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## NECTARY STRUCTURE AND NECTAR IN *SOBRALIA* AND *ELLEANTHUS* (SOBRALIEAE: ORCHIDACEAE)

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**ABSTRACT.** With approximately 200 species, the tribe Sobralieae is a dominant and common Neotropical group of orchids, yet little is known of variation in floral morphology as it relates to their pollination. As currently circumscribed, the tribe includes four genera that differ considerably in flower size and morphology: *Elleanthus*, *Epilyna*, *Sertifera*, and *Sobralia*. Although nectar-foraging pollinators are known for some species, the relationships of pollination to deceit and to nectar production are all poorly understood. We examined pollination-related of nectaries and nectar characteristics (presence/absence, volume, and concentration) for major clades of Sobralieae. Some species produce abundant nectar, but many species offer no reward. When present, nectar is secreted by thickened calli at the lip base. The cells of the nectariferous calli contain starch, which is rapidly converted to sugar during a brief anthesis (often lasting only one day). Most *Sobralia* flowers are relatively large, bee-pollinated, with a gullet-shaped lip, false nectary, large pollinia, and offer no reward. *Elleanthus* flowers are relatively small with a legitimate nectar reward, and most species are hummingbird-pollinated. Hummingbird-pollinated Sobralieae flowers are relatively small, brightly colored in the perianth and/or the subtending bracts, somewhat tubular, with a lip that forms a cup around the callus for storing nectar, and pollinia that are dark and relatively small.

**KEY WORDS:** Deceit, *Elleanthus*, nectar, nectary, pollination, *Sobralia*, Sobralieae

**Introduction.** In Orchidaceae, floral rewards are extremely diverse (van der Cingel, 2001; van der Pijl and Dodson, 1966) and include nectar, oils, resin, wax, food bodies, and even fragrances (Davies and Stpiczyńska, 2008a; Whitten *et al.*, 2007). By far, the most common reward is nectar, which is presented by flowers of varying morphology to many different pollinators: Diptera (flies), Hymenoptera (bees and wasps), Lepidoptera (moths and butterflies), Trochilidae (hummingbirds) in the New World (van der Cingel, 2001; van der Pijl and Dodson, 1966), Nectariniidae (sunbirds) (Johnson *et al.*, 1998) and Zosteropidae (white-eyes) in the Old World (Micheneau *et al.*, 2006).

However, it is estimated that one-third of all species of orchids use deceit strategies (Cozzolino and Widmer, 2005). This high percentage of such deceitful orchids is evidence that pollination by deceit is a

successful adaptive strategy. The evolutionary forces driving deceit pollination are complex and not well understood (Jersáková *et al.*, 2006).

In Sobralieae, known pollinators include various bees and hummingbirds. Some species produce nectar rewards, but others produce no apparent reward. Whereas food-foraging bees are attracted to flowers of diverse morphology, colors, fragrances, nectar guides, and nectar rewards, birds are attracted to nectariferous flowers with bright corollas and/or bracts of contrasting color, and that usually lack fragrances. Most investigated species of *Sobralia* Ruiz & Pav. are reported to be pollinated by a variety of large solitary bees, especially by euglossine bees, whereas hummingbird pollination is known in *Elleanthus* C. Presl and in a few *Sobralia* species (Braga, 1977; Dodson, 1962, 1965; Dressler, 1971, 1976, 2002; Ducke, 1902; Dzedzioch *et al.*, 2003; Fogden and Fogden, 2006; Roubik, 2000;

TABLE 1. Observations of nectar secretion in this study. Although some species were observed and confirmed to have nectar, not all had measurable amounts of nectar. Only sucrose was directly measured. Volumes are in microliters ( $\mu\text{L}$ ) and concentrations are in % sucrose (sometimes noted as  $^{\circ}\text{Bx}$ ). Additional species were sampled and produced nectar, but were too small to measure: *S. ciliata*, *E. lancifolius*, *E. graminifolius*, *E. fractiflexus*, and *E. robustus*.

Species	Sample size, flowers (n)	Mean (avg)	Standard deviation ( $\sigma$ )	Range	Vouchers	Syndrome
<i>Elleanthus aurantiacus</i>	5				none (population sampling)	hummingbird
volume		4.4	1.8	2-7		
concentration		22.8	1.9	21-26		
<i>E. caravata</i>	52				Neubig 202	hummingbird
volume		5.7	2.4	2-10.1		
concentration		24.3	6.9	12-40		
<i>E. cynarocephalus</i>	5				Neubig 247	hummingbird
volume		6	3.1	2-10		
concentration		9.2	6.6	5-21		
<i>E. sodiroi</i>	46				Neubig 246	hummingbird
volume		13.6	6.5	4-31.5		
concentration		15.9	6.4	7-25		
<i>Sobralia bouchei</i>	52				Blanco 3009, Neubig 208	bee
volume		14.1	7.9	2-43		
concentration		21.2	3.3	12-28		
<i>S. callosa</i>	27				Blanco 3021, Neubig 224	hummingbird
volume		6.3	2.4	1.5-12		
concentration		16.3	2	12-19.5		
<i>S. macrophylla</i>	6				Blanco 3022	bee
volume		4.9	2.4	1-8		
concentration		20.6	1.4	18-22		
<i>S. rosea</i>	46				none (population sampling)	bee
volume		8.4	8.6	3-35		
concentration		13.8	3.1	5-19.5		

Roubik and Ackerman, 1987; Singer, 2003; van der Pijl and Dodson, 1966). Molecular data demonstrate that *Sobralia* is not monophyletic (Neubig, 2012; Neubig *et al.*, 2011), and so understanding relationships with pollinators within a phylogenetic context is critical to develop hypotheses of evolution in pollination.

The objectives of this study are to document traits of nectary structure and nectar production relative to other morphological features in *Sobralia* and *Elleanthus* and to relate these features with pollen vectors.

**Materials and Methods.** Observations were primarily made on cultivated plants in greenhouses of the Florida Museum of Natural History over the course of May 2007 through May 2011 and

in Ecuagenera nurseries in Gualaceo, Ecuador, as well as on various natural populations in Ecuador, February 2009. Voucher specimens were deposited at FLAS and QCA herbaria. A list of taxa examined for nectar is presented in Table 1.

*Nectar Volume and Quantity* –. Flowers were examined for nectar presence/absence. If nectar was found, measurements were made of both volume and sucrose concentration. Sucrose concentration was measured with a 0-53 brix Atago refractometer at various times of the day, but primarily at midday and at approximately room temperature (Corbet, 2003). Concentrations are presented in percent sucrose (i.e., equivalent to Degrees Brix, g sucrose per 100 g solution), because

it is a common unit used in nectar and food science (Bolten *et al.*, 1979; Corbet, 2003; Dafni, 1992). Nectar was pipetted and measured with a 0.5–20  $\mu\text{L}$  Rainin micropipetter. Sugar composition and minor nectar constituents such as amino acids (Gottsberger *et al.*, 1984) were not examined in this study.

Most plants were cultivated in a closed greenhouse, and were therefore not exposed to insects or other potential pollinators that might remove nectar. All plants of *Sobralia rosea* and *Elleanthus aurantiacus* were sampled in the wild and could therefore have had their nectar removed (thus modifying nectar volume) by visiting pollinators. Alternatively, rain could have modified nectar volume and concentration in these species. However, except for visitation by pollinators, the occurrence of such factors was the subject of careful inspection and, as far as was possible, controlled experimentation.

**Floral Anatomy** –. Flowers of selected species were fixed in FAA (9 parts 70% ethanol: 0.5 part glacial acetic acid: 0.5 part commercial formalin) for several days and stored in 70% ethanol. Floral tissues were dehydrated in a graded tertiary butanol:ethanol:water series (6 h for each of the following solutions 20:50:30, 35:50:15, 55:45:0, 75:25:0, and two changes of 100% tertiary butanol). Dehydrated tissues were embedded in Paraplast® tissue embedding medium (melting point 56° C) and sectioned with an American Optical 820 rotary microtome at 10  $\mu\text{m}$ . Sections were attached to slides using Haupt's adhesive (1 g gelatin: 100 mL water: 2 g phenol: 15 mL glycerol) and allowed to dry at 30° C for 12 h. Tissues were treated in 3% ferric ammonium sulfate for 20 min, stained in 0.5% Heidenhain's iron-alum hematoxylin for 5–10 min, and counterstained with a 0.01% solution of safranin for 6 h. Stained tissues were dehydrated in a graded ethanol series (95%, 95%, 100%, 100%) for 5 min each and subsequently cleared in two changes of limonene. Coverslips were mounted onto slides using Permount. Observations and photographs were taken with a PixeraPro 150es digital camera attached to a Zeiss Axioskop 40 microscope. Additional hand-cut sections were made of flowers of various species of *Sobralia* and *Elleanthus* to demonstrate variation and the presence of nectaries and cavities. Entire flowers were cleared and/or hand-sectioned, then stained with

Lugol's solution ( $\text{I}_2\text{KI}$ : iodine - potassium iodide) to test for starch. Labella of mature flowers were also hand sectioned in the morning (7 am), at noon, and in the evening (7 pm) and stained with  $\text{I}_2\text{KI}$ . Hand-cut sections of fresh floral tissues were also stained with methylene blue (1% dissolved in  $\text{H}_2\text{O}$ ) for the purpose of indicting cavities and cellular contrast.

To examine cellular detail of the surface of calli, tissues were first pickled in FAA, then dehydrated in a graded ethanol series and dried in a critical point dryer using liquid  $\text{CO}_2$ . Dried samples were then mounted on clean aluminum stubs with double-sided adhesive graphite tabs. Mounted sections were coated with gold-palladium for approximately 60s in an argon vacuum. Sections were photographed digitally using a Hitachi S-4000 scanning electron microscope attached to a computer utilizing Spectrum Mono software.

**Results. Callus Structure** –. The labellar callus is probably not homologous throughout Orchidaceae but is apparently homologous within Sobralieae. The typical callus of most *Sobralia* species consists of two raised ridges borne opposite each other along the length of the labellum base. When seen from the distal end of the labellum, the space between the calli forms a narrow tube (Fig. 1C), which may guide the tongue of a visiting bee, channeling it to the double cuniculus (Fig. 1D–I, see later section for definition). The calli of *S. bouchei* and *S. callosa* differ from both of the previously mentioned types. They are fused and expanded to form a pad on the median portion of the base of the labellum (Figs. 2C, H–I and 3D–H).

In *Elleanthus*, the callus usually consists of two relatively large, globose masses at the base of the lip (Fig. 4D–F, 5 D–E). Exceptions include *E. caravata* and *E. robustus* in which the callus is approximately the same size, but is fused into a single structure (Fig. 4D–F). All investigated species of nectariferous Sobralieae produce nectar from large stores of starch present in the callus.

**Starch** –. All species contained at least some starch. However, the amount of starch and the thickness of the tissue containing the starch varied. In all of the nectar-secreting species, the pad-like or globosse callus contained abundant starch. In species that produced no nectar (*i.e.*, most species of *Sobralia*), the starch was less abundant, and often restricted to the epidermis of

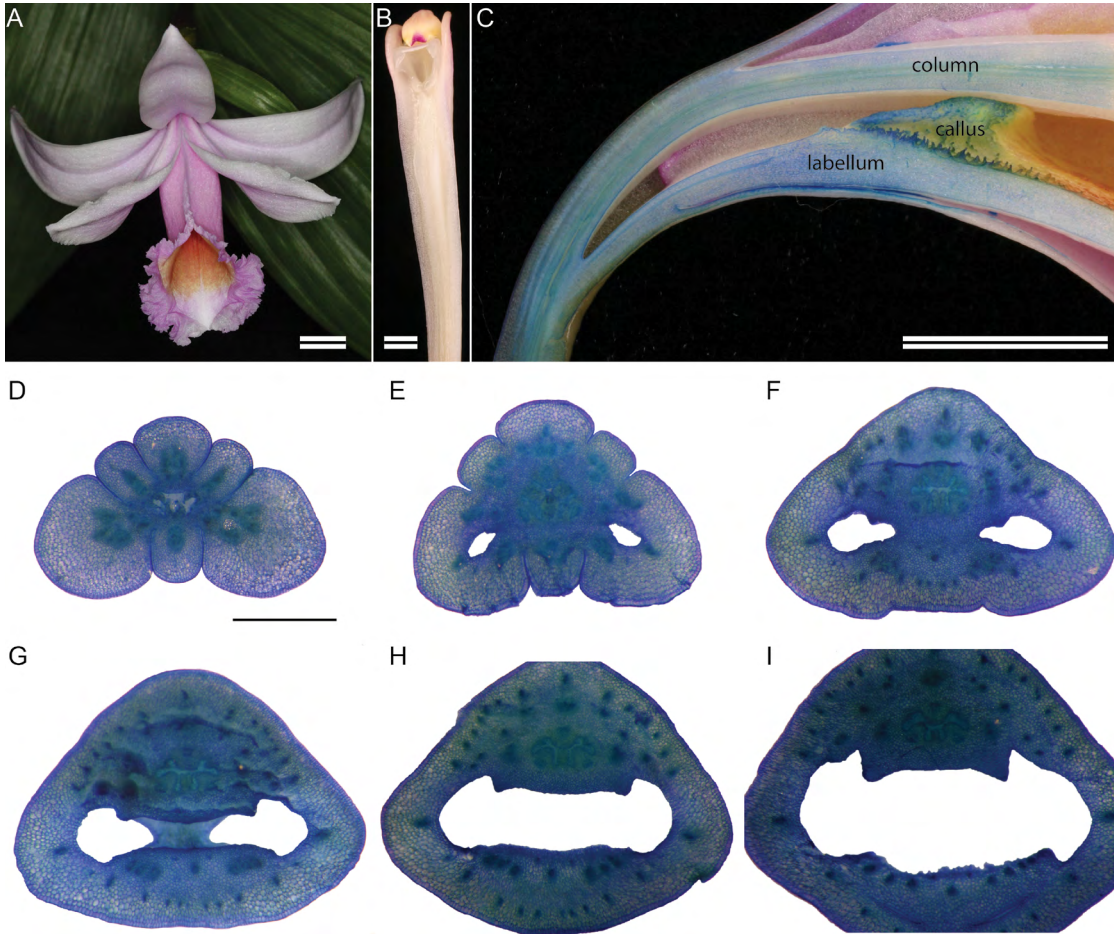


FIGURE 1. *Sobralia decora* (Whitten 3280) flower; a bee-pollinated flower with no nectar reward. All blue surfaces are stained with methylene blue. A. General floral morphology, scale bar = 1 cm. B. Column showing ventral surface with a common elastic rostellum which scrapes the pollinia from the scutellum as the bee exits a flower, scale bar = 3 mm. C. A longitudinal section of the flower, scale bar = 1 cm. Note the long ridged callus. D–I. Serial transverse sections of the pedicel, ovary, and perianth, scale bar = 1 mm. Note the two vacant spaces (double cuniculus) present between the sepals and the column fused to the lip; these form a pair of false nectar spurs.

the callus and epidermal trichomes. Sections of the callus made with a rotary microtome show amyloplasts that exhibited typical birefringent (cross-shaped) patterns when viewed with polarized light (Fig 5F).

**Double Cuniculus** – The double cuniculus is a novel term used here to describe the paired tubes formed between the ovary and the lateral sepals. This paired, tubular, false nectary comprising a double cuniculus was found in *S. chrysostoma*, *S. decora*, *S. gloriana*, *S. macrophylla*, *S. helleri*, *S. klotzscheana*, *S. powellii*, *S. warszewiczii*, and *S. sp.* Species lacking a double

cuniculus include *S. bouchei*, *S. callosa*, *S. crocea*, and *S. rosea*. No species of *Sobralia* sect. *Sobralia*, *Elleanthus*, *Epilyna*, or *Sertifera* examined has a double cuniculus.

**Nectary and Nectar** – Nectar sucrose concentration and volume were measured for four species of *Sobralia* (*S. bouchei*, *S. callosa*, *S. macrophylla*, and *S. rosea*; Table 1) and four species of *Elleanthus* (*E. aurantiacus*, *E. caravata*, *E. cynarocephalus*, and *E. sodiroi*; Table 1). The following species were observed to produce nectar, but the volumes produced were

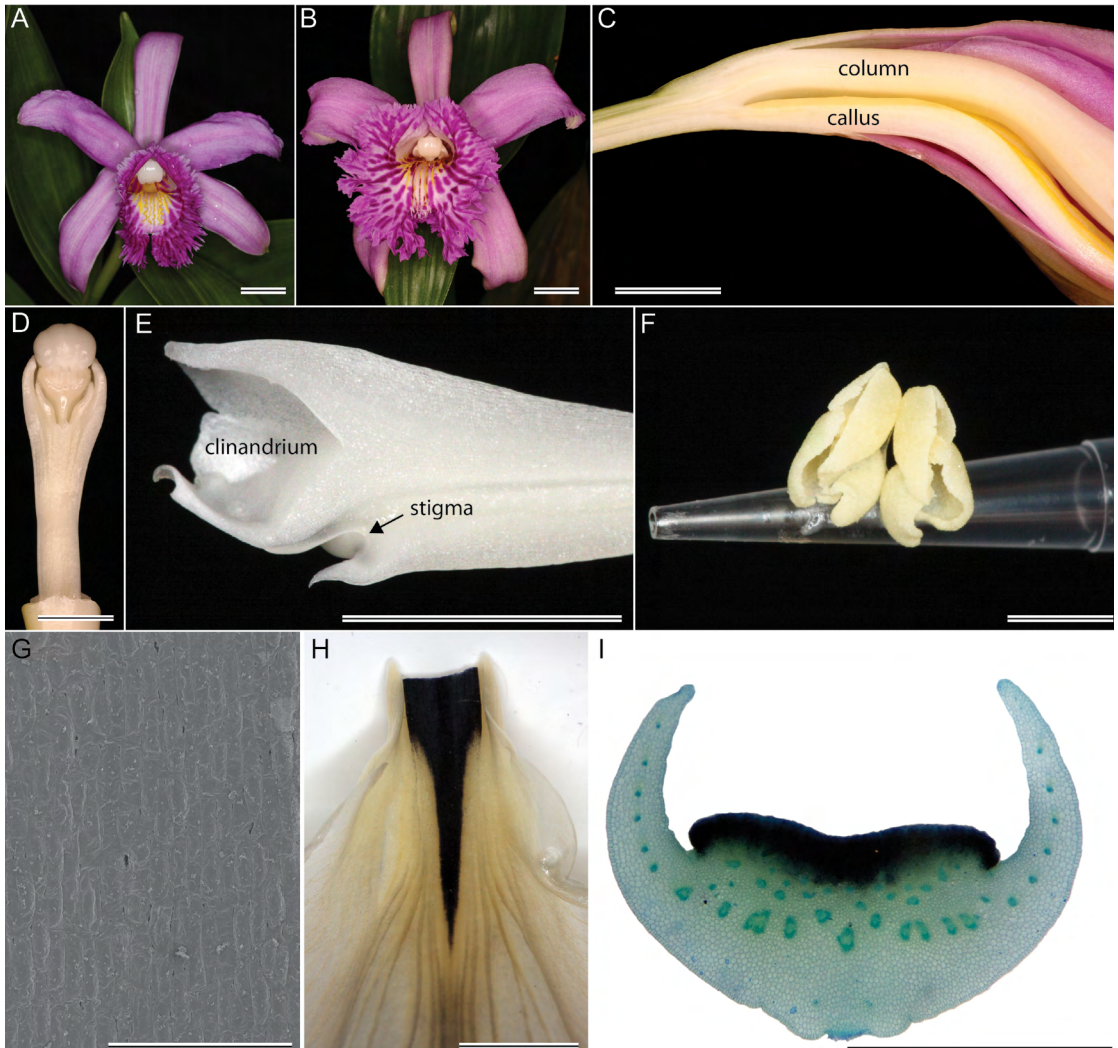


FIGURE 2. Flower of *Sobralia bouchei*, a bee-pollinated flower that produces nectar rewards. A. Frontal view of flower (Blanco 3009), scale bar = 1 cm. B. Frontal view of flower (Neubig 208), scale bar = 1 cm. C. A longitudinal section of the flower, scale bar = 1 cm. D. Ventral view of column showing the distinctive large anther cap and slit-like stigmatic surface differing from almost all other *Sobralia*, scale bar = 1 cm. E. The same column in longitudinal section with the anther removed, scale bar = 1 cm. F. Pollinia. G. SEM of the surface of the callus of lip, showing very different cellular surface texture compared to other *Sobralia* species, scale bar = 1 mm. Note the pores (intercellular spaces), which probably serve to increase surface area for nectar secretion. H. Basal portion of young lip, stained with  $I_2KI$  to indicate starch, precisely outlining the callus, scale bar = 1 cm. This starch is the putative carbohydrate source for nectar secretion. I. Transverse section of lip, showing the same callus with starch stained black from  $I_2KI$  while other tissues are stained with methylene blue solution, scale bar = 0.5 cm. This thick pad represents the fusion of the two distinct calli seen in most other members of the tribe.

too small to be measured: *S. ciliata*, *E. lancifolius*, *E. graminifolius*, *E. fractiflexus*, and *E. robustus*. The following species appear to lack nectar: *S. andreae*, *S. atropubescens*, *S. caloglossa*, *S. chrysostoma*, *S.*

*citrea*, *S. crispissima*, *S. crocea*, *S. decora* (Fig. 1), *S. dichotoma*, *S. doremiliae*, *S. exigua*, *S. gloriana*, *S. helleri*, *S. kerryae*, *S. lacerata*, *S. leucoxantha*, *S. lindleyana*, *S. macrantha*, *S. mandonii*, *S. mucronata*,

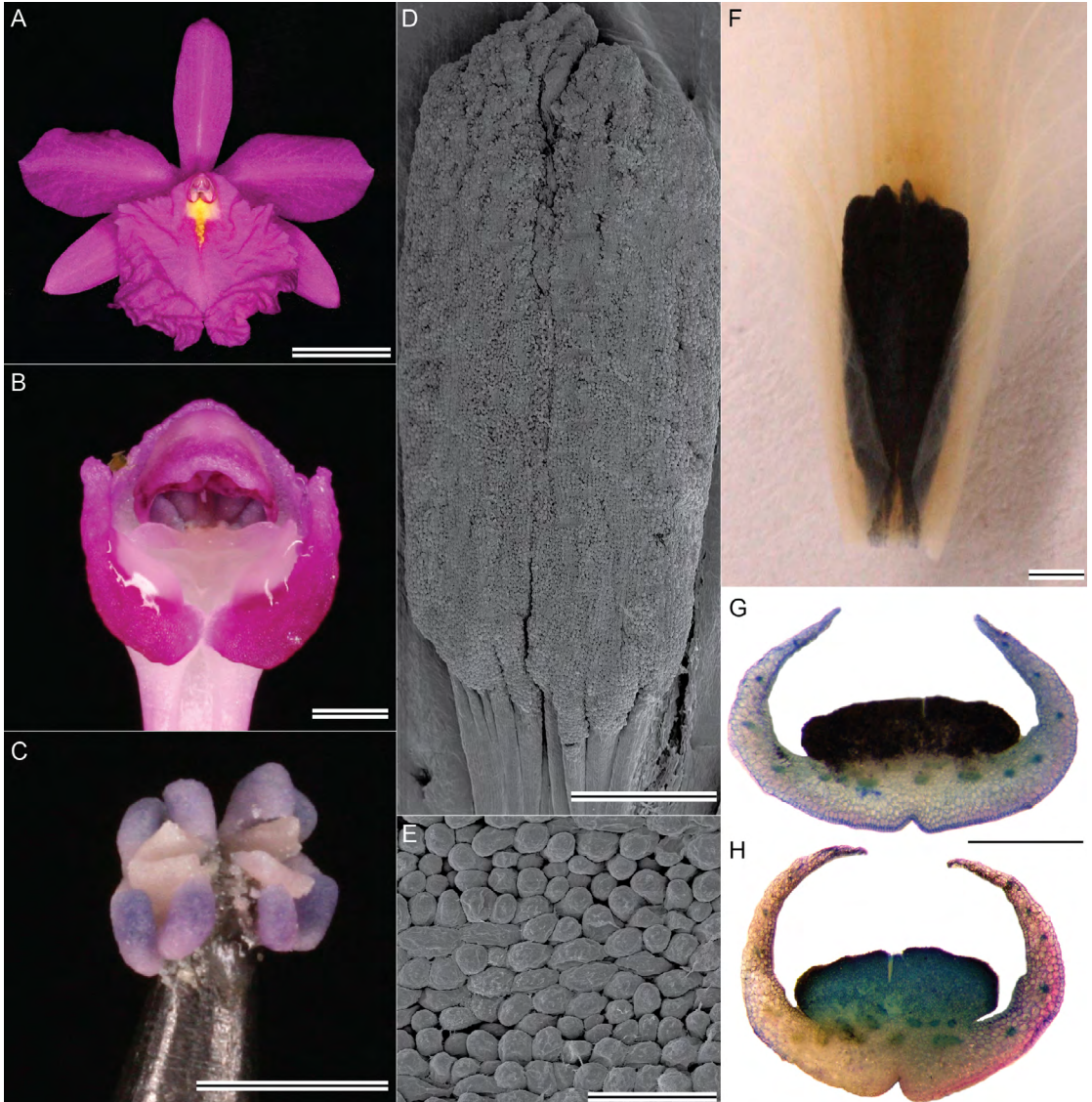


FIGURE 3. Flower of *Sobralia callosa* (Blanco 3021), a hummingbird-pollinated flower that produces nectar rewards. A. Frontal view of this flower, scale bar = 1 cm. B. Frontal view of the column, scale bar = 1 mm. Note the highly differentiated stigmatic orientation of anteriorly-facing surface which would require very different mechanical deposition during the pollination process; the pollinia would be scraped off during entry to the flower, and thus deposited on the stigma. C. Cryptic pollinia, scale bar = 1 mm. D. SEM of the whole callus, scale bar = 1 mm. E. Surface of the callus, showing very different cellular surface texture from *S. bouchei* (Fig. 2), scale bar = 100  $\mu$ m. Note the extremely papillose surface texture which probably serves to increase surface area for nectar secretion. F. Basal portion of young lip, cleared, then stained with  $I_2KI$  to indicate starch, precisely outlining the callus, scale bar = 1 mm. G–H. Transverse sections of the lip and stained with  $I_2KI$  to reveal starch in a young flower (morning) and an old flower (evening), respectively, showing the gradual reduction in starch over time, scale bar = 1 mm.

*S. quinata*, *S. recta*, *S. theobromina*, *S. violacea*, *S. warszewiczii*, and *S. yauaperyensis*. The only other

species of *Sobralia* reported to produce nectar, *S. amabilis*, was not investigated in this study.



FIGURE 4. Flowers of *Elleanthus caravata* (Neubig 202), a hummingbird-pollinated flower that produces nectar rewards.

A. Inflorescence showing the bright color contrast of bract and flower, typical of bird pollination, scale bar = 1 cm. B. Flower showing saccate base where nectar is secreted and stored, scale bar = 1 cm. C. Pollinia showing their relatively small size, dark color, and hard texture, scale bar = 1 mm. D. The callus of the lip in a young flower, stained with  $I_2KI$  indicating the presence of starch, scale bar = 1 mm. E. Longitudinal section of callus, scale bar = 1 mm. F. Transverse section of callus, scale bar = 1mm.

**Discussion.** *Anatomy of floral nectaries and starch* – In orchids, nectar is produced in a variety of structures, including spurs or nectaries derived from the lip callus. The callus is a term given to any raised or sculptured portion of the lip. Although the callus is probably not homologous within Orchidaceae, the ability to produce thickened tissue on various floral parts may

be an exaptation for secreting large amounts of a reward, either nectar or other compounds.

In all species of tribe Sobralieae, there are two calli at the base of the lip, but the calli vary in shape, size, and degree of fusion between species. Darwin (1862) first described the nectary structure of *Elleanthus* as large “balls”, referring to the callus at the base of



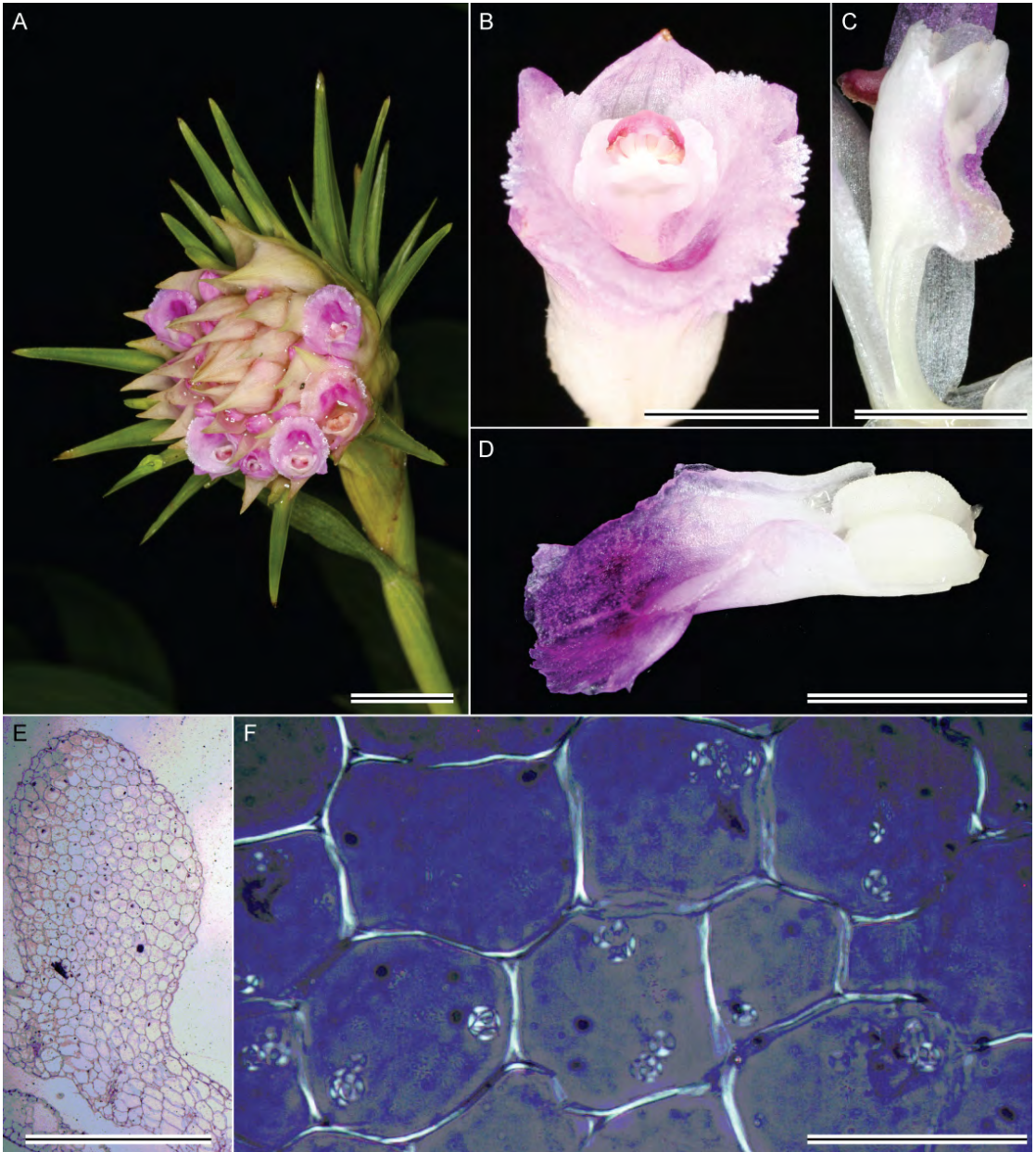


FIGURE 5. Flowers of *Elleanthus sodiroi* (Neubig 246) a hummingbird-pollinated flower that produces nectar rewards. A. Dense capitulate inflorescence, scale bar = 1 cm. B. Frontal view of flower showing the entrance point for the pollinator, scale bar = 0.5 cm. C. Oblique view of ventral surface of the column, scale bar = 0.5 cm. Note the median ridge of the column which forms a “pocket” with the lip. D. Lip of flower showing the two spherical calli at the base, scale bar = 1 cm. E. Transverse section of one callus, scale bar = 1 mm. F. Transverse section of callus under polarized light, scale bar = 10  $\mu$ m. Note the birefringent granules within each cell, indicating the presence of starch.

the labellum. Recently, the anatomy of the callus of *Elleanthus brasiliensis* (Lindl.) Rchb.f. was examined in detail (Nunes *et al.*, 2013) and the callus was

identified as the secretory structure. Veyret (1981), likewise, studied the floral and fruit morphology of *Elleanthus* and *Sobralia*. Veyret did not study the

nectary structure of *Elleanthus*, but did study the unusual fusion of floral parts in *Sobralia*. In that study, a novel structure, referred to as “éperon bifide,” or bifid spur, was identified in *Sobralia sessilis*. Our data indicate that many other species of *Sobralia* also have this bifid spur, which we refer to here as a double cuniculus, in reference to the very similar structure found in Epidendreae, although these structures are not homologous. The double cuniculus probably functions as a pair of parallel false nectaries or pseudo-nectaries for long-tongued insect visitors (*e.g.*, nectar-foraging euglossine bees). No nectar was ever observed within the double cuniculus, nor is the anatomy consistent with metabolically active secretory cells.

Treatment of floral sections with I<sub>2</sub>KI (Figs. 2H-I, 3F-H, & 4D), as well as visualization with polarized light (Fig. 5F), revealed the distribution of starch within floral tissues. The occurrence of starch in the callus is a constant feature among all flowers in the Sobralieae; however, the quantity of the starch is variable. Even species of *Sobralia* that do not secrete nectar will nonetheless accumulate small amounts of starch. Furthermore, the hydrolysis of floral starch might also provide the energy for fragrance production. Those species with the greatest accumulation of starch relative to flower size produce the most nectar. Nunes *et al.* (2013) did not report the presence of starch in *Elleanthus brasiliensis* calli, but these authors may have examined old flowers with exhausted starch reserves, or possibly starch was lost during tissue manipulation. Although their flowers are relatively small compared to those of *Sobralia*, the calli of *Elleanthus* are large compared with the size of their flowers (each being approximately 2-3 mm long) and densely packed with starch, at least prior to anthesis. *Sobralia* calli were more variable both in structure and in terms of starch content. *Sobralia* species that produce nectar (*e.g.*, *S. bouchei*, *S. callosa*, *S. macrophylla*, and *S. rosea*) have two calli that fuse together resulting in the formation of a thickened pad, which prior to anthesis, is densely packed with starch. This starch is no longer present later in the day (Fig. 3G-H) and thus, presumably, acts as a substrate both for nectar sugar production and as a source of metabolic energy for nectar secretion.

We conclude that the callus is the probable source of nectar in Sobralieae based on four observations: 1) In early stages of anthesis, droplets of nectar can be

seen to form directly on the surface of the callus (and not on any other tissues); 2) All nectariferous species have starch-filled calli during the first stages of anthesis; by the onset of floral senescence, the starch is largely exhausted; 3) These calli have a dense cytoplasm that is consistent with cells that move nectar directly through the cell wall; 4) No pores or stomata with underlying vascular tissue (typical of phloem-fed nectaries) were observed on the epidermis of the callus.

A nectariferous callus has been reported in other orchids, but the frequency and distribution of such a structure within the family is poorly documented. A callus that secretes nectar has been demonstrated for *Maxillariella anceps* (Ames & C. Schweinf.) M.A. Blanco & Carnevali (Davies *et al.*, 2005), *Stenorrhynchos* Rich. ex Spreng. (Galletto *et al.*, 1997), and in some other orchid groups (Davies and Stpiczyńska, 2008a). Many orchids are known to accumulate starch for various secretory purposes relating to pollination (*e.g.*, fragrance production in Stanhopeinae). Starch accumulation followed by depletion associated with nectar secretion also has been found in other orchids such as *Scaphyglottis* Poepp. & Endl. (Stpiczyńska *et al.*, 2005a), *Acianthera* Scheidw. (de Melo *et al.*, 2010), *Limodorum* L. (Figueiredo and Pais, 1992), *Epipactis* Zinn (Pais and Figueiredo, 1994), in multiple species in subfamily Orchidoideae (Galletto *et al.*, 1997; Stpiczyńska *et al.*, 2005b), and among other plant families (Durkee, 1983). Based on its ubiquity, it would appear that having a fixed reserve of starch is advantageous for the rapid production of floral secretions, whether they are fragrance (Curry *et al.*, 1991) or nectar.

The ultrastructure of floral nectaries (Fahn, 1979; Vassilyev, 2010), together with the transport and secretion of nectar (Pacini and Nepi, 2007) is generally well understood. In orchids, the anatomy of structures that secrete floral rewards (including nectaries, osmophores, elaiophores, and resin-secreting structures) has only been studied recently, and for only a small number of orchid species (Davies and Stpiczyńska, 2008a; Davies and Stpiczyńska, 2008b; Davies *et al.*, 2005; Stpiczyńska, 2003; Stpiczyńska *et al.*, 2003, 2005a; Stpiczyńska *et al.*, 2010). Based on these studies, floral secretions are produced by diverse anatomical structures. The secretion of nectar onto the surface of the flower can be achieved in two

main ways. The first is via stomata in the epidermis overlying the nectary. The second is via the cell walls of the epidermis. It is this latter method that seems to predominate in Sobralieae, because there are virtually no stomata or hairs on the nectar-secreting surface of the callus (Figs. 2G & 3D-E). The epidermis of most species that we examined (except for *S. bouchei*) were highly papillose, with no intercellular spaces to increase the surface area through which secretion could occur. In *S. bouchei*, the callus surface, as viewed using SEM, was relatively glabrous comprising brick-shaped cells with narrow intercellular spaces (Fig. 2G).

Some orchids (*Aerangis* Rchb.f. and *Platanthera* Rich.) and non-orchids (*Brassica napus* L.) have the ability to reabsorb the sugars secreted in unconsumed nectar (Burquez and Corbet, 1991; Koopowitz and Marchant, 1998; Stpicyńska, 2003). There is no evidence to support this type of reabsorption in Sobralieae.

*Nectar concentration and volume* –. Nectar concentration and volume are two traits that are thought to be linked to the class of pollinator (Baker and Baker, 1983). Hummingbird-pollinated flowers are thought to produce relatively large volumes of dilute nectar, whereas bee-pollinated taxa produce comparatively smaller volumes of more concentrated nectar (Bolten and Feinsinger, 1978; Hainsworth and Wolf, 1972, 1976; Pyke and Waser, 1981).

The sugar concentrations of nectar have been studied extensively for various angiosperm groups, but not for Sobralieae. Many studies have demonstrated that there are differences between the floral nectar of flowers having different pollinators. For example, the range of sucrose concentrations for solitary bee nectar is 16-50%, whereas that for hummingbirds is 13-30% (Baker, 1975; Baker and Baker, 1983). These ranges tend to overlap by a considerable margin and the immediate difference occurs only in the upper range of concentrations for bees. The largest difference between pollinators is the relative ratio of sucrose-glucose and fructose, but again, there is considerable overlap.

It has been suggested that hummingbird-pollinated flowers “never” have high ratios of glucose and fructose (*i.e.*, their nectar contains relatively high concentrations of sucrose; Baker and Baker, 1983). A

relatively high ratio of sucrose was found in a broad sampling of hummingbird-pollinated plants in Costa Rica (Stiles and Freeman, 1993). Therefore, there is a substantial degree of overlap in nectar volume and its sugar concentration relative to the type of pollinator. As hummingbird pollination is often a relatively derived condition within predominantly insect-pollinated groups (Beardsley *et al.*, 2003; Kay *et al.*, 2005), it is reasonable to assume that hummingbirds select for a specific type of nectar. More recently, the hypothesis of nectar preferences in hummingbirds has been challenged by more recent studies (Johnson and Nicolson, 2007). Similar trends in sugar ratios, as they relate to pollinators, have also been reported for *Ipomoea* (Galletto and Bernardello, 2004), as well as in other plant groups (Burke *et al.*, 2000; Galletto *et al.*, 1998), however, these studies found no significant differences in nectar composition between plants having different pollinators. Other surveys involving many unrelated plants have shown variable nectar concentrations for hummingbird-pollinated taxa (McDade and Weeks, 2004). Similarly, our observations show that sucrose concentration in Sobralieae is highly variable and were not related to pollination syndrome (Table 1; Figs. 6). We did not analyze the sugar composition of nectar nor the ratios of the individual sugars. Nevertheless, our data revealed differences in nectar volume between pollinator classes. Many of the hummingbird-pollinated species produced smaller volumes (perhaps because each inflorescence bears numerous small flowers), generally approximately 6  $\mu$ L per flower (except for *E. sodiroi*, which produced as much as 32  $\mu$ L nectar per flower). Bee-pollinated flowers, such as those of *S. bouchei* and *S. rosea*, produced more nectar, an average yield of 8.4-14.1  $\mu$ L per flower. Conversely, *S. macrophylla* produced very little nectar, and although we examined approximately 50 flowers from several different plants (n=6), only rarely was nectar observed.

The majority of documented pollinators of *Sobralia* are nectar-foraging euglossine bees (Apidae: Euglossini). The nectar viscosity of some euglossine bee-pollinated plants other than orchids has been studied relative to the length of the proboscis of the pollinating bee (Borrell 2005, 2006). Borrell (2007) also measured sugar concentrations from euglossine

bee crops and from various euglossine nectar sources and found that orchid bees harvest nectars with 34%–42% sucrose, independent of body size. Borrell (2006) speculated that long nectar spurs may be a mechanism by which flowers conserve nectar while remaining attractive to traplining bee visitors. Our analyses of *Sobralia* nectar produced lower values than those conducted by Borrell.

Bee-pollinated species of Sobralieae produced relatively low-viscosity nectar (*i.e.*, *S. bouchei*, *S. macrophylla*, and *S. rosea*), whereas species of *Sobralia* having deceit strategies produced no observable nectar. A larger sampling of *Sobralia* species that are bee-pollinated, yet produce nectar, would be difficult, since so few species of the genus produce rewards. Nevertheless, it is likely that at least a few, hitherto unexamined species, produce nectar, and Romero (1998) has reported the occurrence of pseudopollen in *S. liliastrum* Lindl., suggesting that both mimicry and rewards other than nectar may occur in certain species of this genus.

*Nectar Deceit* –. Many orchids have “gullet flowers” that produce no nectar, *e.g.*, *Cattleya* (Dressler, 1981) and *Cochleanthes* (Ackerman, 1983). *Sobralia*, like many food-deceit orchids, probably takes advantage of a general floral bauplan that is attractive to a wide variety of pollinators. This is termed generalized food deception (Jersáková *et al.*, 2006), and the mechanism is apparently frequent and sometimes referred to as pollinator naiveté (Ackerman, 1986). Most *Sobralia* species exhibit generalized food deception. Food deception based on generalized foraging behavior has been demonstrated for many orchids (see Jersáková, Johnson, & Kindlmann (2006) for a detailed list of such groups) and most *Sobralia* species exhibit this strategy. Narrow pollinator specificity also exists in many orchids (Schiestl and Schluter, 2009), but is not known for any pollination system found in Sobralieae.

Whether pollination is achieved by rewards or deceit, floral structural adaptation is necessary for effective pollination. Orchids have a plethora of structures for presenting nectar to pollinators, especially long-tongued insects. Some members of tribe Vandaeae (especially *Angraecum*) have long tubular spurs (formed from an invagination of the lip) that are associated with hawk moth pollination (van

der Cingel, 2001). In some orchids, a cuniculus is formed by the fusion of a hypanthium-like structure, as in the Laeliinae (*e.g.*, *Brassavola* R.Br.), and forms a single tube serving much the same function as the spurs in *Angraecum* (Stpiczyńska *et al.*, 2010). In several genera of Zygopetalinae, a gap at the base of the lip leads into a rolled, tubular backswept sepal that forms a false spur (Ackerman, 1983). Even though these structures may not be homologous, they all have a similar function, namely to facilitate pollination, either by deceit or through the production of a legitimate reward. Most species of *Sobralia* deceive the pollinator in that they have a ridged callus that forms a tube that serves as a funnel and guides the proboscis of the pollinator deep into the “double cuniculus” embedded within the ovary (Fig. 1).

The double cuniculus is unusual among orchids and is found only in part of the core group of *Sobralia* (Neubig, 2012; Neubig *et al.*, 2011). It comprises an open channel that runs between the lateral sepals and the ovary and can extend up to several centimeters into the latter (Fig. 1). This is perhaps the most significant feature of the double cuniculus. All the flowers having a double cuniculus that we examined offered no nectar reward, neither at the callus, as is typical of other Sobralieae, nor within this cunicular region, deep inside the ovary. Because *Sobralia* usually has a typical gullet-shaped flower (zygomorphic, with a tubular lip and nectar guides), and because it produces no nectar, this double cuniculus is interpreted as being a pair of false nectaries. This probably contributes to the effectiveness of the deceit, especially in the case of long-tongued bees, and in particular, Apidae (Danforth *et al.*, 2006). This interpretation is supported by the fact that the width of the individual tubes of the double cuniculus exceeds the width of the proboscis of known bee pollinators (*e.g.*, euglossines). We speculate that the deep double cuniculus induces the bee to probe further into the throat of the flower, thereby increasing the likelihood of effective pollination (Nilsson, 1988). Because long-tongued euglossine bees are the most commonly observed pollinators of *Sobralia*, this length-mediated deceit probably contributes significantly to pollinator selection.

Based on the fact that euglossines have the longest proboscises of any Neotropical bee subtribe, we hypothesize that any *Sobralia* species that possesses

cunicular tubes that penetrate deeply into the ovary is likely to be pollinated by nectar-foraging euglossine bees (male or female). Even nectariferous species, such as *Sobralia rosea*, have very large flowers with a particularly long, tubular throat (~5 cm), at the base of which occurs a true nectary favoring pollination by long-tongued bees.

*Future directions* – Detailed observations of floral morphology, anatomy, and secretions cannot substitute for careful field studies of pollination biology, but they may contribute to a hypothesis that can inform and prioritize fieldwork. The most glaring gaps in our knowledge relate to plant-pollinator relationships at the species level, especially for the white-flowered species of *Elleanthus* sect. *Elleanthus*, sect. *Chloidelyna*, and *Epilyna*. Verification of hummingbird pollination in other taxa, such as *Sertifera*, *Sobralia ciliata*, *S. callosa*, and *S. crocea*, is also critical for accurate interpretation of the number of modifications to this derived pollination syndrome.

The pollinators of the small, white-flowered species of *Elleanthus* and *Epilyna* are still not known. These species include *E. lancifolius* (sect. *Elleanthus*), all of sect. *Chloidelyna* (e.g., *E. fractiflexus*, *E. graminifolius*, *E. linifolius*, *E. poiformis*, and *E. stolonifer*), *E. caricoides*, and all of *Epilyna*. These flowers are even smaller than those of typical hummingbird-pollinated species, and have no bright colors, and therefore, it is highly unlikely that they attract or can be pollinated effectively by hummingbirds. These species have yellow pollinia and very small quantities of nectar (<1 µL), and it has been speculated that they are pollinated by small, nectar-seeking moths, such as

Noctuidae (C. Dodson, pers. comm.).

Most intriguing is the species *S. rarae-avis* (and the putatively closely related *S. madisonii* and *S. infundibuligera*, neither of which were examined morphologically in this study); their nocturnal fragrance is suggestive of pollination by hawkmoths or crepuscular bees, pollinator classes hitherto unknown for Sobralieae. The advent of increasingly cheap and portable digital video cameras should prove useful in documenting visits by pollinators.

By elucidating a greater number of specific plant-pollinator interactions for selected clades of Sobralieae, a more fine-tuned appreciation of the evolution of pollination-related floral features might be obtained, and recent molecular phylogenetic studies can be used to provide the evolutionary context for mapping such features (Neubig, 2012; Neubig *et al.*, 2011).

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