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1	Physiological and Behavioural Effects of Fruit Toxins					
2	on Seed-Predating versus Seed-Dispersing Congeneric Rodents					
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22 Summary

23 Fleshy, ripe fruits attract seed dispersers but also seed predators. Although many fruit consumers (legitimate seed dispersers as well as seed predators) are clearly exposed to 24 plant secondary compounds (PSCs), their impact on the consumers' physiology and 25 foraging behaviour has been largely overlooked. Here, we document the divergent 26 27 behavioural and physiological responses of three congeneric rodent species in the Middle East, seed dispersers versus seed predators, to fruit consumption. The fruit pulp of the 28 29 desert plant Ochradenus baccatus contains high concentrations of glucosinolates (GLSs). These GLSs are hydrolyzed into active toxic compounds upon contact with the 30 myrosinase enzyme released from seeds crushed during fruit consumption. Acomys 31 russatus and A. cahirinus share a desert habitat. Acomys russatus acts as an O. baccatus 32 33 seed predator, and A. cahirinus circumvents the activation of the GLSs by orally expelling vital seeds. We found that between the three species examined, A. russatus was 34 35 physiologically most tolerant to whole fruit consumption and even A. minous, which is evolutionarily naïve to O. baccatus, exhibits greater tolerance to whole fruit consumption 36 37 than A. cahirinus. However, like A. cahirinus, A. minous may also behaviourally avoid 38 the activation of the GLSs by making a hole in the pulp and consuming only the seeds. 39 Our findings demonstrate that seed predators have a higher physiological tolerance than seed dispersers when consuming fruits containing toxic PSCs. The findings also 40 41 demonstrate the extreme ecological/evolutionary lability of this plant-animal symbiosis to 42 shift from predation to mutualism and vice versa.

44 Introduction

Fleshy, ripe fruit pulp is a nutritious and easily digestible reward that facilitates seed dispersal by animals (Herrera, 1982; Corlett and Lucas 1990; Jordano, 2000; Izhaki, 2002a). However, fruits are likely to attract not only seed dispersers but also seed predators since seeds are also a valuable food source (Corlett and Lucas 1990; Kelt et al. 2004). Eventually, the nature of the plant-animal interaction is determined by the consumers' fruit eating strategy (Corlett and Lucas 1990; Jordano 2000; Dominy and Duncan 2005).

52 Variability in fruit eating can be a function of multiple parameters, such as age, 53 satiation/hunger, feeding location, seasonality, or availability of other food or water (Levey, 1987; Corlett and Lucas 1990; Gautier-Hion et al. 1993; Kaplin and Moermond 54 55 1998; Lambert 1999; Samuni-Blank et al. 2012). In addition, fruits may contain plant secondary compounds (PSCs) to deter seed predators or alter the behaviour of the 56 57 consumer (Cipollini and Levey 1997, Samuni Blank et al. 2012). In folivorous herbivores, evolutionary pressure has resulted in various behavioural and physiological 58 adaptations for coping with PSCs, thus altering plant-herbivore interactions (McArthur et 59 al. 1991; Dearing et al. 2005; Torregrossa et al. 2011). Similar adaptations to fruit PSCs 60 may also influence fruit eating strategies and thus affect plant-animal interactions. 61

Glucosinolates (GLSs) are a common class of fruit PSCs. Intact GLSs are 62 63 generally nontoxic and have limited effect in defending the plant against herbivores and 64 pathogens. The defensive properties of GLSs are enhanced upon their hydrolysis by the myrosinase enzyme causing the release of a toxic aglycone (Hopkins, 2009). These 65 unstable molecules rearrange into several toxic biological compounds (Kjaer, 1976; Das, 66 67 2000; Wittstock and Halkier 2002). Generally, GLSs and myrosinases are 68 compartmentalized to prevent toxicity to the plant, and only mix upon damage to tissue (Rask et al. 2000; Hopkins, 2009). The myrosinase-GLS system, also known as the 69 70 mustard oil bomb, provides plants with an effective defense against generalist herbivores (Wittstock and Halkier 2002). 71

72 The GLS-myrosinase defense system is employed in the fleshy fruit of 73 Ochradenus baccatus (Resedaceae), a common inhabitant of wadis and depressions in 74 Israeli desert regions (Wolf and Shmida 1995, 1997; Wolf and Burns 2001; Bronstein et 75 al. 2007). The fleshy fruits of O. baccatus attract a number of consumers because of its high water and sugar content (Bronstein et al. 2007; Spiegel and Nathan 2007; 2011). 76 77 However, the seeds of O. baccatus also represent a valuable nutrient source, as they are rich in protein (~25% of dry mass). To prevent seed predation, the fruits of O. baccatus 78 defend themselves with a unique arrangement of the mustard oil bomb, where GLSs are 79 80 found in the pulp and the myrosinase enzyme is found in the seeds (Samuni-Blank et al. 2012). Thus, activation of the mustard oil bomb only occurs if consumers crush both 81 seeds and pulp together. 82

83 Ochradenus baccatus naturally co-occurs with two congeneric rodent species (Acomys russatus and A. cahirinus) that differ in their interactions with the plant. Acomys 84 85 russatus usually consumes the O. baccatus fruit as a whole, acting as a seed predator, while A. cahirinus consumes only the pulp and spits the seeds, thus acting as a seed 86 87 disperser (Samuni-Blank et al. 2012, 2013). These interactions result in differential exposure to activated GLSs between species: A. russatus consumes activated GLSs by 88 89 masticating the seeds, while A. cahirinus consumes unactivated GLSs by expelling the 90 seeds. We hypothesize that the variation in physiological adaptations to fruit PSCs drive 91 these plant-animal interactions or vice versa. We also predict that toxin avoiders will 92 show low physiological tolerance to whole fruit consumption and that the frequency of 93 toxin avoiders within a species will be indicative of its tolerance.

94 The aim of the present study was to test the effects of fruits PSCs (pulp and seeds) 95 on fruit consumers by preventing behavioural circumvention, while testing the 96 physiological impacts of fruit PSCs on seed dispersers versus seed predators. For comparative purposes, we also documented the fruit eating behaviour and physiological 97 98 responses to fruit PSCs of the Crete spiny mouse (A. *minous*), which is endemic to the 99 Mediterranean island of Crete and evolutionarily naïve to O. baccatus fruits. We 100 presented each species with diets containing intact or activated GLSs, and monitored 101 various physiological parameters such as body mass, food intake, dry matter digestibility and serum markers of liver damage. We predicted that the seed predator, *A. russatus*,
would be physiologically more adapted to consume activated toxins, as it regularly does
so in the wild. We also predicted that consumption of food containing activated GLSs
would result in liver damage.

106

107 Materials and Methods

108 Fruit collection

Fruits were collected from the Almog junction site (31°48'N, 35°27'E) located near the Dead Sea and kept at -20°C. For experiments in which separation of pulp and seeds was needed, fruits were kept at 4°C and pulp was manually separated from the seeds within two days of collection and then kept separately at -20°C Stored fruits maintained active myrosinase and GLSs content (Samuni-Blank et al. 2012). *Ochradenus baccatus* fruiting occurred year-round, with high peaks in May and November 2008-2012. Whole fruits and fruit pulp were thawed before use.

116 Animals and Maintenance

The experimental protocols were approved by the University of Haifa Committee of 117 Animal Experimentation (Permit number 096/08). A. cahirinus (but see Volobouev et al. 118 2007 for a discussion on the A. cahirinus-dimidiatus complex) and A. russatus were from 119 captive breeding colonies (at least three generations in captivity) originally established 120 from individuals trapped in the vicinity of the Dead Sea (31°28'N, 35°23'E). Animals 121 122 were maintained at the Department of Biology and Environment at the University of Haifa, Oranim. Adult individuals of A. minous were live-trapped in the vicinity of 123 124 Heraklion, Crete, a few days prior to the experiment using Sherman folding traps placed under rocks. All captured A. minous individuals were maintained at the Natural History 125 126 Museum of Crete (Ministry of Environment, Energy and Climate Change, Permit number 127 117272/586).

Prior to experiments, animals were fed rodent chow (Koffolk serial no. 19510) and fresh carrots as a source of free water, *ad libitum*. During the experiments, animals were housed individually in standard mouse cages (21 X 31 X 13 cm) in a temperature-

131 controlled room $(25 \pm 2^{\circ}C)$ under a 12:12 h light-dark cycle. All animals were naïve to O.

132 *baccatus* prior to the experiment.

133 Fruit eating behaviour of A. minous

To determine the fruit eating behaviour of A. minous, we placed each single animal (n =134 16) in a cage with five fruits over-night. The cage floor was examined the next morning 135 for intact fruit parts (pulp or seeds) and the fruit eating behaviour was classified as one of 136 three types: "Whole fruit" - eating pulp and seeds simultaneously and leaving no remains; 137 "Pulp" - eating the pulp and leaving the seeds; "Seed" - eating the seeds and leaving the 138 139 pulp. Individuals that exhibited more than one fruit eating behaviour were classified 140 according to their dominant strategy. For example, if one individual ate two whole fruits and only the seeds of the remaining three fruits, then the behaviour was classified as 141 "Seed". We define avoiders as individuals that consume either pulp or seeds, but not 142 143 both. We define confronters as individuals that consume the whole fruit, pulp and seeds simultaneously. We recorded the number of intact seeds dropped to the cage floor as well 144 as the number intact defecated seeds (in the feces). Similar experiments were previously 145 146 performed on the two other Acomys species (Samuni-Blank et al. 2013).

147 Physiological Effects of Fruit Toxins

148 To examine the physiological effects of different fruit eating strategies on the three rodent species, we performed controlled feeding trials. Diets included rodent chow (Koffolk 149 150 serial no. 19510) mixed with homogenized pulp (Pulp) or pulp and crushed seeds (Mash; Table 1). The fruit pulp was free of myrosinase and contained only intact GLSs. The 151 152 mash diet combined the GLSs from the pulp and the myrosinase enzyme from the seeds and therefore contained activated GLSs. To allow the animals to acclimate to the fruit 153 154 diet, they were fed a 25% fruit diet (pulp or mash; wet weight) for one day combined with 75% chow, followed by 50% fruit mixed with 50% chow for three additional days. 155 156 Wet food pellets (~50% water content) were prepared from the mashed food and given to 157 the rodents within an hour of preparation (n = 6-8 individual rodents per treatment).

Each experiment lasted four days. During the experiment, body mass and food intake were measured daily. Excreta and food leftovers were collected every day from the plastic cage floor, dried (50°C for 24 h), and weighed. Dry matter (DM) digestibility of food consumed by an animal was calculated from its DM food intake and fecal DM output as: DM digestibility = (DM food intake – fecal DM output)/ DM food intake.

163 The effect of activated GLSs on the liver was studied through measurements of liver enzymes and liver function test. On the last day of the experiment, animals were 164 euthanized with CO₂ and immediately dissected. Blood samples were collected by heart 165 puncture and centrifuged (15 min at 2500 rpm). Blood serum was collected and stored at 166 167 -20°C. Samples were analyzed for alkaline phosphatase (ALP) and alanine aminotransferase (ALT). ALT is an intracellular enzyme of hepatocytes and its 168 appearance in blood is indicative of possible liver damage. Acomys cahirinus and A. 169 170 russatus samples were also analyzed for albumin, gamma-glutamyl transpeptidase 171 (GGT), total bilirubin (Bili. Tot.) and direct bilirubin (Bili. Dir.). Serum levels of all 172 these enzymes and proteins are known to increase when hepatobiliary damage occurs 173 (Ozer et al. 2008). All samples were analyzed at the Laboratory of Clinical Biochemistry 174 in Rambam Medical Center, Haifa.

175 Statistical analyses

To test for differences between the fruit eating behaviour of A. minous, we used a one 176 177 proportion Z test. Physiological responses on the last day of the diets (body mass, DM digestibility, liver enzymes, etc.) were compared using two-way ANOVA (with species 178 and diet as main effects) followed by Tukey's Honestly-Significant-Difference (HSD) and 179 180 linear regression. We also used t-test to compare the slopes of two regression lines. The data were tested for normality prior to statistical comparisons. To evaluate the dispersion 181 of individuals' values from the mean values within each of the species, we calculated the 182 183 coefficient of variation (CV = 100*S.D./mean, %) of the three Acomys species under the 184 two diet treatments. Sample size (n = 5-6 per species) was kept consistent for each of the parameters within each of the diets. For all CV calculations, we used the same sample 185

size, in cases of unequal sample sizes, values from larger groups were removed atrandom.

In all cases, significance level was set at P < 0.05. All data were reported as means ± standard error (SE). Statistical analysis was conducted using SPSS 19.0 (SPSS, USA).

190

191 **Results**

192 Fruit eating behaviours

Acomys russatus exhibited the "Whole fruit" behaviour (confronter, consumed pulp and 193 seeds mashed together; movie available at: http://y2u.be/RcLDPst87vs) whereas A. 194 195 *cahirinus* exhibited the "Pulp" behaviour (avoider, consumed the pulp and spat the seeds; movie available at: http://y2u.be/25XI_mtglPU) (Samuni-Blank et al. 2013). Five 196 individuals of A. minous left the fruit untouched, and were not included in the analysis. 197 Interestingly, the most common fruit eating behaviour of A. minous differed from that of 198 the other two Acomys species (Table 2). Eight out of eleven individuals employed, at 199 200 least once, the "Seed" behaviour, by making a hole in the pulp, eating the seeds, and leaving the pulp on the cage floor (movie available at: http://y2u.be/yvHL7oA0HbM). 201 202 The "Seed" behaviour was the dominant behaviour for six of these eight. From these six 203 individuals, four individuals exclusively exhibited the "Seed" (avoider) behaviour and the 204 remaining two individuals alternated between the "Seed" and "Whole fruit" behaviours.

A smaller proportion of A. minous (36.6%) preferred the "Whole fruit" (confronter) 205 206 behaviour (movie available at: http://y2u.be/ghVA7Ibhu8c). Half used the "Seed" strategy exclusively and the others alternated between "Seed" and "Whole fruit" 207 behaviour. Only a single individual used only the "Pulp" (avoider) behaviour (movie 208 available at: http://y2u.be/rb1vC3EbnP8). There was no significant difference between 209 these feeding behaviours (avoider vs. confronter; Z = 0.89, n = 11, N.S). For all 210 individuals, no intact seeds were found in the feces. Overall, we characterized A. minous 211 as avoider since more than 50% of the individuals avoided the consumption of the whole 212 213 fruit and its activated PSCs.

214 Physiological Effects of Fruits

Body mass: Final body mass (% of initial) differed significantly among species ($F_{2, 39} = 29.5, P < 0.001$) and between diets ($F_{1, 39} = 32.5, P < 0.001$). In addition, there was a marginal significant diet*species interaction ($F_{2, 39} = 3.1, P = 0.05$). Post-hoc Tukey's HSD tests showed that on pulp diet *A. cahirinus* and *A. russatus* maintained the lowest and highest body mass, respectively (P < 0.05). Similarly, when fed mash diet, *A. cahirinus* lost significantly more (~20%) of its initial body mass (P < 0.05), compared to *A. minous* and *A. russatus*, which lost ~15% and ~10%, respectively (Table 3).

Food intake: DM food intake (% body mass/day) differed significantly among species ($F_{2, 39} = 26.4$, P < 0.001) and between diets ($F_{1, 39} = 23.9$, P < 0.001). There was a significant interaction between species and diet ($F_{2, 39} = 4.1$, P < 0.05). Post-hoc Tukey's HSD tests showed that *A. minous* maintained DM intake on the mash diet, while the other two species exhibited a significant decrease (P < 0.05) in DM intake on the mash diet compared to the pulp diet. In addition, *A. cahirinus* DM intake on the mash diet was significantly (P < 0.05) lower than that of the other two rodents (Table 3).

Digestibility: There were no significant differences in DM digestibility (Table 3) among the species ($F_{2, 37} = 0.09$, N.S), between diets ($F_{1, 37} = 1.03$, N.S), and no significant interaction ($F_{2, 37} = 0.49$, N.S.).

There was a positive correlation between body mass at the end of the trial (% of initial) and total DM food intake for *A. cahirinus* and for *A. russatus*, and a marginally positive correlation for *A. minous*, while on the pulp diet (Fig. 1A). There were no significant correlations between intake and final body mass for any species while on the mash diet. The slopes of regression lines differed significantly for pulp and mash diets in all three *Acomys* species combined ($t_{43} = 3.02$, P < 0.01; Fig. 1B).

Serum biochemical assay: To our knowledge, we were the first to document serum markers of liver damage of any *Acomys* species. Serum alanine aminotransferase (ALT) activity differed significantly by diet ($F_{1, 38} = 7.9, P < 0.01$), but not by species ($F_{2, 38} = 0.1$, N.S), and there was a significant diet*species interaction ($F_{2, 38} = 3.6, P < 0.05$) (Fig. 2A). Inspecting the figure, it was apparent that serum ALT was elevated about two times in *A. cahirinus* and *A. minous* when eating mash as compared with pulp, whereas *A. russatus* showed no change on the two diets. Despite the significant effects overall in the
ANOVA, none of the post-hoc pair-wise comparisons (Tukey's HSD tests) were
significant.

There was a significant difference in serum alkaline phosphatase (ALP) activity (Fig. 2B) among species ($F_{2, 36} = 8.9$, P = 0.001) but not between diets ($F_{1, 36} = 1.8$, N.S) nor was there a diet*species interaction ($F_{2, 36} = 1.6$, N.S). Post-hoc Tukey's HSD tests showed that *A. russatus* had significantly lower ALP levels than the other two species (P< 0.05). All other comparisons were not significant.

For *A. cahirinus* and *A. russatus*, gamma-glutamyl transpeptidase (GGT) activity was significantly different among species ($F_{1, 28} = 11.7$, P = 0.002) but not between diet treatments ($F_{1, 28} = 0.8$, N.S), and the diet*species interaction was not significant ($F_{1, 28} =$ 0.4, N.S). For albumin, total bilirubin (Bili. Tol.) and direct bilirubin (Bili. Dir.), there were no significant differences (P > 0.05) between diet treatments, among species or nor was there a diet*species interactions (Table 4).

258 **Coefficient of variation:** The coefficient of variation (CV) of all parameters 259 measured for each the three species and the two diets revealed that for eight out of ten 260 different CV values *A. russatus* had the lowest scores (Table 5).

261

262 **Discussion**

The mustard oil bomb products are known to have numerous physiological effects on animals, such as reduction in growth and in food intake, as well as damage to the liver (Duncan and Milne 1992; Duncan and Milne 1993; Kim et al. 1997; Sørensen et al. 2001). In the present study, we demonstrated that the activated GLSs of *O. baccatus* differentially affected the performance of seed dispersers and predators.

Two of the three studied species, *A. cahirinus* and *A. russatus*, were previously reported to exhibit very different feeding behaviours: *A. cahirinus* separate the pulp from the seeds to avoid the GLSs activation while *A. russatus* consume the whole fruit, pulp and seeds. Here, we report that a third species, *A. minous*, evolutionary naïve to *O. baccatus* and closely related to *A. cahirinus*, showed a different dominant behaviour:
puncturing a hole in the pulp and consuming only the seeds or consuming the whole fruit.
This species tended to avoid the consumption of hydrolyzed GLSs (like *A. cahirinus*), but
at the same time acted as a seed predator (like *A. russatus*).

276 In line with its behaviour, A. minous also incurred greater physiological effects than A. russatus. The physiological results were consistent with each of the species' fruit 277 278 eating strategies: species with a high percentage of PSC avoiders, i.e. individuals spitting 279 seeds or eating only the seeds (Table 2), exhibited a more negative effect of the mash diet 280 on body mass maintenance. After four days on the mash diet, A. russatus (the confronter) maintained ~ 90% of its initial body mass, while avoiders A. minous and the seed 281 disperser A. cahirinus maintained only ~ 85% and ~ 80% of their initial body mass, 282 283 respectively. Change in body mass was correlated with food consumption (Fig. 1B) and 284 all species consumed more DM on pulp diet than on mash diet (Table 3).

285 Although A. minous maintained DM intake when feeding on the mash diet, individuals eating that diet lost significantly more body mass than those eating only pulp. 286 This mass loss may stem from the higher energetic costs of detoxifying the hydrolyzed 287 products of the GLSs. Detoxification costs are relatively high, and can be as high as 45% 288 289 of the basal metabolic costs in specialist herbivores (Sorensen et al. 2005). Accordingly, 290 positive correlations between body mass and food intake for each species were found only on the pulp diet (Fig. 2A). Detoxification costs may also explain the different slopes 291 292 of the regression lines for the pulp and the mash diets across all *Acomys* species (Fig. 293 2B).

The very low DM intake of *A. cahirinus*, resulting in significant loss of body mass when feeding on the mash diet, could be explained by the bitter taste of the mustard oil bomb, which serves as a deterrent (Fenwick et al. 1983a,b; Garcia-Bailo et al. 2009). Indeed, taste is a valuable tool to estimate food quality and bitter taste perception prevents animals from ingestion of toxic compounds (Chandrashekar et al. 2000; Nelson et al. 2001; Mueller et al. 2005; Chandrashekar et al. 2006). Previous studies have

demonstrated that rodents are deterred by bitter taste and also that bitter taste inhibits 300 food intake (Glendinning et al. 1990; Scalera, 1992). However, different mice species 301 302 may have different taste sensitivities (Glendinning et al. 1990). This could partly explain 303 the higher DM intake values and thus the maintenance of higher body mass of A. russatus 304 and A. minous when feeding on a mash diet. Also, since individuals of A. minous in this 305 study were captured in the wild and thus were exposed to a wider range of diets and environmental changes, it is reasonable that they will show a greater tolerance compared 306 307 to captive-bred A. cahirinus.

Fleshy ripe pulp is used by plants as a nutritious and easily digestible reward for seed dispersing animals (Herrera, 1982; Corlett and Lucas 1990; Jordano, 2000; Izhaki, 2002b). We propose that digestibility did not differ between the pulp and mash diets because of the low percentage of seeds (~8%; wet weight) compared to pulp (~92%) in *O. baccatus* fruits.

As an index to the toxicity of the diets, we tested for serum levels of a set of 313 enzymes and proteins known to increase when hepatobiliary damage occurs (ALT, ALP, 314 GGT, albumin, and bilirubin; Ozer et al. 2008). Toxins can cause acute liver cell necrosis, 315 316 upon damage liver cells, aminotransferases enzymes are released into the bloodstream, 317 and their activity is elevated in the serum (Ozer et al. 2008; Hyder et al. 2013). Indeed, 318 for A. cahirinus and A. minous, ALT levels were almost two fold higher on the mash diet 319 than on the pulp diet. For A. cahirinus there was also an increase in ALP levels on the mash diet. However, on mash diet, A. cahirinus refused to eat (total consumption over 320 four days of the experiment was less than 1.5 g) and lost almost 20% of its initial body 321 322 mass. Severe weight loss is known to be associated with the nonalcoholic fatty liver 323 disease (NAFLD; Salt 2004). Mice with NAFLD have been shown to possess significantly higher levels of ALT and ALP but not albumin (Wang et al. 2011). Thus, 324 the severe weight loss on mash diet can also be the reason for the elevated levels in A. 325 326 *cahirinus* and *A. minous* (who lost more than 15% of its initial body mass) in which the 327 majority of individuals would typically avoid the consumption of the whole fruit and its 328 activated PSCs.

329 The seed predator A. russatus displayed the least variability (i.e., lowest CVs) out of all the study species, while on different diets and within the various parameters. In 330 331 addition, A. russatus appeared more tolerant to O. baccatus diets in terms of defending 332 body mass when feeding on the mash diet, which contained the toxic components of the mustard oil bomb. This result is consistent with other studies showing the ability of A. 333 russatus to maintain body mass under various conditions (Shkolnik and Borut 1969; Kam 334 and Degen 1993; Gutman et al. 2006). Physiologically, the unique adaptive mechanisms 335 of the diurnal A. russatus for desert survival (Haim and Borut 1981; Haim et al. 1994; 336 Haim et al. 2005; Ehrhardt et al. 2005; Levy et al. 2011) may also explain their low 337 variability under the different treatments across various examined parameters. 338

Despite more than three generations of O. baccatus naivety, the seed predator, A. 339 russatus, was the least affected by fruits PSCs, suggesting its higher physiological 340 tolerance to the activated GLSs, while A. cahirinus, the seed disperser, was most 341 342 negatively affected. The ability of the wild-caught, naïve A. minous, and the captive-bred A. cahirinus to behaviourally circumvent the activation of the GLSs, and the 343 344 physiological ability of A. russatus to tolerate the mustard oil bomb, demonstrate the wide ecological/evolutionary lability of this plant-animal symbiosis to shift from 345 346 predation to mutualism and vice versa.

348 List of abbreviations

349	ALP	Serum alkaline phosphatase
350	ALT	Serum alanine aminotransferase
351	Bil. Dir.	Direct bilirubin
352	Bil. Tot.	Total bilirubin
353	DM	Dry matter
354	GGT	Serum gamma-glutamyl transpeptidase
355	GLSs	Glucosinolates
356	NAFLD	Nonalcoholic fatty liver disease
357	PSCs	Plant secondary compounds

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364

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525

Figure 1. Change in body mass as a function of total dry matter (DM) intake over all four days of the experiment. (A) *A. cahirinus* (n = 16), *A. minous* (n = 13) and *A. russatus* (n = 16) feeding on pulp or mash of *O. baccatus*. (B) Change in body mass as a function of dry matter (DM) intake on the pulp *vs*. the mash diet, combining all three species.

531

Figure 2. Serum biochemical assay of *A. cahirinus* (n = 8 for each diet), *A. minous* (n = 533 5-6 for each diet) and *A. russatus* (n = 8 for each diet) on pulp (white bars) and mash (gray bars) *O. baccatus* fruits for (A) Alanine aminotransferase (ALT) and (B) Alkaline phosphatase (ALP). Data are presented as means \pm SE.

536

537 **Table 1.** Diet components (%).

538

Table 2. Summary of the fruit-eating behaviour of the three congeneric species of *Acomys.* *5 other individuals did not consume any part of the fruit and were excluded.

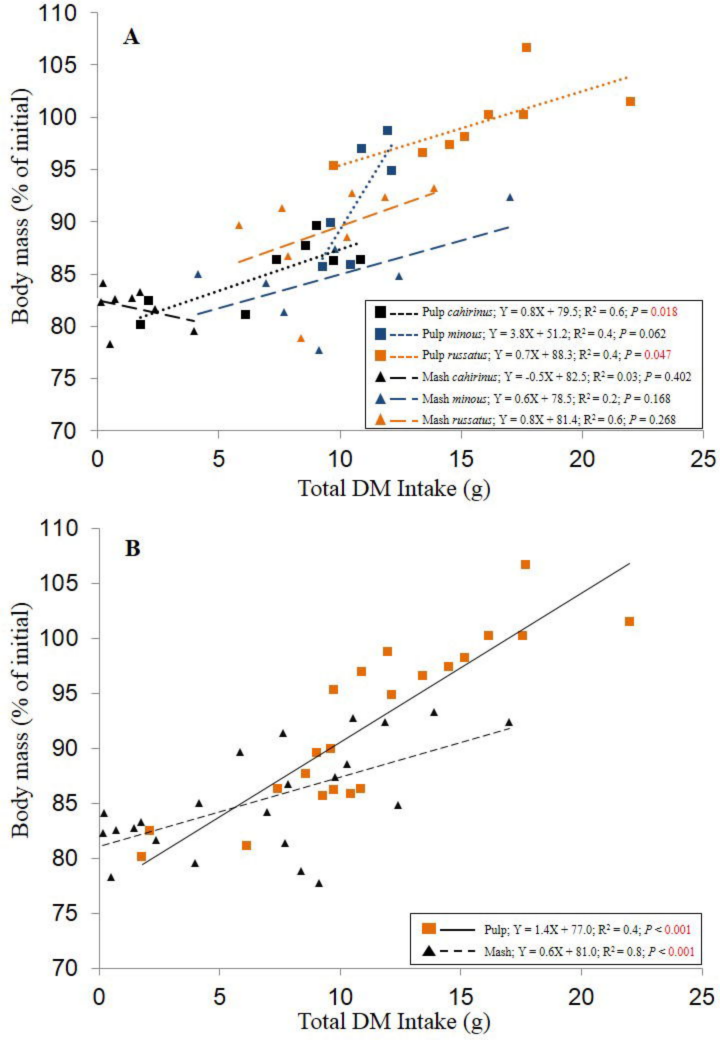
542**Table 3.** Summary of the feeding trials. Body mass (% of initial), dry matter intake (%543body mass/day) and dry matter digestibility (%) of *A. cahirinus*, *A. minous*, and *A.*544*russatus* on day 4 of the trial. Similar letters adjacent to means indicate no significant545difference (P > 0.05) among the means (two-way ANOVA followed by Tukey's HSD).546Data are means \pm SE.

547

Table 4. Markers of liver damage of *A. cahirinus* (n = 8 for each diet) *A. russatus* (n = 8 for each diet) on different diets. The parameters measured are gamma-glutamyl

transpeptidase (GGT), albumin, total bilirubin (Bili. Tol.) and direct bilirubin (Bili. Dir.)
Data are means ± SE.

Table 5. Coefficient of variation (%) of the three *Acomys* species in the two diet treatments of the different parameters: body mass, dry matter intake, dry matter digestibility, alkaline phosphatase (ALP) and alanine aminotransferase (ALT). The lowest score for each test within each diet is marked in bold.



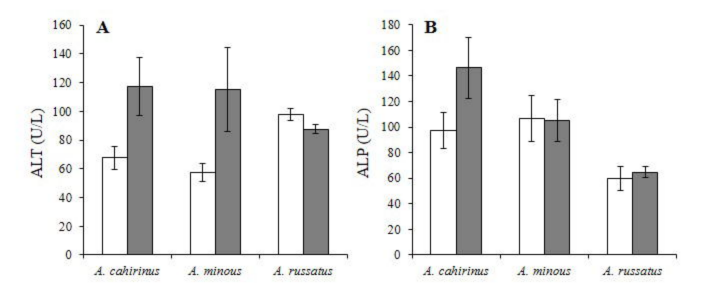


Table 1.

		Mash	Pulp
Day 1	Chow	50	50
	Water	25	25
	Pulp	23	25
	Seeds	2	0
Day 2-4	Chow	50	50
	Water	0	0
	Pulp	46	50
	Seeds	4	0

Table 2	2.
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	Ν	Pulp	Seed	Whole	Reference
		(%)	(%)	(%)	
A. cahirinus	43	76.7	4.6	18.6	Samuni-Blank et al. 2013
A. minous	11*	9.1	54.5	36.4	Current study
A. russatus	43	11.6	0	88.4	Samuni-Blank et al. 2013

Table	e 3 .
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Parameters	Diet	A. cahirinus	Ν	A. minous	Ν	A. russatus	N
Body Mass	Pulp	$85.0 \pm 1.2^{\text{CD}}$	8	92.0 ± 2.3 ^B	6	99.5 ± 1.3 ^A	8
	Mash	$81.8\pm0.7^{\rm D}$	8	$84.8 \pm 1.7^{\text{CD}}$	7	89.2 ± 1.3^{BC}	8
Food intake	Pulp	$3.3\pm0.9^{\rm \ B}$	8	$5.9\pm1.0^{\rm \ AB}$	6	$8.6\pm0.5^{\rm A}$	8
	Mash	$0.6\pm0.2^{\rm C}$	8	$5.3\pm0.7^{\rm \ B}$	7	$4.1\pm0.6^{\rm \ B}$	8
Digestibility ^{N.S.}	Pulp	84.9 ± 2.0	8	82.9 ± 2.1	6	81.3 ± 1.1	8
	Mash	80.3 ± 5.6	6	80.6 ± 2.6	7	81.8 ± 2.0	8

Table 4.	
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Parameters	Diet	A. cahirinus	A. russatus
GGT (U/L)	Pulp	12.3 ± 1.4	8.9 ± 1.7
	Mash	12.0 ± 0.8	7.0 ± 0.5
Albumin (g/dL)	Pulp	1.1 ± 0.0	1.2 ± 0.1
	Mash	1.1 ± 0.0	1.1 ± 0.0
Bili. Tot. (mg/dL)	Pulp	0.1 ± 0.0	0.1 ± 0.0
	Mash	0.1 ± 0.0	< 0.05
Bili. Dir. (mg/dL)	Pulp	< 0.01	< 0.01
	Mash	< 0.01	< 0.01

Table	5.
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Parameters	Diet	Coefficient of variation (%)			
rarameters	Diet	A. cahirinus	A. minous	A. russatus	
Body mass	Pulp	3.07	6.15	1.94	
	Mash	1.92	5.92	5.88	
Food intake	Pulp	68.93	38.74	17.41	
	Mash	41.36	37.18	45.23	
Digestibility	Pulp	7.77	6.29	2.21	
	Mash	17.04	9.30	6.38	
ALP	Pulp	29.20	37.38	11.3	
	Mash	44.82	34.46	20.57	
ALT	Pulp	40.26	25.95	13.18	
	Mash	49.50	62.07	9.87	