See discussions, stats, and author profiles for this publication at: https://www.researchgate.net/publication/262523769

Experimental evidence for bird pollination and corolla damage by ants in the short-tubed flowers of Erica halicacaba (Ericaceae)

Article in South African Journal of Botany · January 2011

DOI: 10.1016/j.sajb.2011.11.003

CITATION:	S	reads 48	
4 autho	rs , including:		
	Jeremy Midgley University of Cape Town 238 PUBLICATIONS 8,085 CITATIONS	.	Phoebe BarnardPacific Biodiversity Institute and University of147 PUBLICATIONS 1,265 CITATIONS
	SEE PROFILE		SEE PROFILE
	Robert Simmons		
	University of Cape Town		
	117 PUBLICATIONS 1,378 CITATIONS		
	SEE PROFILE		

Some of the authors of this publication are also working on these related projects:



Damara Tern conservation View project

Project

Black Harrier conservation in South Africa View project

All content following this page was uploaded by Robert Simmons on 23 May 2014.

AUTHOR QUERY FORM

	Journal: SAJB	Please e-mail or fax your responses and any corrections to: E-mail: <u>corrections.esil@elsevier.spitech.com</u> Fax: +1 619 699 6721
ELSEVIER	Article Number: 743	

Dear Author,

Please check your proof carefully and mark all corrections at the appropriate place in the proof (e.g., by using on-screen annotation in the PDF file) or compile them in a separate list. Note: if you opt to annotate the file with software other than Adobe Reader then please also highlight the appropriate place in the PDF file. To ensure fast publication of your paper please return your corrections within 48 hours.

For correction or revision of any artwork, please consult http://www.elsevier.com/artworkinstructions.

Any queries or remarks that have arisen during the processing of your manuscript are listed below and highlighted by flags in the proof. Click on the 'Q' link to go to the location in the proof.

Location in article	Query / Remark: <u>click on the Q link to go</u> Please insert your reply or correction at the corresponding line in the proof
<u>Q1</u>	Please confirm that given names and surnames have been identified correctly.
<u>Q2</u>	Please check the telephone number of the corresponding author, and correct if necessary.
<u>Q3</u>	Missing Fig. 1 citation was inserted here. Please check if appropriate.

Thank you for your assistance.

SAJB-00743; No of Pages 1

ARTICLE IN PRESS

ELSEVIER

Available online at www.sciencedirect.com

SciVerse ScienceDirect

SOUTH AFRICAN JOURNAL OF BOTANY

South African Journal of Botany xx (2011) xxx

Highlights

www.elsevier.com/locate/sajb

Experimental evidence for bird pollination and corolla damage by ants in the short-tubed South African Journal of Botany xxx (2011) xxx-xxx flowers of Erica halicacaba (Ericaceae) R.C. Turner^{a,*}, J.J. Midgley^b, P. Barnard^c, R. Simmons^d, S.D. Johnson^a ^a School of Life Sciences, University of KwaZulu-Natal Pietermaritzburg, Private Bag X01, Scottsville 3209, South Africa South Africa ^b Department of Botany, University of Cape Town, Rondebosch 7701, South Africa South Africa

^d Percy Fitzpatrick Institute, Department of Zoology, University of Cape Town, Rondebosch 7701, South Africa

► We investigate bird pollination and floral robbery in *Erica halicacaba*. ► We use exclosure experiments, bird mist-netting and observation to determine the avian pollinator. ► We use predator exclusion experiments to determine that ants rob flowers.

0254-6299/\$ -see front matter © 2011 SAAB. Published by Elsevier B.V. All rights reserved. doi:10.1016/j.sajb.2011.11.003

SAJB-00743; No of Pages 7



ARTICLE IN PRESS

Available online at www.sciencedirect.com

SciVerse ScienceDirect

SOUTH AFRICAN JOURNAL OF BOTANY

South African Journal of Botany xx (2011) xxx-xxx

www.elsevier.com/locate/sajb

Experimental evidence for bird pollination and corolla damage by ants in the short-tubed flowers of *Erica halicacaba* (Ericaceae)

R.C. Turner^{a,*}, J.J. Midgley^b, P. Barnard^c, R. Simmons^d, S.D. Johnson^a

^a School of Life Sciences, University of KwaZulu-Natal Pietermaritzburg, Private Bag X01, Scottsville 3209, South Africa

^b Department of Botany, University of Cape Town, Rondebosch 7701, South Africa

^c South African National Biodiversity Institute, Kirstenbosch, Cape Town, South Africa ^d Percy Fitzpatrick Institute, Department of Zoology, University of Cape Town, Rondebosch 7701, South Africa

Received 31 July 2011; received in revised form 22 November 2011; accepted 22 November 2011

11 Abstract

9 10

12 Unrelated plants pollinated by similar animals tend to show convergent evolution of floral traits. Floral syndromes have been used successfully 13to develop hypotheses about pollination systems but can be misleading when plants have unusual floral morphology or mechanisms of pollen transfer. A case in point is Erica halicacaba, a local endemic shrub on the Cape Peninsula of South Africa. Its short-tubed greenish-yellowish 14 15 flowers with a narrow aperture have been considered to be insect pollinated. However, field observations, selective exclusion experiments and 16 analysis of pollen loads all indicate that its primary pollinator is the Orange-Breasted Sunbird (Anthobaphes violacea). These sunbirds were com-17 mon visitors and netted individuals carried large numbers of *E. halicacaba* pollen tetrads on their culmens, rather than on their head plumage as is typical for Cape Erica species with longer tubular flowers. Plants of E. halicacaba from which vertebrates were excluded had a lower incidence of 18 anther tripping and set significantly fewer seeds than those exposed to both birds and insect visitors. Nectar in E. halicacaba is present in small 1920concentrated amounts and, typical of sunbird-pollinated plants, is dominated by sucrose. Flowers often had holes in their corollas and we identified ants, specifically Myrmicaria nigra, as the likely agents. This was supported by reduced rates of corolla damage on branches from which crawling 21insects such as ants were experimentally excluded. These findings indicate that considerable caution should be applied when attempting to predict 22pollinators from floral syndromes, and highlight the role that crawling insects can play in damaging flowers. 23

²⁴ © 2011 SAAB. Published by Elsevier B.V. All rights reserved.

25 Keywords: Anthobaphes violacea; Bird pollination; Erica halicacaba; Ericaceae; Floral Syndromes; Nectar Robbing; Nectar; Pollination Biology

26

27 **1. Introduction**

The concept of floral syndromes can be traced to Kölreuter's 28761) descriptions of plant-pollinator interactions and floral 29morphology (Fenster et al., 2004) (Fig. 1). It has been controver-O330 sial on account of its association with a typological school of pol-31lination biology that was not grounded in evolutionary biology, 32and because there has been uncertainty about the degree of spec-33 34 ificity in pollination systems (Johnson and Steiner, 2000; Pauw, 2006; Waser et al., 1996). Fenster et al. (2004) defined a floral 35

* Corresponding author. Tel.: +27 217832427.

E-mail address: rossct@telkomsa.net (R.C. Turner).

syndrome as "a suite of floral traits, including rewards, associated ³⁶ with the attraction and utilization of a specific group of animals as ³⁷ pollinators", and further emphasised the importance of classify- ³⁸ ing pollinators into functional groups according to the selective ³⁹ pressures they exerted, as these typically influence evolution of ⁴⁰ suites of floral traits. It is in this sense that Hargreaves et al. ⁴¹ (2004) proposed that "floral syndromes can be seen as patterns ⁴² of convergent evolution that are useful for developing testable ⁴³ hypotheses about pollination systems". ⁴⁴

Workers have emphasised the usefulness of floral syn-45 dromes (De Merxem et al., 2009; Hargreaves et al., 2004; 46 Pauw, 2006) and it is a feature of many pollination studies 47 that pollinators are initially inferred from the floral morphology 48

0254-6299/\$ -see front matter 0 2011 SAAB. Published by Elsevier B.V. All rights reserved. doi:10.1016/j.sajb.2011.11.003

2

ARTICLE IN PRESS



Fig. 1. (A) *Anthobaphes violacea* pollinating *E. halicacaba*. Scale 20 mm. (B) *Erica halicacaba* in typical habitat. Scale 20 mm. (C) Pollen on culmen of juvenile Orar easted Sunbird. Scale 20 mm. (D) Scalloped area on underside of culmen with pollen amalgam. Scale 20 mm. (E) *Myrmicaria nigra* causing floral damage to arr *pralicacaba* flower. Scale 10 mm. (F) Fertilised flower. Scale 20 mm. (G) Male visitor with pollen on culmen. Scale 20 mm. All photos R.C. Turner, except (G) Rob Simmons.

of plants (e.g. Kleizen et al., 2008; Manning and Goldblatt, 49 2005; Turner et al., 2011). In the majority of these studies, pre-50dictions have been confirmed by later observation, thus the pre-51dictive value of floral syndromes can be high. However, the 52utility of floral syndromes has been questioned (Ollerton et al., 532009) and several studies have demonstrated that there are limits 5455to the power of predictive paradigms, especially in specialised single-pollinator mutualisms (Castellanos et al., 2003; De 56Merxem et al., 2009; Johnson, 1995; Olesen and Valido, 2003; 57Pauw, 1998). A case in point is Pauw's (1998) study of bird-58pollinated in Microloma sagittatum (L.) R.Br. (Apocynaceae), a 59species which possesses some traits consistent with bird-60 61pollination, such as the firm-textured, unscented, tubular, red

flowers with accessible perching posts, but, also traits usually as- 62 sociated with insect pollination, such as a short floral tube with a 63 narrow entrance. Ollerton (1998) noted that these traits had "ex- 64 cluded *M. sagittatum* from being recognized as a bird- 65 pollinated plant — these characteristics did not fit preconceived 66 ideas of what such a plant should look like". Similarly, de 67 Merxem et al. (2009) concluded their study of variable flower 68 tube length in *Tritoniopsis revoluta* (Burm.f.) Goldblatt by com- 69 menting that "although syndromes may provide clues about pro- 70 spective pollinators, they are not always a fail-safe way of 71 predicting all of a flowers' important visitors". 72

Bird-pollinated flowers typically have brightly coloured 73 flowers, often red, orange and sometimes pink (Anderson, 74

ARTICLE IN PRESS

2005; Johnson, 1995; Manning and Goldblatt, 2005; Pauw, 751998). The question of why so many bird-pollinated flowers 76 are red has been explored by Raven (1972) and more recently 77 reviewed by Rodríguez-Gironés and Santamaría (2004), who 78 focussed upon the energetics of, and competition between, pol-79 linator guilds to explain why red flowers were favoured by 80 birds and purple flowers by bees. They emphasised that certain 81 bee species can see the red end of the light spectrum and that 82 birds can detect purple flowers, and that an explanation could 83 therefore not be as simple as "bees can't see the colour red". 84 The Cape flora contains numerous examples of tubular Erica spe-85 cies with brightly-coloured, red, pink, white, orange, yellow and 86 green flowers, with approximately 50% of putatively bird-87 pollinated species having more than two colour forms, and in sev-88 eral cases, bicoloured and tricoloured forms (Oliver and Oliver, 89 2002; Rebelo et al., 1985). A noteworthy example is Erica viscaria 90 L. subsp. viscaria, which has flowers "12-20 mm-long, tubular, 91 non-viscid, hairy or postulate, red, pink, purple, white, yellowish 92or green, in some cases bicoloured - pink with a white mouth or 93 red with a yellow mouth" (Oliver and Oliver, 2002). While 94Rebelo et al. (1985) concluded that the Orange Breasted Sunbird 95 was the sole pollinator of ornithophilous Cape Erica species, re-96 cent studies by Geerts and Pauw (2009, 2010) have shown that 97 Southern Double-collared Sunbirds (Cinnyris chalybea) visit 98 flowers of Erica cruenta Sol., Erica discolor Andrews and Erica 99 perspicua Wendl., suggesting that it is not only one specialist pas-100 serine nectarivore that has imposed selective pressure upon bird-101 pollinated Cape Erica species. 102

Baker and Baker (1983, 1990) proposed that hummingbirds 103 104 and passerines such as sunbirds select for different nectar properties in the flowers they pollinated. Johnson and Nicolson 105(2008) suggested that a more useful paradigm involved group-106 ing birds into subcategories of specialist and generalist pollina-107tors. They reported that flowers adapted for pollination by 108 specialized passerine nectarivores, whether hummingbirds or 109 sunbirds, typically had small amounts of sucrose-rich nectar, 110 and that flowers adapted for generalist bird pollinators typically 111 had large amounts of dilute nectar with very low sucrose content. 112 Barnes et al. (1995) showed that 29 of the 37 ornithophilous 113 Erica species they studied had sucrose-dominant nectars with a 114 mean sucrose proportion of $93.8\pm6.2\%$ (mean \pm SD). As their re-115sults were not consistent with the original ideas of the Bakers, 116 they concluded that pollination syndromes could not be deduced 117 from nectar types. However, their finding of sucrose-dominated 118 119 nectar in *Erica* is actually consistent with the more recent finding that plants pollinated by specialized passerine nectarivores tend 120to have sucrose-dominated nectar (Johnson and Nicolson, 2008). 121

The floral morphology of Erica halicacaba L₁ is unique 122within the genus Erica because of its gooseberry-shaped 123flowers (halicacaba=gooseberry. Gr.) with adpressed lobes 124125and its pollinator has been unknown since the taxon was described by Linnaeus in 1760. Rebelo et al. (1985) categorised 126426 south-western Cape *Erica* species "according to their prin-127 cipal putative pollinating agents, based on the shape of the 128 flowers and field observations", suggesting the possibility of 129 bird pollination in E. halicacaba (p. 276), although they subse-130131quently classified the species as insect pollinated in the same

paper (p. 279). Their suggestion was that large flying hymenopterans, e.g. carpenter bees, could reliably pollinate flowers by forcing open the adpressed corolla lobes. Based on observa-134 tions of cultivated *E. halicacaba* plants at the Kirstenbosch Na-135 tional Botanical Garden, Rebelo et al. (1985) observed that bees chewed holes in the bases of corollas in order to rob nectar, and Oliver and Oliver (2000) suggested that the large flowers may serve as an "overnight or cool weather shelter" for insects, presumably resulting in fertilization of flowers. However, preliminary field visiting flowers of *E halicacaba* led us to hypothesize that this spetices, which is a member of a guild of at least ten *Erica* species with similar short-tubular, yellow to greenish-yellow flowers, is specialtive for pollination by sunbirds.

2. Materials and methods

Field studies took place during October and November of 148 2009 on Glen Cairn Ridge, southern Cape Peninsula, South 149 Africa (34.147136S, 18.427021E; 190 m). Approximately 150 250 mature plants were available for study purposes. As pol-151 len tetrads of *E. halicacaba* could potentially be confused 152 with those of other *Erica* species, we checked the study site 153 for other co-flowering *Erica* species and Cape taxa known 154 to have pollen in tetrads.

Erica halicacaba is endemic to the Cape Peninsula (South 157 Africa, Western Cape), occurring on rocky ridges and mountain 158 tops from Table Mountain to Paulsberg in the Cape of Good 159 Hope Nature Reserve. In rockier fire refugia individual plants 160 may attain heights of more than 2 m tall, with gnarled, woody 161 trunks more than 0.15 m in diameter, or form mats up to 2 m^2 162 in area on vertical cliffs. Plants are generally floriferous, mature 163 individuals bearing many hundreds to thousands of gooseberry- 164 shaped, greenish-yellow flowers with a mean corolla length of 165 22.5 mm (20.0–24.5 mm; n=25). Peak flowering occurs from 166 July to November and is variable between localities, depending 167 on aspect, habitat and microclimate. Far more than 2000 plants 168 occur in at least 20 locations and the species is classified as a 169 taxon of least concern (LC) in the 2010 IUCN South African 170 Red Data Plant List (Raimondo et al., 2009). 171

2.3. Pollinator observations and bird mist netting 172

We observed pollinator-plant interactions during the early 173 morning (6–10 am) and late afternoon (4–8 pm) for 12 days, 174 from 19 to 30 November 2009.

Birds were netted on 26 November 2009, using Ecotone 176 12 m, 5-shelf nylon mist-nets with 16 mm mesh. Nets were positioned approximately 30 m from our observation point, a 178 rocky outcrop containing multiple mature *E. halicacaba* individuals. Mist nets were constantly monitored for bird-captures 180 by three fieldworkers for a period of approximately 4 h. 181

Please cite this article as: Turner, R.C., et al., Experimental evidence for bird pollination and corolla damage by ants in the short-tubed flowers of *Erica halica-caba* (Ericaceae), S. Afr. J. Bot. (2011), doi:10.1016/j.sajb.2011.11.003

146

4

ARTICLE IN PRESS

Culmens were swabbed for pollen using sections of double-182 sided adhesive tape ± 5 cm long. The non-adhesive protective 183 backing covering the adhesive surface used for sampling was 184 replaced after culmen swabbing. Netted birds were immediately 185 swabbed while still in mist nets and then disentangled for col-186 lection of biometric data and ringing. Individual birds were re-187 leased less than 10 min after initial capture. Pollen swabs were 188 examined in the laboratory at 45-60×magnification under a 189 Vickers Light Stereoscopic microscope. Pollen grains were 190identified and counted. 191

192 2.4. Selective exclusion experiments

To determine whether birds were important for seed produc-193 tion in E. halicacaba, and also whether plants were capable of 194 autonomous seed production, we selectively excluded either 195vertebrates only or all flower visitors. This was done on 10 sep-196 arate mature plants at different localities within the population. 197Selective exclusion of vertebrates was achieved by covering 198 flowering branches with wire "chicken-mesh" exclosures 199(mesh aperture $\pm 15 \times 17$ mm), and exclusion of all visitors 200 was achieved with bridal veil exclosures (aperture ± 0.8 mm 201 diam.). Uncovered inflorescences were used as controls. Trea-202 ted inflorescences were examined one month later to assess 203rates of visitation and seed set. Visitation to flowers of many 204Erica species, including E. halicacaba, is easily scored by de-205termining if the anther rings have been tripped (Geerts and 206Pauw, 2010; Turner et al., 2011). Mature, unvisited flowers 207have an untripped anther ring, i.e. adjacent anther thecae remain 208 209 connected laterally. When manipulated, anther rings break (are triggered) and pollen is visibly ejected from apical anther pores. 210Visits by Orange Breasted Sunbirds, as observed in the field 211 during this study, result in tripped anther rings. Seed set was 212 established by dissection of fruits. Developing seeds are distin-213guishable from ovules by their larger size and "honey-combed" 214testa. 215

216The effects of selective exclosure on the proportion of flowers on each treated branch that had tripped anthers or 217 which developed fruits was analyzed using generalized linear 218 models with a binomial error distribution, implemented in 219 220 PASW Statistics 18 (SPSS, Chicago, USA). Plant was used as a blocking factor and was entered in the model before treat-221 ment. Model significance was assessed using likelihood ratio 222 tests and posthoc pairwise comparisons of means were con-223ducted using the Šidák procedure. Mean proportions and asym-224 225 metrical standard errors were obtained by back-transformation from the logit scale. 226

227 2.5. Role of crawling insects in corolla damage

Preliminary observations suggested that damage to flowers in the form of holes pierced in corollas was effected by ants. Specimens of the ant species observed to be commonly responsible for corolla damage were collected for identification at the Iziko Museum, Cape Town. To assess the frequency of corolla damage by ants, the mean percentage of flowers per branch experiencing active corolla damage by ants (ants active around holes in the corolla) was calculated for the mature flowers on 10 235 branches on 10 randomly chosen plants throughout the 236 population. 237

To determine if holes in the corolla were made by flying in- 238 sects, such as bees, as was suggested by Rebelo et al. (1985), or 239 by crawling insects, such as ants, sticky Plantex[®] was applied 240 around the stems of ten freestanding flowering branches on 241 ten separate plants throughout the population to exclude all 242 crawling insects. Flowers were examined one week later to 243 quantify floral damage in the form of holes in corollas. 244 Untreated branches on the same plants were used as controls. 245 The proportion of flowers robbed for the two groups was ana- 246 lyzed using generalized linear models with a binomial error dis- 247 tribution. Plant was used as a blocking factor and was entered 248 in the model before treatment. Model significance was assessed 249 using likelihood ratio tests. Mean proportions and asymmetrical 250 standard errors were obtained by back-transformation from the 251 logit scale. 252

2.6. Nectar properties	253
------------------------	-----

Nectar was sampled from 114 flowers from 114 separate 254 plants throughout the population, using disposable, calibrated 255 1 ml micro-syringes. Nectar sugar concentrations were mea-256 sured using two Eclipse handheld refractometers (Bellingham 257 and Stanley Ltd.), one capable of taking measurements of up 258 to 50% and the other up to 80% sugar concentration. Nectar 259 sampling took place between 7 am and midday. Individual nec-260 tar samples were stored in separate eppendorfs, and four of 261 these samples later spotted onto four 7.0 cm Whatman filter pa-262 pers and dried for later determination of constituent free sugars 263 using HPLC methods, as described by Brown et al. (2009). 264

265
2

266

3.1. Pollinator observations

Birds thoroughly "worked" plants of *E. halicacaba*, often 267 concealed within the well-branched shrubs for up to 30 min 268 while visiting flowers. Movement from one plant to another 269 was observed during all fieldwork periods. Birds used the stur- 270 dy main and side-branches, as well as flowers, as perches. Birds 271 approached flowers from above, below and side-on, manipulat- 272 ing individual flowers, which are attached to the plant by a ro- 273 bust pedicel (± 3 mm long and 1 mm diam.). Birds were not 274 observed to rob flowers by piercing holes in corollas. Pollen 275 placement was on the lower third of the culmen (Plate 276 G) and also in a small depression at the base on the underside 277 of the lower culmen (Plate 278

Fourteen *A. violacea* individuals were netted on 26th November 279 2009. All fourteen birds had *E. halicacaba* pollen tetrads on their 280 culmens and the mean number of pollen tetrads per bird sampled 281 was 79 (range 20–200). The tetrads were assumed to be from 282 *E. halicacaba* since no other *Erica* species were flowering on 283 Glen Cairn Ridge at that time, and as the three other Cape gen-284 era known to have pollen tetrads were likewise absent, viz. 285 *Typha capensis* (Rohrb.) N.E.Br. (Typhaceae), *Cytinus* 286

ARTICLE IN PRESS

sanguineus (Thunb.) Fourc. (Cytinaceae), and Drosera capensis L. (Droseraceae) (Copenhaver, 2005). Pollen from only one
other plant species was recorded on all of the 14 culmens: yellow
monads of Saltera sarcocolla (L.) Bullock.

291 3.2. Selective exclusion experiments

Selective exclosure of flowering branches significantly de-292creased the frequency of flowers with tripped anther rings 293 $(\chi^2 = 274.1, df = 2, P = < 0.001;$ Fig. 2A). Each treatment was 294also significantly different, indicating a strong effect of the 295exclosures on rates of pollinator visitation. Thus 91.5% of 296 open inflorescences (controls) had flowers with tripped anther 297 rings; 31.7% of flowers in the chicken wire treatment group 298had tripped anther rings, birds presumably occasionally manag-299 ing to penetrate chicken wire exclosures; but only 1.5% of 300 flowers in the bridal veil treatment (all visitors excluded) had 301 tripped anther rings (these may have been disturbed when the 302 exclosure was applied) and only 0.7% of flowers (2 flowers) 303 set seed. Thrips were observed in several individual flowers but 304owing to their small size (< 2 mm long and <0.5 mm wide) 305 (Picker et al., 2004) they are not capable of displacing anther 306 rings of the larger southern African Erica species (Turner et al., 307 2011), especially those of the robust-flowered E. halicacaba. 308

Selective exclosure of flowering branches had a similar negative effect on the frequency of flowers that set seed $(\chi^2 = 287.1, df = 2, P < 0.0001; Fig. 2B)$ and all treatments were significantly different (Tukey test). Seed set occurred in 87.1% of open controls; in 17.2% of vertebrate-excluded flowers; and in 0.01% of the flowers from which all visitors were excluded.

316 3.3. Role of crawling insects in corolla damage

317No visits by flying insects, such as *Apis mellifera* L. (Hymenoptera), carpenter bees (Xylocopa spp., Hymenoptera), or blis-318 ter beetles (Meloidae), were observed during this study (\pm 96 319 hours). The main ant species observed to be responsible for co-320 rolla damage was identified as Myrmicaria nigra Mayr at the 321 Iziko Museum, Cape Town. M. nigra was observed to damage 322 flowers of E. halicacaba by chewing holes on the corolla 323 throughout the study population, during all fieldwork periods. 324 Tiny bird claw puncture holes in corollas, as a result of perch-325 ing, are enlarged and exploited by M. nigra in order to rob nec-326 tar. The mean (\pm SD) percentage of flowers actively robbed by 327 ants was 65 ± 5.0 . 328

Floral damage by crawling insects was significantly reduced by application of Plantex[®] to freestanding branches $(\chi^2 = 146.1, df = 1, p < 0.0001)$. The mean percentage of flowers pierced was 1.0 (upper se: 0.66; lower se: 1.36) for Plantex[®] treated plants versus 64.1 (upper se: 4.86; lower se 4.59) for control plants.

335 *3.4. Nectar properties*

The mean (\pm SD) nectar standing crop volume was 12.0 \pm 337 21.0 µl (range 1.25–100 µl). The mean (\pm SD) nectar sugar



Fig. 2. The effects of selective exclusion experiments on anther tripping (A) and fruit set (B) in *Erica halicacaba*. Values are back-transformed means and standard errors. Means that have different capital letters are significantly different.

concentration was $36.4 \pm 11.9\%$ (range 19-63.5%). The nectar 338 was sucrose-rich, with sucrose ($55.55 \pm 1.2\%$), glucose (17.2 ± 339 0.4%) and fructose ($25.8 \pm 0.6\%$). 340

4. Discussion

Data supporting the hypothesis that flowers of *E. halicacaba* 342 are adapted for pollination by Orange Breasted Sunbirds in- 343 clude repeated observation of legitimate pollination behaviour 344 with no observed robbery by birds; large numbers of pollen tet- 345 rads recorded on culmens of all mist-netted birds; high visita- 346 tion rates and seed set in open control flowers versus 347 vertebrate-excluded and bridal veil exclosures; and small vol- 348 umes of sucrose-rich nectar consistent with trends in other 349 flowers pollinated by sunbirds (Johnson and Nicolson, 2008). 350

341

6

ARTICLE IN PRESS

We showed experimentally that floral damage could be re-351duced to negligible levels (1% as opposed to 64.1% in open 352controls) by application of Plantex[®] to exclude crawling floral 353 predators, including Myrmicaria nigra ants, which we observed 354 and recorded to be responsible for extensive damage of flowers 355 in the wild (Plate $= \lim_{x \to \infty} E_x$). Ants are known nectar robbers 356 (Fritz and Morse, 1981; Haber et al., 1981; Herrera et al., 357 1984) and as such, may have neutral or negative effects on 358 plant fitness, influence floral evolution, have impacts upon 359 plant population demographics, and influence the stability of 360 established mutualisms (Irwin et al., 2001). In particular, Irwin 361 and Brody (1998) found that hummingbirds visited significantly 362 fewer plants in which "heavy" experimental robbing treatments 363 had been performed. While we did not study the impact of floral 364 365 damage upon visitation and seed set, open control treatments showed high visitation and pollination rates by A. violacea despite 366 presence of and damage by ants, suggesting that ants do not alter 367 the visitation dynamics of A. violacea to flowers of E. halicacaba. 368 Furthermore, while corolla damage by ants was significant in our 369 study population, the stochastic nature of floral robber-plant inter-370 actions (cf. Irwin and Brody, 1998) are such that the ex situ obser-371 vations by Rebelo et al. (1985) of bees as floral robbers may well 372 hold for other populations of E. halicacaba on the Cape Peninsula. 373 Nevertheless, our study shows that ants can cause extensive dam-374 age to the flowers of Cape plants. 375

Geerts and Pauw (2009) emphasised the correlation be-376 tween culmen length and flower length in "short-" and "long-377 billed" sunbird pollination syndromes. Mean corolla lengths of 378 *E.* halicacaba and *S.* sarcocolla flowers are 22.5 mm (n=25) 379 380 and 23.5 mm (15–28 mm; n=25) respectively, which correspond with reported A. violacea culmen lengths of 20-23 mm (Rebelo, 3811987, in Geerts and Pauw, 2009). Geerts and Pauw (2009) also 382 reported that short-billed sunbirds showed a tendency to rob lon-383 ger tubular flowers for which their culmen length was too short. 384 Our observations of zero robbery of E. halicacaba flowers by 385 A. violacea suggest that no mismatch of plant-pollinator mor-386 phology exists within the mutualism (de Merxem et al., 2009). 387

In our study, pollen was deposited on the lower third of cul-388 mens in all 14 netted birds (Plate = ig G). To reach nectar 389 without robbing, a bird could only enter through the adpressed 390 corolla lobes, tripping the anther ring in the process. This is in 391 contrast to pollination syndromes of hummingbird-pollinated 392 flowers (Castellanos et al., 2003), and to longer-tubed Cape 393 Erica and other species in which pollen is typically placed on 394 395 the head plumage of visiting birds (Geerts and Pauw, 2009; Johnson, 1995; Manning and Goldblatt, 2005). Of further inter-396 est concerning pollen placement is the small depressed area at 397 the base of the underside of the lower culmen (Plate =398 D): here we regularly (64% of netted birds) recorded an amal-399 gam of white E. halicacaba and yellow S. sarcocolla pollen 400 401 grains. Owing to the adpressed corolla lobes, repeated visits by birds to flowers result in pollen loads from previous visits 402being pushed towards the base of the bill, where they collect 403 in the excavated area. Hummingbird-pollination of short-tubed, 404 urn-shaped flowers of Brazilian Ericaceae also involves pollen 405 placement on the culmens of birds (Freitas et al., 2006). In the 406 407 light of the very few flowering bird-pollinated plant species on

Glen Cairn ridge during October and November, as well as our records of only two pollen types on culmens of mist-netted birds, we can deduce that the nectar of *E. halicacaba* and *S. sarcocolla* were an important energy source for resident Orange Breasted Sunbirds at that time.

We conclude that experimental field data and *in situ* obser-413 vation have been important in determining both the pollination 414 system of *E. halicacaba*, as well as the contribution of crawling 415 insects, especially ants, to flower damage. 416

417

423

Acknowledgements

We wish to thank South African National Parks for permis- 418 sion to perform ecological studies within the Table Mountain 419 National Park; Dr. Hamish Robertson of the Cape Town Iziko 420 Museum for identification of *Myrmicaria nigra*; and Esme 421 Beamish for assistance with mist-netting. 422

References

Anderson, B., 2005. Specialized bird perch aids cross-pollination. Nature 435, 424 41.

- Baker, H.G., Baker, I., 1983. Floral nectar sugar constituents in relation to pol-426
 linator type. In: Jones, C.E., Little, R.J. (Eds.), Handbook of Experimental 427
 Pollination Biology. Van Nostrand- Reinhold, New York, pp. 117–141. 428
- Baker, H.G., Baker, I., 1990. The predictive value of nectar chemistry to the 429 recognition of pollinator types. Israel Journal of Botany 39, 157–166.
 430
- Barnes, K., Nicolson, S.W., Van Wyk, B.-E., 1995. Nectar Sugar Composition 431 in Erica. Biochemical Systematics and Ecology 23 (4), 419–423. 432
- Brown, M., Downs, C.T., Johnson, S.D., 2009. Pollination of the red hot poker 433 *Kniphofia caulescens* by short-billed opportunistic avian nectarivores. 434 South African Journal of Botany 75, 707–712. 435
- Castellanos, M.C., Wilson, P., Thomson, J.D., 2003. Pollen transfer by hum- 436 mingbirds and bumblebees, and the divergence of pollination modes in 437 *Penestemon*. Evolution 57 (12), 2742–2752.
- Copenhaver, G.P., 2005. A compendium of plant species producing pollen tet- 439 rads. Journal of the North Carolina Academy of Science 121 (1), 17–35. 440
- De Merxem, D.G., Borremans, B., De Jäger, M.L., Johnson, T., Jooste, M., 441 Ros, P., Zenni, R.D., Ellis, A.G., Anderson, B., 2009. The importance of 442 flower visitors not predicted by floral syndromes. South African Journal 443 of Botany 75, 660–667. 444
- Fenster, C.B., Scott, W., Armbruster, Wilson, P., 2004. Pollination syndromes 445 and floral specialization. Annual Review of Ecology, Evolution, and Systematics 35, 375–403. 447
- Freitas, L., Galetto, L., Sazima, M., 2006. Pollination by hummingbirds and 448 bees in eight syntopic species and a putative hybrid of Ericaceae in South-449 eastern Brazil. Plant Systematics and Evolution 258, 49–61. 450
- Fritz, R.S., Morse, D.H., 1981. Nectar parasitism of Asclepias syriaca by ants: 451 effect on nectar levels, pollinia insertion, pollinaria removal and pod production. Oecologia 50, 316–319. 453
- Geerts, S., Pauw, A., 2009. Hyperspecialization for long-billed pollination in a 454 guild of South African plants: the Malachite Sunbird pollination syndrome. 455 South African Journal of Botany 75, 699–706. 456
- Geerts, S., Pauw, A., 2010. Easy technique for assessing pollination rates in the 457 genus Erica reveals road-impact on bird pollination in the Cape Fynbos, 458 South Africa. Austral Ecology. doi:10.1111/j.1442-9993.2010.02201.x. 459
- Haber, W.A., Frankie, G.W., Baker, H.G., Baker, S., Koptur, S., 1981. Ants 460 Like Flower Nectar. Biotropica 13 (3), 211–214. 461
- Hargreaves, A.L., Johnson, S.D., Nol, E., 2004. Do floral syndromes predict 462 specialization in plant pollination systems? An experimental test in an 463 "ornithophilous" African Protea. Oecologia 140, 295–301.
- Herrera, C.M., Herrera, J., Espadaler, X., 1984. Nectar thievery by ants from 465 southern Spanish insect-pollinated flowers. Insectes Sociaux 31 (2), 466 142–154. 467

ARTICLE IN PRESS

- Irwin, R.E., Brody, A.K., 1998. Nectar robbing in *Ipomopsis aggregata*: effects
 on pollinator behaviour and plant fitness. Oecologia 116, 519–527.
- Irwin, R.E., Brody, A.K., Waser, N.M., 2001. The impact of floral larceny on
 individuals, populations, and communities. Oecologia 129, 161–168.
 doi:10.1007/s004420100739.
- Johnson, S.D., 1995. Bird pollination in South African species of *Satyrium* (*Orchidaceae*). Plant Systematics and Evolution 203, 91–98.
- Johnson, S.D., Nicolson, S.W., 2008. Evolutionary associations between nectar
 properties and specificity in bird pollination systems. Biology Letters 4,
 49-52.
- Johnson, S.D., Steiner, K.E., 2000. Generalization versus specialization in plant
 pollination systems. Tree 15 (4), 140–143.
- Kleizen, C., Midgley, J.J., Johnson, S.D., 2008. Pollination Systems of *Colchicum* (Colchicaceae) in southern Africa: evidence for rodent-pollination. Annals of Botany 747–755.
- Kölreuter, J.G., 1761. Vorläufige Nachrichten von einigen das Geschlecht der
 Pflanzen betreffenden Versuchen und Beobachtungen. Gleditschischen
 Handlung, Leipzig.
- Manning, J.C., Goldblatt, P., 2005. Radiation of pollination systems in the Cape
 genus *Tritoniopsis* (Iridaceae: Crocoideae) and the development of bimodal
 pollination strategies. International Journal of Plant Sciences 166 (3),
 459–474.
- Olesen, J.M., Valido, A., 2003. Lizards as pollinators and seed dispersers: an
 island phenomenon. Trends in Ecology & Evolution 18 (4), 177–181.
- 492 Oliver, E.G.H., Oliver, I.M., 2000. Field Guide to the Ericas of the Cape Peninsula.
 493 National Botanical Institute. ISBN 1-919684-23-9.
- Oliver, E.G.H., Oliver, I.M., 2002. The genus *Erica* (Ericaceae) in southern Africa: taxonomic notes 1. Bothalia 32 (1), 37–61.

Soon and the second sec

496 Ollerton, J., 1998. Sunbird surprise for syndromes. Nature 394, 726-727.

- Ollerton, J., Alarcón, R., Waser, N.M., Price, M.V., Watts, S., Cranmer, L., Hingston, 497
 A., Peter, C.I., Rotenberry, J., 2009. A global test of the pollination syndrome hypothesis. Annals of Botany 103, 1471–1480.
- Pauw, A., 1998. Pollen transfer on bird's tongues. Nature 394, 731–732. 500
- Pauw, A., 2006. Floral syndromes accurately predict pollination by a special- 501 ized oil-collecting bee (*Rediviva peringueyi*, Melittidae) in a guild of 502 South African orchids (Coryciinae). American Journal of Botany 93, 503 917–926. 504
- Picker, M., Griffiths, C., Weaving, A., 2004. Insects of South Africa. Struik, 505 Cape Town. ISBN 1-77007-0613. 506
- Raimondo, D., Von Staden, L., Foden, W., Victor, J.E., Helme, N.A., Turner, 507 R.C., Kamundi, D.A., Manyama, P.A., 2009. Red List of South African 508 Plants. South African National Biodiversity Institute, Pretoria. Strelitzia 25. 509
- Raven, P.H., 1972. Why are bird-visited flowers predominantly red? Evolution 510 26, 674.
- Rebelo, A.G., 1987. Bird Pollination in the Cape Flora. In: Rebelo, A.G. (Ed.), 512 A Preliminary Synthesis of Pollination Biology in the Cape Flora. CSIR, 513 Pretoria, pp. 83–108. 514
- Rebelo, A.G., Siegfried, W.R., Oliver, E.G.H., 1985. Pollination syndromes of *Erica* 515 species in the south-western Cape. South African Journal of Botany 51, 270–280. 516
- Rodríguez-Gironés, M.A., Santamaría, L., 2004. Why Are So Many Bird 517 Flowers Red? PLoS Biology 2 (10), e350. doi:10.1371/journal.- 518 pbio.0020350 Published online 2004 October 12. 519
- Turner, R.C., Midgley, J.J., Johnson, S.D., 2011. Evidence for rodent- 520 pollination in *Erica hanekomii* (Ericaceae). Botanical Journal of the Linne- 521 an Society 166 (2), 163–170. doi:10.1111/j.1095-8339.2011.01139.x. 522
- Waser, N.M., Chittka, L., Price, M.V., Williams, N.M., Ollerton, J., 1996. Gen- 523 eralization in pollination systems, and why it matters. Ecology 77, 524 1043–1060. 525

526 Edited by JC Manning