0001), with peaks higher than expected in March, April, and May (wet season) and lower than expected in September to December (the last phase of the wet season and the dry season; Figure 3).

Similarly, the frequency of sightings of *F. klingbeili* was also significantly uneven ($\chi^2 = 56.73$, df = 11, P < 0.0001), with peaks higher than expected in November, December, and January (dry season) and negative peaks in June to September (wet season; Figure 3). There was no correlation between monthly sighting frequency of and the number of rainy days per month in *F. higginsi* (r_s = 0.049, P = 0.879). However, the activity of *F. klingbeili* was inversely correlated to the monthly number of rainy days (r_s = -0.633, P < 0.05). Monthly activity patterns differed significantly between species (χ^2 = 48.1, df = 10, P < 0.0001).

253

254 3.3 | Habitat

255 *3.2.1.* Habitat at sighting site scale

256	Although the two species occurred in various habitats, particularly in SEF (Figure 4), there were
257	significant interspecific differences (χ^2 = 25.6, df = 4, P < 0.0001). Fornasinius higginsi was observed more
258	frequently in MAF, while F. klingbeili was more commonly found in DRS (Figure 4). Habitat niche breadth
259	was much broader in <i>F. klingbeili</i> compared to <i>F. higginsi</i> , with respective values of 4.596 and 2.733 (B_s).
260	Except for a few individuals found dead along roads and paths, most individuals from both species
261	(including preyed specimens) were discovered under hollow shafts with middens of Dendrohyrax
262	interfluvialis (F. klingbeili) and of D. dorsalis (F. higginsi), particularly Dialium guineense trees. However, F.
263	klingbeili was also found in rocky sites where there were Procavia capensis kerstingi middens, particularly
264	in the DRS habitat. Height above ground of tree cavities where hyraxes made their middens (and therefore
265	used by these beetles) ranged from 5 to 13 m. This made it impossible for us to examine the different
266	hollows. However, we identified seven trees regularly used by hyrax in Cote d'Ivoire, 11 in Ghana, and 32 in
267	Togo. In these, we found remains of <i>F. higginsi</i> in the immediate vicinity of six trees in Cote d'Ivoire
268	(85.7%), nine in Ghana (81.8%), while <i>F. klingbeili</i> were found nearby nine of the surveyed trees in Togo
269	(28.1%).

270

271 3.2.2. Habitat at landscape scale

272 Land use categories for each beetle occurrence site of the two Fornasinius species at the landscape 273 scale are shown in Table 1. The two species differed significantly in terms of the percent coverage of the 274 various land use categories in their respective presence sites (Kruskal Wallis ANOVA, P < 0.0001), with 275 significant differences being for: (i) tree cover representing a much higher percentage of land use in F. 276 higginsi occurrence sites and (ii) bare areas and built up areas appearing with significantly higher 277 percentages in *F. klingbeili* occurrence sites (Figure 5):

12

279 **4** | **DISCUSSION**

280 4.1 General considerations

281 The two species of *Fornasinius* beetles examined in this study showed both similarities and 282 significant statistical differences in their ecological traits. These similarities likely stem from conservative 283 traits associated with the evolutionary history of Goliathiinae beetles. These traits include aspects of 284 population structure, such as similar patterns in body size distribution among individuals, as well as 285 morphometrics, specifically the larger size in terms of total length of males compared to females (sexual-286 size dimorphism). In other genera of the Goliathini, males also tend to be larger than females primarily due 287 to their hypertrophic cephalic processes (e.g. genus Dicronorhina, Lekkerkerk & Krikken, 1986 and De 288 Palma, 2011; genus Goliathus, Jiang et al., 2012; genus Mecynorhina, Christiansen, 2013 and De Palma et 289 al., 2024; genus Eudicella, Vendl et al., 2018) These processes serve as signals of a mating system based on 290 male-male combat for access to females (Björkman et al., 2009). Sexual selection likely contributes to the 291 increase in male body size, as larger body size and success in sexual combat are often correlated in larger 292 animals (Shine, 1978, 1994; Berry & Shine, 1980).

293 We observed that males outnumbered females, although this difference was statistically significant 294 only in *F. higginsi*. A skewed operational sex ratio may be associated with intense sexual competition 295 among these beetles. This could be because females are a limited resource, confined to a relatively small 296 area where many males are actively competing. However, our estimates of sex ratios may be biased by our 297 sampling modality. In fact, we have considered only the specimens encountered along the transects, in 298 which the sex-ratio may have been unbalanced to favor males due to their lower philopatry and habit to 299 flutter and disperse in search of females with which to mate. Conversely, females are more philopatric and 300 may remain in the nests of hyraxes rather than disperse. In support of this hypothesis, the data collected by 301 one of us (M. De Palma) inside a hyrax midden in Ghana indicate that the majority of the specimens were 302 females.

304 We frequently observed these beetles under, or in the vicinities of, trees that housed hyrax 305 middens, but they were rarely found in areas without hyrax. This suggests that the beetles are confined to 306 small areas within dense forest habitats, unlike other Goliathiinae species that are more widely distributed. 307 Because we couldn't access the various tree cavities with hyrax middens, our survey likely underestimated 308 the abundance of these beetle species. We did, however, find remains of *F. higginsi* near almost all (>90%) 309 of the trees used by Dendrohyrax interfluvialis, indicating that this species is widespread wherever the 310 hyraxes accumulate their middens. Local hunters in eastern Ghana also suggest that multiple Fornasinius 311 specimens, both adults and larvae, may inhabit each tree cavity used by hyrax individuals for their middens. 312 So, although their density may be locally high, they are confined to small areas within the appropriate 313 habitat. The percentage of trees with hyraxes that also have F. klingbeili was significantly lower (<30%), 314 suggesting that this species is either less common than F. higginsi or less frequently associated with hyrax 315 middens. Currently, it is unknown whether Fornasinius species are obligate hosts of hyrax species, as we 316 found them in some sites apparently without any hyrax populations. However, the presence of hyraxes is 317 often associated with the presence of these beetles. Nor can it be ruled out that these beetles may also 318 occasionally associate with the latrines of other forest mammals, and that this explains the reason for the 319 observation of individuals in sites without the apparent presence of hyraxes. For instance, we observed in 320 southern Nigeria two adults of the closely-related Fornasinius aureosparsus at least than 1 m from a African 321 civet's (Civettictis civetta) latrine, suggesting that they were just dispersing out of it. However, the storage 322 time of the latrines is essential for the maintenance of a stable population of beetles. Therefore, it can be 323 hypothesized that the latrines of any other species can only be used by *Fornasinius* species occasionally and 324 in suboptimal environmental conditions.

The use of hyrax middens by *Fornasinius* beetles is similar to that of other cetonids that use bird nests for larval development (Choi et al., 2018; Zbyryt & Oleksa, 2018), and that may be even obligate hosts of their avian nests (Zbyryt & Oleksa, 2018). Tropical cetoniinae may also develop in termite nests (Touroult & Le Gall, 2013), but we have not observed any *Fornasinius* specimen nearby termite nests.

329 Hyrax species, both Dendrohyrax spp. and Procavia capensis, are scattered throughout West Africa, 330 which influences the distribution of Fornasinius species. Further investigation of hyrax middens may reveal 331 whether the association between these beetles and mammals is obligatory or facultative. The occurrence 332 of F. klingbeili in Dendrohyrax middens was previously reported by Maquart & Malec (2017). It's worth 333 noting that Hegemus pluto and Argyrophegges kolbei from East Africa also use the middens of Heterohyrax 334 brucei and Dendrohyrax dorsalis, respectively (Di Gennaro, 2014), indicating that these two Fornasinius species are not unique among goliathini beetles in having their life cycle linked to hyrax middens. 335

336

337

4.3 Habitat and biogeography implications

338 Our study documented that F. klingbeili occurs in dry open forests in the Kara region in central 339 Togo, as well as in dense deciduous forests in the Togo hills and in the lowland dry forest in the Togodo 340 area. Thus, it is apparently a habitat generalist, as suggested by the relatively high value of its habitat niche 341 breadth. Conversely, F. higginsi shows less generalist behaviour in habitat use, typically dwelling in dense 342 forests. These patterns are clearly confirmed also at the landscape scale: the occurrence sites of K. higginsi 343 were characterized by a higher percentage of tree cover areas and less of bare areas and built-up areas. 344 The distribution of *F. higginsi* closely follows that of *Dendrohyrax dorsalis* in Cote d'Ivoire and that of *F.* 345 klingbeili follows the distribution of Dendrohyrax interfluvialis in Togo and even in eastern Ghana as it 346 seems to be present only east of the Volta lake and the former Volta river (Oates et al., 2022). Like other 347 species of the same genus (Milner & Harris, 1999), the two Dendrohyrax species are moist evergreen and 348 semi-deciduous forest dwellers (Jones, 1978; Djossa et al., 2012). Therefore, the density of the two 349 Fornasinius species is likely to be higher in moist evergreen and semideciduous forests than in other habitat 350 types. These two types of forests are also those with higher Cetoniinae species diversity in Ghana (Mudge 351 et al., 2012). Given that D. interfluvialis occurs between the Volta and Niger Rivers, including the south of 352 the Republic of Benin throughout the "Dahomey Gap" (sensu Salzmann and Hoelzmann, 2005) and even close to Benin City in western Nigeria (Oates et al., 2022), and that F. klingbeili has already been recorded 353

from relatively open dry forest-savannah mosaics in Togo (Maquart & Malec, 2017; this study), we predict that *F. klingbeili* may also occur in Benin and western Nigeria, where its presence has not been reported so far. At the same time, it is very likely that *F. higginsi* is widespread also in Liberia and eastern Sierra Leone, in the regions where the forest cover is still adequate.

358 Based on the above-considerations, the distribution of *F. higginsi* should be very similar to the 359 classic pattern of the West African species of the Upper Guinean Forest block (Oates et al., 2004; Mallon et 360 al., 2015), while that of F. klingbeili would not be limited to the Dahomey Gap but could perhaps also 361 penetrate the humid forests of the Lower Guinean forest block, west of the Niger River. From a 362 biogeographical point of view, all Fornasinius species have, in Africa, a parapatric distribution. In this sense, 363 the two species studied are no exception. Unfortunately, there is no data on the separation zones between 364 the ranges of the two target species in eastern Ghana. These areas have been enormously modified by the 365 creation of Lake Volta, the largest artificial lake in the world, and the natural landscape has changed so 366 much that it is impossible nowadays to reconstruct the dynamics of their past distribution on a fine scale.

367

368 **4.4. Seasonal activity patterns**

369 Both Fornasinius species were year-round active. Although adult sightings occurred almost every 370 month, a more in-depth analysis, controlling for field effort, revealed substantial interspecific differences: 371 F. higginsi was more frequently encountered on the ground during the wet season, and F. klingbeili during 372 the dry season. We hypothesize that the individuals found on the ground are those that disperse in search 373 of new mating sites, and therefore, their monthly frequency of sightings may correlate with their above-374 ground activity intensity. Since these beetles are found year-round inside the hyrax middens (tree cavities 375 often placed at over 10 m from the ground), we cannot exclude the possibility that the number of 376 individuals found on the ground (often just recently dead) may not directly correlate with their activity 377 intensity inside the tree cavities.

378 Niche theory cannot explain the different activity patterns we uncovered in the two species. In fact, 379 the nearest presence localities for the two species (Kwahu escarpment as the easternmost site for F. 380 higginsi and Hohoe and the Kadjebi district as the westernmost sites for K. klingbeili) are about 155 km 381 apart, now separated by Lake Volta. Therefore, it seems that the former Volta river and the westernmost 382 part of the Dahomey Gap separated the ranges of these two species. However, in the gap area between the 383 two Fornasinius, both Dendrohyrax interfluvialis and Procavia capensis are known to occur (Kingdon et al., 2013; Oates et al., 2022). Therefore, we cannot exclude the possibility that the ranges of the two species 384 385 may be closer than they appear, or even be sympatric in a few sites. As regards the interspecific 386 relationships between these two species, there are no data available. However, their distribution can be 387 predicted to be strictly parapatric due to highly competitive potential occurring between the two species. 388 In fact, both have a very peculiar niche (inhabiting hyrax middens) and hyrax populations are scattered in 389 the environment, probably providing a limited resource to these beetles. Nonetheless, it would be 390 interesting to verify whether, in the areas where both Dendrohyrax interfluvialis and Procavia capensis do 391 co-occur, also both *Fornasinius* species may coexist, perhaps by partitioning the niche using each a given 392 species of hyrax. However, this hypothesis remains purely speculative.

393

394 **4.5 Conservation implications**

395 According to our observations, we conclude that the survival of these mammals will protect 396 Fornasinius populations because their life cycle is closely linked to three hyrax species, D. interfluvialis, D. 397 dorsalis, and P. capensis. Although opportunistically eaten by people, these hyrax species are not easily 398 found by human hunters and are not among the main prey species hunted for meat (Oates et al., 2022). 399 This is likely due to their cryptic, mainly nocturnal and arboreal habits, as well as their solitary foraging 400 activities (Djossa et al., 2012). Furthermore, Dendrohyrax spp. and Procavia capensis are not threatened in 401 Togo (Amori et al., 2016) or in West Africa in general (Oates et al., 2022), suggesting that neither of the two 402 Fornasinius species are currently facing serious threats. Additionally, these beetles are not collected by

403 local communities due to their low cultural and/or subsistence value. While the international entomological 404 market may exploit the wild populations of both species (which are sold at a relatively high price in the 405 northern world), it is unlikely to pose a serious threat to these species at the global scale. However, the loss 406 of forest habitat, particularly the cutting of hollow shafts used by hyrax individuals, may be a local threat as 407 it could affect their mammalian "hosts" (Oates et al., 2004; Poorter et al., 2004; Critical Ecosystem 408 Partnership Fund, 2015). Therefore, habitat protection is crucial for the conservation of these beetle species. Overall, although some Fornasinius populations might be locally threatened by deforestation or 409 410 perhaps overhunting of hyraxes, we conclude that for both species a red listing of LC/NT should be 411 adequate.

412

413 **4.5 Limitations of the study and future steps of the research**

414 The present study, although consisting of a sample size sufficient to conduct robust statistical 415 analyses, is certainly limited by the fact that the research was carried out in only a few presence sites 416 within the ranges of the two Fornasinius species. This was because, while Fornasinius species were common at sites of occurrence, it was nevertheless difficult to find new locations of occurrence, especially 417 418 for F. klingbeili. For many of the statistical analyses carried out by us, considering the number of individuals 419 observed in the same site as independent would have caused biases due to pseudoreplication (Heffner et 420 al., 1996; Chaves & Chaves, 2010) given that the various individuals probably all came from a single source 421 (a certain hyrax midden within a specific hollow tree). Therefore, the significance of our conclusions may be 422 affected by the small number of surveyed sites.

Regarding future studies, it will be necessary to verify the generality of our observations with a larger sample of presence locations. Additionally, exploring the ecological relationships between hyraxes and *Fornasinius* beetles in more detail will be appropriate, for example, by using technologies such as camera traps placed near hyrax middens. These future studies will have to verify whether the cohabitation

between hyraxes and *Fornasinius* beetles is obligated, and above all if, as it seems at the current state of
our research, each species of *Fornasinus* is linked to a specific species of *Dendrohyrax*.

429	Increasing collaboration between mammalogists and entomologists is essential to better study the
430	ecology and conservation of both hyraxes and Fornasinius beetles. Both are subject to the same type of
431	threat (habitat loss) but also face species-specific threats (overhunting for subsistence in the case of
432	hyraxes and harvesting for the international entomological market in the case of beetles). Finally, a formal
433	"threats analysis" approach, as recently performed on threatened turtles (Luiselli et al., 2024a, 2024b),
434	would be extremely important for defining future management and conservation planning.

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444	
445	
446	CONFLICT OF INTEREST
447	The authors declare no conflict of interest.
448	
449	DATA AVAILABILITY STATEMENT

450 The data that supports the findings of this study are available on request from the corresponding 451 author. The data are not publicly available due to privacy or ethical restrictions.

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TABLE 1. Summary of the habitat features of each occurrence site of the two Fornasinius species at the
landscape scale. Symbols: Tree cover areas (TRC); Shrubs cover areas (SHR); Grassland (GRS); Cropland
(CRP); Vegetation aquatic or regularly flooded (VGA); Sparse vegetation (SPA); 7 - Bare areas (BAR); Built up
areas (BLT). Geographic coordinates are not given for conservation reasons.

Genus	Species	Country	Locality	TRC	SHR	GRS	CRP	VGA	SPA	BAR	BLT
Fornasinius	higginsi	Ghana	Ankasa Forest reserve	99.30%	0.01%	0.00%	0.69%	0.00%	0.00%	0.00%	0.00%
Fornasinius	higginsi	Ghana	Kwahu	86.35%	0.04%	4.04%	9.41%	0.00%	0.00%	0.00%	0.00%
Fornasinius	higginsi	Ghana	Kwahu plateau	76.10%	0.00%	4.09%	19.80%	0.00%	0.00%	0.00%	0.00%
Fornasinius	higginsi	Ghana	Kyebi	52.42%	0.00%	2.58%	36.99%	3.63%	0.00%	0.00%	4.09%
Fornasinius	higginsi	Ghana	Kyebi	80.35%	0.00%	0.50%	16.48%	1.14%	0.00%	0.00%	1.41%
Fornasinius	higginsi	Cote d'Ivoire	Tai Forest	99.33%	0.00%	0.00%	0.67%	0.00%	0.00%	0.00%	0.00%
Fornasinius	higginsi	Cote d'Ivoire	Tai Forest	100.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%	0.00%
Fornasinius	higginsi	Cote d'Ivoire	Banco	54.00%	0.00%	0.00%	15.81%	25.05%	0.00%	0.00%	2.76%
Fornasinius	higginsi	Cote d'Ivoire	Danané	86.84%	0.00%	2.41%	10.73%	0.03%	0.00%	0.00%	0.00%
Fornasinius	klingbeili	Тодо	Badou	74.74%	0.16%	3.60%	6.36%	0.00%	0.00%	0.00%	15.10%
Fornasinius	klingbeili	Тодо	Kara	5.63%	1.60%	3.40%	11.53%	0.00%	0.00%	0.06%	77.79%
Fornasinius	klingbeili	Тодо	Togodo forest	13.08%	2.66%	83.44%	0.83%	0.00%	0.00%	0.00%	0.00%
Fornasinius	klingbeili	Togo	Bénali	91.42%	1.58%	0.64%	6.36%	0.00%	0.00%	0.00%	0.00%
Fornasinius	klingbeili	Тодо	Kpété Béna	91.42%	1.58%	0.64%	6.36%	0.00%	0.00%	0.00%	0.00%
Fornasinius	klingbeili	Тодо	Klouto	82.25%	1.56%	2.08%	13.51%	0.00%	0.00%	0.00%	0.61%
L	1	I		1						[1

- **FIGURE 1.** The two study species: *Fornasinius higginsi* (a) female, b) male) from Banco forest, Cote d'Ivoire,
- 553 Fornasinius klingbeili (c) female, d) male) from Missahohé forest, Togo



- **FIGURE 2** Map of Cote d'Ivoire, Ghana and Togo showing the sites of quantitative transects of *Fornasinius*
- *higginsi* and *F. klingbeili* during the present study, and the landuse of the region.



- 560 FIGURE 3. Monthly activity patterns of Fornasinius higginsi and F. klingbeili expressed as number of
- individuals observed during the study transects versus those expected on the basis of the relative monthly
- 562 field effort. For statistical details, see the text

Fornasinius higginsi (n = 77)



Fornasinius klingbeili (n = 39)



563

FIGURE 4. Distribution of the two *Fornasinius* species across habitat types at the sighting site scale.

566 Symbols: DRS = dry savannah, WES = wet savannah, DRF = dry forest, PLT = plantation, FTP = forest-

567 plantation mosaic, MAF = mature (and pristine) forest, SEF = wet secondary or altered forest.



FIGURE 5. Means and Standard Deviation of the land use categories of each occurrence site of the two *Fornasinius* species at the landscape scale. Symbols: Tree cover areas (TRC); Shrubs cover areas (SHR);
Grassland (GRS); Cropland (CRP); Vegetation aquatic or regularly flooded (VGA); Sparse vegetation (SPA); 7
Bare areas (BAR); Built up areas (BLT).



Fornasinius klingbeili



Land use category

574

APPENDIX

TABLE A1 Field effort (number of days) per month, spent by the team within the range of the two study

578 species.

	Ghana + Cote d'Ivoire	Тодо	TOTAL
	F. higginsi	F. klingbeili	
January	5	40	45
February	4	42	46
March	6	44	50
April	12	47	59
May	3	55	58
June	1	54	55
July	5	46	51
August	7	53	60
September	9	61	70
October	6	67	73
November	5	48	53
December	5	52	57
TOTAL	68	609	677

- **FIGURE A1.** Population structure-by-size in the two *Fornasinius* species studied here. For statistical details,
- see the text. Yellow = females; purple = males.

Fornasinius higginsi – males = 69, females = 36



Fornasinius klingbeili – males = 25, females = 19



583

FIGURE A2. Monthly activity of *Fornasinius higginsi* and *F. klingbeili* (expressed as percentage of observed 586 individuals per month with Standard Error). For statistical details, see the text

