Comparative anatomy of the nectary spur in selected species of Aeridinae (Orchidaceae)

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INTRODUCTION

Aeridinae is a large and diverse subtribe containing some 1350 species in 103 genera distributed throughout the warm-temperate and tropical regions of Asia, Australia and eastern Pacific Islands, with Acampe Lindl. and Taeniophyllum Blume extending as far west as tropical Africa. Mostly epiphytes, members of this subtribe are characterized by their monopodial growth and highly developed velamen radicum. Pollinaria comprise two or four hard pollinia with obvious stipe and viscidium, and many genera possess a column-foot and spurred labellum (Topik et al., 2005, 2006). Although circumscription of Aeridinae has now been clearly defined (e.g. Cameron et al., 1999; Chase et al., 2003), prior to the use of molecular techniques (Topik et al., 2005, and references therein; Kocyan et al., 2008) our understanding of relationships within this subtribe was frustrated by morphological diversification and parallelism (Garay, 1972; Dressler, 1993).

Molecular, phylogenetic analyses tentatively support the monophyly of Aeridinae. However, the position of Sedirea Garay & H.R. Sweet is unresolved and it is claimed that molecular analyses and pollinaria studies indicate that it is sister to the whole subtribe and to the Phalaenopsis alliance, respectively (Topik et al., 2005, 2006; Kocyan et al., 2008). On the basis of phylogenetic analyses, Ascocentrum Schltr. and Papilionanthe Schltr. are currently assigned to the Aerides alliance and Schoenorchis Reinw. ex Blume is placed in the Pelatantheria alliance (Topik et al., 2005). Stereochilus Lindl. is also closely related to Aerides Lour. and belongs to the Stereochilus—Smitinandia—Vandopsis—Pomatocalpa—Trichoglottis clade (Kocyan et al., 2008).

Despite the enormity of Aeridinae and the morphological diversity of its flowers, generally, anatomical investigation of the nectary spur, which is so characteristic of many of its members, has been neglected. A short spur (<3 cm) is thought to represent the apomorphic state for this particular subtribe and both loss and elongation (>3 cm) of the nectary spur are considered to have occurred several times in its evolution (Topik et al., 2005). Spur length, however, although reflecting the overall length of the flower, is often related to the length of pollinator mouthparts (T. Yukawa, Tsukuba Botanical Garden, National Science Museum, Tsukuba, Japan, unpubl. res. -- cited in Topik et al., 2005).

In Aerides, two different flower types occur; those with open
apertures and those with hidden apertures to the nectary spur, and these appear to have evolved at least twice in geographically distinct regions. Even so, floral morphology of this genus is probably determined more by pollinator-driven selection than by phylogeny (Kocyan et al., 2008).

Reports of Aeridinae pollination are scarce. However, it appears that their pollinators are diverse and that these orchids employ a wide range of pollination strategies. Most species are entomophilous. For example, *Phalaenopsis* Ridl., *Vanda* Jones ex R. Br. and *Aerides odorata* Lour. are said to be pollinated by carpenter bees (*Xylocopinae*), and *Papilionanthe teres* (Roxb.) Schltr. by *Xylocopa latipes* (Carr, 1928; Dressler, 1990; van der Cingel, 2001; Kocyan et al., 2008). *Sarcochilus* R. Br. and *Pomatocalpa* Breda, Kuhl & Hasselt are pollinated by *Carbonaria* bees, whereas *Amesiella* Schltr. ex Garay and *Neofinetia* Hu are night-scented and possibly moth-pollinated (Ackerman, 1984; Dressler, 1990; van der Cingel, 2001; T. Yukawa, Tsukuba Botanical Garden, National Museum, Tsukuba, Japan, unpub. res. – cited in Topik et al., 2005). Beetles pollinate short-spurred or spur-less species, as well as *Peristeranthus* T.E. Hunt and *Trudelia cristata* (Vanda cristata Wall. ex Lindl.) and, in the case of *Cottonia* Wight., this may involve pseudocopulation (van der Cingel, 2001). Other genera, such as *Ascocentrum* Schltr. and *Renanthera* Lour., are said to be ornithophilous (Dressler, 1990; van der Cingel, 2001), and Slade (1980) reported the pollination of the hybrid orchid *Ascocentrum* ‘Sagrik Gold’ [A. miniatum (Lindl.) Schltr. × A. curvifolium (Lindl.) Schltr.] by an unidentified honeyeater (Meliphagidae). Autogamy is apparently common and occurs in *Schoenorchis paniculata* Blume, *Taeniophyllum hasseltii* Rchb.f. and a species of *Thrixspermum* Lour., whereas in *Cleisostoma parishii* (Hook.f.) Garay and *Papilionanthe longicornu* [Aerides uniflora (Lindl.) Garay], it is wind-assisted (van der Cingel, 2001).

In the present paper, we compare the anatomical organization of the nectary spur of a range of Aeridinae taxa (*Ascocentrum*, *Papilionanthus*, *Schoenorchis*, *Sedirea*, *Stereochilus*) that differ in flower size, relative spur dimensions and pollinator, as a prelude to a wider investigation of ornithophilous orchids in that their flowers are weakly zygomorphic, and that their pollen dimorphism is probably determined more by pollinator-driven selection than by phylogeny. Furthermore, we compare the nectary spur organization of ornithophilous *Ascocentrum* spp. with that of Neotropical, bird-pollinated *Maxillariinae* Benth., *Laeliinae* Benth. and *Oncidiinae* Benth. to determine whether these unrelated taxa exhibit pollinator-mediated convergence (Davies and Sticzynska, 2008, and references therein).

**MATERIALS AND METHODS**

Nectary spurs of seven species of Aeridinae were investigated: *Ascocentrum amplexiculum* (Roxb.) Schltr. var. aurantiacum Pradhan (accession no. KLD200701), *Ascocentrum curvifolium* (Lindl.) Schltr. (accession no. KLD200901), *Ascocentrum garayi* Christenson (accession no. KLD200702), *Papilionanthe vandarum* (Rchb.f.) Garay (accession no. S19990240), *Schoenorchis gemmata* (Lindl.) J.J. Sm. (accession no. KLD200703), *Sedirea japonica* (Rchb.f.) Garay & H.R. Sweet (accession no. S20070249) and *Stereochilus dalatensis* (Guillaum) Garay (accession no. KLD200704). Those specimens whose accession numbers are prefixed ‘S’ were obtained from the Swansea Botanical Complex, UK, whereas the remainder came from the second author’s collection. Abbreviations for authority names of plants follow Brummitt and Powell (1992). Nectar spurs were examined by light microscopy (LM), scanning electron microscopy (SEM) and transmission electron microscopy (TEM).

In each case, nectary tissue was fixed in 2.5% glutaraldehyde/4% formaldehyde in phosphate buffer (pH 7.4; 0.1 M) for 4 h at 4°C and carefully washed three times in phosphate buffer. It was then post-fixed in 1% osmium tetroxide solution at 0°C for 1.5 h, washed in distilled water and dehydrated using a graded ethanol series. Finally, material was infiltrated and embedded in LR White resin. Following polymerization at 60°C, sections were cut at 60 nm for TEM using a Reichert Ultracut-S ultramicrotome and a glass knife, stained with uranyl acetate and lead citrate (Reynolds, 1963), and examined using either an FEI Tecnai G2 Spirit Bio TWIN or Zeiss Leo 912AB transmission electron microscope, at an accelerating voltage of 120 kV.

Semi-thin sections (0.9–1.0 μm thick) were prepared for LM and stained with 0.25% toluidine blue O in 0.25% (w/v) aqueous sodium tetraborate solution (TBO). Hand-cut sections of fresh material were tested for the presence of starch and lignin with IKI solution and acidified phloroglucinol, respectively. Ruthenium red was used to test for the presence of acidic polysaccharides and mucilage, whereas alcoholic Sudan III was used to test for lipids (Jensen, 1962). Auramine O was also used to test for the presence of lipids, and the staining reaction was examined by means of a Nikon Eclipse 90i microscope equipped with fluorescein isothiocyanate filter. Autofluorescence of plastid chlorophyll in *P. vandarum* was investigated using a Nikon Optiphot II fluorescence microscope with UV-2B filter. In each case, control sections were used. The thickness of the spur wall was measured at its proximal end and micrometry and photomicrography were undertaken using a Nikon Eclipse 600 microscope with Screen Measurement version 4.21 software.

Fixed spurs were also cut longitudinally to examine the epidermis lining the lumen. They were subsequently dehydrated in acetone, subjected to critical-point drying using liquid CO₂, sputter-coated with gold and examined by means of a TESCAN/VEGA LMU scanning electron microscope, at an accelerating voltage of 30 kV.

**RESULTS**

Those members of Aeridinae investigated differed in both their floral morphology (including that of the spur) and their spur anatomy. Spurs varied in the width of the lumen, and the thickness of their walls and that of the outer tangential wall of the secretory epidermis. Moreover, whereas in some species the epidermis lining the lumen was glabrous, in others it was papillose, pubescent or hirsute, and there were also obvious differences in the structure of the epidermal cuticle and spur vasculature. The distribution of these characters and quantitative data are presented in Table 1.

**Ascocentrum**

*Ascocentrum* spp. have features characteristic of ornithophilous orchids in that their flowers are weakly zygomorphic,
**Table 1. Comparison of micromorphological characters (mean with range in parentheses) of the spur in selected Aeridinae**

<table>
<thead>
<tr>
<th>Character studied</th>
<th>Ascocentrum ampullaceum var. aurantiacum</th>
<th>Ascocentrum curvifolium</th>
<th>Ascocentrum garayi</th>
<th>Papilleanthe vandarum</th>
<th>Schoenorchis gemmata</th>
<th>Sedirea japonica</th>
<th>Stereochilus dalatensis</th>
</tr>
</thead>
<tbody>
<tr>
<td>Spur length (mm)</td>
<td>6.00 (5.69–6.30)</td>
<td>6.00 (5.73–6.27)</td>
<td>5.90 (5.66–6.20)</td>
<td>16.00 (15.94–16.12)</td>
<td>1.43 (1.30–1.58)</td>
<td>12.50 (8.00–14.00)</td>
<td>2.83 (2.76–2.92)</td>
</tr>
<tr>
<td>Thickness of spur wall (µm)</td>
<td>250.92 (223.51–330.51)</td>
<td>266.30 (249.70–281.60)</td>
<td>354.80 (245.80–457.50)</td>
<td>368.13 (285.05–454.45)</td>
<td>97.3 (68.00–167.80)</td>
<td>448.85 (379.30–590.00)</td>
<td>227.19 (140.20–285.11)</td>
</tr>
<tr>
<td>Number of vascular bundles</td>
<td>6–10</td>
<td>10</td>
<td>10</td>
<td>22</td>
<td>2</td>
<td>6–10</td>
<td>6</td>
</tr>
<tr>
<td>Dimensions of secretory epidermal cells (µm)</td>
<td>36.71 × 31.52</td>
<td>17.98 × 16.40</td>
<td>19.50 × 23.70</td>
<td>30.95 × 14.83</td>
<td>10.68 × 14.37</td>
<td>24.30 × 31.0</td>
<td>15.75 × 20.54</td>
</tr>
<tr>
<td>Thickness of outer cell wall of secretory epidermis (µm)</td>
<td>3.61 (2.17–3.36)</td>
<td>7.40 (6.10–8.90)</td>
<td>1.90 (0.80–2.10)</td>
<td>1.05 (0.81–1.37)</td>
<td>0.91 (0.40–1.29)</td>
<td>3.39 (2.90–4.13)</td>
<td>1.85 (1.04–2.33)</td>
</tr>
<tr>
<td>Presence of subepidermal collenchyma</td>
<td>+</td>
<td>+</td>
<td>+</td>
<td>–</td>
<td>–</td>
<td>–</td>
<td>–</td>
</tr>
<tr>
<td>Length of secretory hairs (µm)</td>
<td>37.42 (16.30–56.80)</td>
<td>46.22 (16.40–66.50)</td>
<td>28.12 (12.50–34.70)</td>
<td>260.55 (71.27–397.09)</td>
<td>–</td>
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<td>–</td>
</tr>
</tbody>
</table>
diurnal, lack fragrance and nectar guides, and are pink, orange or red with cryptic anther caps (Figs 1A, B, 3A, B and 5A, B).

Ascocentrum ampullaceum var. aurantiacum. This species has weakly zygomorphic, scentless, red–orange flowers with purple iridescence and orange–yellow column and spur. The labellum is simple and ligulate and uniformly orange–yellow (Fig. 1A, B). The entrance to the saccate spur is scarlet and laterally compressed (Fig. 1B, C). The anther cap is orange with a scarlet central region.

The wall of the spur is composed of several layers of cells. Short, unicellular hairs (mean length = 37.42 μm) occur predominantly in the middle part of the spur, whereas distally, the epidermis lining the lumen is glabrous and composed of flattened cells covered with non-striated cuticle (Fig. 1C–F). Nectar residues were observed on the surface of epidermal cells, especially those at the distended, distal part of the spur (Fig. 1E, F). The secretory hairs are densely distributed (Fig. 2A). Epidermal cells (both flattened cells and unicellular hairs) have thick cellulosic walls, and the outer, tangential

FIG. 1. Habit of the flower and micromorphology of the spur of *Ascocentrum ampullaceum* var. *aurantiacum*; (C–F) scanning electron micrographs. (A) Red–orange flower showing saccate nectary spur (arrows). Labellum marked with an asterisk. Scale bar = 2.5 mm. (B) Bisected flower showing relative positions of column, labellum (asterisk) and nectary spur (arrow). Scale bar = 2 mm. (C) Middle region of spur with secretory hairs. Scale bar = 150 μm. (D) Distal part of spur with glabrous epidermis. Scale bar = 300 μm. (E, F) Nectar residue on surface of glabrous and hirsute part of spur. Scale bars = 30 and 25 μm, respectively.

Key to abbreviations used in all figures: C, cuticle; CL, cuticular layer; CW, cell wall; ER, endoplasmic reticulum; D, dictyosome; m, mitochondrion; N, nucleus; n, nectar residue; P, plastid; Ph, phloem; st, starch; sv, secretory vesicle; V, vacuole; Vb, vascular bundle; Xy, xylem.
walls are particularly pronounced (Fig. 2B–G). The cuticle stains with Sudan III, TBO and auramine O (Fig. 2C–G), but with auramine O, fluorescence of the cuticle is much greater for the flat, epidermal cells than for the hairs (Fig. 2D). The nucleus is located in the basal part of the hair, and the parietal cytoplasm contains leucoplasts (Fig. 2B, G). Epidermal cells located at the bottom of the spur contain numerous vacuoles that stain purple with TBO.
indicating the presence of phenolic compounds. Similar compounds are also present in the small, thick-walled collenchyma cells that compose the one- to two-layered, subepidermal region (Fig. 2E, F). Primary pit-fields, containing numerous plasmodesmata, connect epidermal and subepidermal collenchymatous cells (Fig. 2H). Deep-seated parenchyma contains elongated idioblasts, whose vacuolar contents stain intensely with TBO. The spur is supplied with several collateral vascular bundles comprising xylem and phloem in equal proportions (Fig. 2E).

The secretory epidermis contains granular cytoplasm with numerous mitochondria, rough endoplasmic reticulum (RER).
profiles, dictyosomes and abundant secretory vesicles (Fig. 2I–K), as well as leucoplasts of irregular shape that contain small starch grains. The latter also occur in subepidermal cells (Fig. 2L).

Ascocentrum curvifolium. Ascocentrum curvifolium also has weakly zygomorphic flowers (Fig. 3A) lacking fragrance. All tepals are orange–red with purple iridescence (Fig. 3A, B). The throat of the flower, as well as the lateral lobes of the labellum, is yellow. The latter are inflated, permitting only a narrow channel into the spur. The mid-lobe of the labellum is immovable and arranged at 90° to the floral tube. Proximally, the nectary spur is yellow, but orange with purple iridescence distally (Fig. 3B).

As in A. ampullaceum var. aurantiacum, hairs up to 66.5 μm in length are found only at the middle part of the spur, the rest of the lumen being glabrous and lined by flattened epidermal cells (Fig. 3C–F). Nectar residues are present on the surface of the hairs and at the bottom of the spur (Fig. 3C, E, F).

In transverse section, the wall of the spur is fleshy and multilayered (Fig. 4A; Table 1). Small epidermal cells and hairs lining the lumen of the spur display extraordinarily thick, cellullosic walls and a thick cuticle (Fig. 4A–F). The cuticle is smooth, without visible pores and cracks (Fig. 3E). It stains intensely with TBO, Sudan III and auramine O (Fig. 4C–F) and, as in the previous species, it is the cuticle overlying flat, epidermal cells that fluoresces most (Fig. 4E, F). Cells comprising the 1–2 layers directly beneath the epidermis are collenchymatous, with thick walls (Fig. 4A–F). These walls are not lignified and stain intensely with ruthenium red. Epidermal and subepidermal cells have dense cytoplasm with relatively large nuclei (Fig. 4C). The granular, vesiculate cytoplasm contains numerous mitochondria and RER profiles.
Numerous small, collateral, vascular bundles, comprising equal proportions of xylem and phloem, occur in deeply seated parenchyma, whereas idioblasts containing phenolic compounds are found close to the outer epidermis.

Ascocentrum garayi. Flowers of *A. garayi* are yellow–orange with dark brown anther caps (Fig. 5A, B). They lack fragrance and nectar guides and the entrance to the spur, which contains copious nectar, is narrow (Fig. 5B, C). Nectar is secreted by trichomes arising from the epidermis lining the lumen of the saccate spur. They are present only in the middle part of the spur, where the latter becomes constricted (Fig. 5C–E). The rest of the lumen is lined with glabrous epidermal cells. The plastids do not accumulate starch (Fig. 6H). Cell walls of the secretary epidermis are relatively thick (mean = 1.9 μm), composed of cellulose and have a thick cuticular layer (Fig. 6B–E, G). The cuticle overlying both epidermal cells and trichomes is smooth but lamellate, without visible striations, and lacks pores and cracks. Although it stained with Sudan III, it did not fluoresce following treatment with auramine O (Fig. 6E, F).

Both epidermal cells and trichomes have dense, parietal cytoplasm and relatively large nuclei (Fig. 6A, B–D). The plastids do not accumulate starch (Fig. 6H). The nectary is supplied with collateral vascular bundles (Fig. 6A, C), but here, unlike the other *Ascocentrum* spp. investigated, phloem predominates. The ground parenchyma contains idioblasts whose vacuoles stain strongly with TBO (Fig. 6A, C) and sometimes these contain raphides.

**Papilionanthe vandarum**

The predominantly white flowers of *P. vandarum* are fragrant and are thought to be pollinated by moths. The
labellum is three-lobed and the spur is long, straight or curved, and slender (Fig. 7A, B). Long, unicellular, clavate hairs (up to 397.09 μm in length) are present throughout the spur. However, they are particularly long and more densely arranged along the inner, adaxial surface of the spur (Fig. 7C, D). Between these are occasional, bicellular hairs (Fig. 7E). Epidermal cells lining the spur lumen are small and covered by cuticle with evident striations. Striations are also present at the base of the hairs, whereas the cuticle covering the rest of the hair is almost smooth and lacks pores and distensions. Nectar residues occur on the surface of both hairs and epidermal cells (Fig. 7D, E).

In transverse section, the wall of the nectary spur is thick and multilayered (Table 1). Epidermal cells, and hair cells lining the lumen, contain parietal cytoplasm and numerous, but starchless, leucoplasts (Fig. 8A–G). The cuticle covering the secretory hairs and flat epidermal cells does not fluoresce with auramine O (Fig. 8H). Below the secretory epidermis are 2–3 layers of small, parenchymatous cells. Cellulosic cell walls are relatively thin (Fig. 8A–D) and contain numerous primary pit-fields with plasmodesmata that connect the epidermis and subepidermal parenchyma. The ground parenchyma contains numerous, large vascular bundles with prominent phloem (Fig. 8A).

Epidermal cells and hairs contain dense cytoplasm with abundant mitochondria and vesicles (Fig. 8I–L). Vesicles are also present in the periplasmic space (Fig. 8J). As well as typical chromoplasts, large leucoplasts with very densely and regularly packed membranes and numerous plastoglobuli, or peripherally situated osmiophilic regions, may be present (Fig. 8I–L). Vacuoles may also contain small, dark precipitates and vesicles (Fig. 8I).

**Schoenorchis gemmata**

The white mid-lobe of the labellum predominates in this very small, fragrant, rose–purple flower. Sepals are white or purple with white apices, and the spur and lateral lobes of the labellum are white, marked with purple. This species is entomophilous, the labellum forming a platform for alighting.
pollinators. The nectary spur, although short, is proportionally large and has a narrow entrance (Fig. 9A, B). It is straight, but expanded proximally. The column is very short and fleshy. The spur wall is significantly thinner than that of other species investigated (Table 1). The epidermis lining the lumen is glabrous, and the cuticle is distended (Figs 9B, C and 10A–E). These distensions occur on the cell surface, but are particularly abundant at cell junctions (Figs 9C and 10B–E). Epidermal cells, like the underlying parenchyma, are small with dense cytoplasma and large nuclei, leucoplasts and several small vacuoles (Fig. 10A–C, F). The spur is supplied by only two vascular bundles, each containing more phloem than xylem (Fig. 10A–C, F). Ground parenchyma contains numerous idioblasts with phenolic contents and/or prominent raphides (Fig. 10C–E). TEM shows that leucoplasts have an electron-dense stroma, with small starch grains (Fig. 10F). The thin, cellulosic wall has a reticulate cuticular layer, as well as a layer that becomes detached (Fig. 10G).

The cuticle does not fluorescence on treatment with auramine O (Fig. 10E).

Sedirea japonica

Like all the other entomophilous species investigated, flowers of Sedirea japonica are more strongly zygomorphic than those of Ascocentrum spp. They have pale, greenish white tepals and column. However, whereas the dorsal sepal and petals are entirely greenish white, bases of lateral sepals are barred purple–brown. The white labellum is marked at the margin with rose-coloured blotches that perhaps function as nectar guides, and is developed to a greater extent than the labella of Ascocentrum spp., the mid-lobe being expanded distally (Fig. 11A, B). The greenish white spur is approximately twice as long as it is wide at its widest point, and forwardly curving. The anther cap is cream-coloured. Flowers are very fragrant by day.

Fig. 7. Habit of flower and spur of Papilionanthe vandarum; (C–E) scanning electron micrographs. (A, B) Zygomatic, white flower with cream anther cap and curved nectary spur (arrow). Scale bars = 5 mm. (C) Spur lumen showing relative position of secretory hairs. Scale bar = 0.5 mm. (D) Secretory hairs with nectar residues on their surface and epidermal cells with striated cuticle (arrows). Scale bar = 80 μm. (E) Bicellular, secretory hairs with nectar residues. Scale bar = 25 μm. Abbreviations: see Fig. 1.
Fig. 8. Histology and ultrastructure of spur of *Papilionanthe vandarum*: (A–H) light micrographs; (I–L) transmission electron micrographs. (A) Section of spur wall showing epidermis, underlying tissues and vascular bundle. Scale bar = 50 μm. (B) Glabrous epidermis cells with nectar residues upon its surface (arrow). (C) Transverse section through epidermis and secretory hairs showing parietal cytoplasm with plastids. (D) Epidermal cells with striated cuticle (solid arrows). Pit-fields are present in epidermal and subepidermal cells (open arrows). (E) Epidermis with secretory, unicellular hair cut longitudinally and showing smooth cuticle. (B–E) Scale bars = 20 μm. (F) Unicellular, secretory hairs with large nuclei and numerous plastids in parietal cytoplasm. (G) Hairs showing autofluorescence and that plastids lack chlorophyll. (F, G) Scale bars = 25 μm. (H) Secretory hair following treatment with auramine O, yet showing no fluorescence. Scale bar = 25 μm. (I) Transverse section of hair showing cytoplasm with mitochondria and plastids. Vacuole contains vesicles and dark precipitates. Scale bar = 4 μm. (J) Plastid with numerous, dense lamellae and osmiophilic regions. Scale bar = 1.5 μm. (K) Detail of cytoplasm with nucleus and chromoplasts. Scale bar = 1 μm. (L) Parietal cytoplasm with plastid, mitochondria and secretory vesicles. Scale bar = 1.5 μm. Abbreviations: see Fig. 1.
The epidermis lining the spur is glabrous to minutely papillose (Figs 11C–F and 12A–F). These epidermal cells are elongate, and the centrally placed, conical papillae (Fig. 11E, F) stain with TBO (Fig. 12A–C). The cuticle is finely striated (Fig. 12C), but lacks pores (Fig. 12D–F). It stains intensely with Sudan III, but only slightly with auramine O (Fig. 12E, F). Cell walls, in particular radial walls and outer tangential walls adjacent to the lumen, are thick (Fig. 12B, C). The cytoplasm of epidermal cells and underlying parenchyma contains many large chloroplasts (Fig. 12C–E). Large idioblasts, close to the outer epidermis, stain strongly with TBO and often contain raphides (Fig. 12A). The spur is supplied with several large, vascular bundles and phloem predominates. Generally, xylem parenchyma occurs in each vascular bundle, but only 4–5 small, xylem vessels are present (Fig. 12B).

TEM reveals that the cytoplasm is granular, with abundant mitochondria, RER profiles, dictyosomes and vesicles that accumulate in the outermost cytoplasm (Fig. 12G, H). Leucoplasts have few internal membranes, and osmiophilic precipitates can be found in vacuoles. Primary pit-fields with plasmodesmata are frequent in the walls of the epidermis and adjoining parenchyma (Fig. 12G).

**Stereochilus dalatensis**

This entomophilous species produces simple racemes of small, fragrant, pale pink flowers with rose-coloured labella. The tepals are strongly reflexed and the column is vertical, so that the angle between it and the more or less horizontal, fleshy labellum approaches 90°. The anther cap is rose-coloured, whereas the nectary spur is white, flushed pink, relatively wide, saccate, fleshy and conical (Fig. 13A).

In *S. dalatensis*, a median septum divides the nectary spur longitudinally into two loculi or lumina (Figs 13B, C and 14A). A small, proximal protuberance is present in the loculus (Fig. 13B). The epidermis lining the lumen is glabrous and covered with a smooth cuticle. Where rows of adjacent epidermal cells run parallel to each other, the cuticle often becomes distended, and this coincides with the position of longitudinal walls, thus emphasizing cell shape (Fig. 13D, E). Furthermore, nectar residues may be present on the surface of epidermal cells (Fig. 13E).

The wall of the spur is multilayered and relatively thick (Fig. 14A; Table 1). The spur is supplied by six, collateral vascular bundles, with equal development of xylem and phloem (Fig. 14B). Epidermal cells have a thick outer wall that is heavily cuticularized (Fig. 14C–G), but the cuticle does not
fluoresce with auramine O (Fig. 14F). These cells contain parietal cytoplasm with mitochondria, vesicles, free ribosomes and ER profiles, but dictyosomes are seldom encountered. Large nuclei and amyloplasts are frequently present (Fig. 14C–E), and chromoplasts occur in the underlying parenchyma. TEM observations confirmed the presence of cutinized, reticulate layers in the outer, tangential cell wall of epidermal cells, and that the distended cuticle lacks pores and cracks (Fig. 14G).

A ‘Saftdecke’ or nectar cover to the spur appears to be present in Ascocentrum garayi, Schoenorchis gennata and Stereochilus dalatensis (Figs 5C, 9B and 13B).

**DISCUSSION**

Nectary spurs of the species investigated differ greatly in shape and length, from very short, saccate spurs in Stereochilus and Schoenorchis, moderately long spurs in Ascocentrum and Sedirea, to long spurs in Papilionanthe. Furthermore, spur morphology is considered taxonomically significant (Kocyan et al., 2008), with spur length being correlated with that of the mouth-parts of the pollinator. Remarkably, the presence of a spur, even in deceptive species lacking a reward, can be sufficient to attract pollinators (Bell et al., 2009).

Species of Ascocentrum possess weakly zygomorphic, red or orange flowers with cryptic anther caps. Their flowers lack nectar guides and fragrance. These features are characteristic of ornithophilous species (van der Pijl and Dodson, 1969; Proctor and Yeo, 1973; Proctor et al., 1996; van der Cingel, 2001; Ortega-Olivencia et al., 2005; Cronk and Ojeda, 2008; Davies and Stpiczynska, 2008, and references therein; Stpiczyńska et al., 2009). Cryptic anther caps may also facilitate pollination. The presence of
anther caps and pollinaria on beaks of birds usually evokes a bill-cleaning response and thus pollinia are often lost or destroyed. However, many ornithophilous species (some 50% of hummingbird-pollinated taxa) have blue, grey, brown, cream or greyish white, cryptic anther caps, and these are thought to illicit a lesser response than their more conspicuous, yellow counterparts (Dressler, 1971; Topik et al., 2006). As a result, transfer of pollinaria to the stigma is more likely. Moreover, the nectary of all Ascocentrum species studied has a collenchymatous subepidermis, similar to that found in other ornithophilous species (Stpiczyn’ska and Davies, 2006; Stpiczyn’ska et al., 2004, 2005, 2009). Thus, morphologically, anatomically and in terms of colour [which in A. ampullaceum var. aurantiacum and A. curvifolium resembles the floral colour combinations of presumed hummingbird-pollinated Ornithidium coccineum (Jacq.) Salisb. ex R. Br., O. sophronitis Rchb.f. and Hexisea imbricata (Lindl.) Rchb.f. (Stpiczyn’ska et al., 2004, 2005, 2009)], all the Ascocentrum species studied are thought to be ornithophilous. Unfortunately, direct observations of bird pollination are rare. Nevertheless, Slade (1980) recorded an unidentified honeyeater pollinating flowers of the hybrid Ascocentrum ‘Sagarik Gold’ growing in his garden in Vanuatu (New Hebrides), and Whitten et al. (2007) assert
that certain brightly coloured species of *Ornithidium* (Salisb.) ex R. Br. are indeed bird-pollinated.

Conversely, zygomorphy was more pronounced in the flowers of all other species investigated. They were all fragrant, with a proportionally more expanded mid-lobe to the labellum than has *Ascocentrum*. They lacked red pigmentation, cryptic anther caps and subepidermal collenchyma, and this is consistent with characteristics of entomophilous species (van der Pijl and Dodson, 1969; van der Cingel, 2001; Davies and Stpiczyńska, 2008).

Their nectary spurs showed considerable variation in length. Topik *et al.* (2005) reported that short-spurred species of Aeridinae are mainly pollinated by bees and beetles, whereas those species having long spurs are generally pollinated by moths. Petaloid spurs are also present in other plant groups such as *Aquilegia* L. and *Impatiens* L., and it has been shown that intra- and inter-specific variation in the spur length of *Aquilegia* is an adaptation for pollination by a variety of different pollinators (Hodges, 1997; Kramer and Hodges, 2010).

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**Fig. 12.** Histology and ultrastructure of spur of *Sedirea japonica*: (A–F) light micrographs; (G, H) transmission electron micrographs. (A) Section through nectary spur showing lumen lined with glabrous epidermis and vascular strands embedded in parenchyma. Scale bar = 100 μm. (B) Detail of spur wall with secretory epidermis and vascular bundle. Scale bar = 50 μm. (C) Epidermal cells with thick radial walls and fine cuticular striations (arrow). (D) Hand-cut section treated with IKI, showing epidermal and subepidermal cells containing chloroplasts. (E) A similar section stained with Sudan III and showing epidermis with thick cuticle. (C–E) Scale bars = 20 μm. (F) Weakly fluorescent cuticle covering secretory epidermis. Scale bar = 25 μm. (G) Parietal cytoplasm of secretory epidermis and subepidermal cell containing ER profiles, mitochondria, dictyosomes, secretory vesicles and plasmodesmata in cell wall (arrow). Scale bar = 1 μm. (H) Detail of parietal cytoplasm of secretory epidermal cell containing plastids, dictyosomes and mitochondria. Scale bar = 0.5 μm. Abbreviations: see Fig. 1.
Consequently, it is speculated that the species of *Stereochilus*, *Schoenorchis* and *Sedirea* investigated here, all of which are brightly coloured in shades of purple, pink and white, and all of which produce nectar and fragrance by day, are mainly pollinated by hymenoptera. Flowers of *Papilionanthe vandarum*, however, are mainly white, with relatively long spurs, indicating that this species is probably pollinated by moths (van der Pijl and Dodson, 1969; van der Cingel, 2001).

A ‘Saftdecke’ or nectar cover, often formed by the lateral compression of the spur (Sprengel, 1793 – cited in Kocyan et al., 2008), appears to be present in certain species, including both ornithophilous (e.g. *Asocentrum garayi*) and entomophilous (e.g. *Schoenorchis gemmata* and *Stereochilus dalatensis*) taxa. Once thought by Sprengel to protect against raindrops, these nectar covers have since been interpreted by others as protection against nectar thieves or evaporation.

Of the species investigated, only *Stereochilus dalatensis* has a bilocular nectary spur, the loculi or lumina being separated by a median, longitudinal septum whose function remains unknown. In all other cases, the spur was unilocular. The inner surface of the nectary spurs of *Schoenorchis gemmata*, *Stereochilus dalatensis* and *Sedirea japonica* is glabrous to minutely papillose. Spurs possessing a glabrous inner epidermis have also been recorded for other bee-pollinated species, such as *Aerides crassifolia* C.S.P. Parish ex Burb. (Kocyan et al., 2008). Moreover, *S. dalatensis* has a protuberance within the lumen, similar

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Fig. 13. Inflorescence and spur of *Stereochilus dalatensis*; (B–E) scanning electron micrographs. (A) Zygomorphic, pink and rose-coloured flowers with saccate spurs (arrows). Scale bar = 5 mm. (B) Bisected spur showing two loculi (lumina) separated by a median, longitudinal septum with proximal protuberance (arrow). Scale bar = 0.5 mm. (C) Detail of the two spur lumina lined with glabrous epidermis. Scale bar = 200 μm. (D) Epidermal cells showing distension of cuticle at cell junctions (arrows). Scale bar = 50 μm. (E) Secretory epidermis showing distension of cuticle (arrows) and nectar residues. Scale bar = 10 μm. Abbreviations: see Fig. 1.
to that described for certain species of *Aerides* (Chen and Wood, 2009). By contrast, the internal surface of the nectary spur of *Ascocentrum* species and *Papilionanthe vandarum* is pubescent to hirsute. These hairs are thought to increase the total surface area for nectar secretion and re-absorption (Davies and Stpiczynska, 2008, and references therein). That trichomes can occur within the spurs of certain rewardless species, such as *Dactylorhiza fuchsii* (Druce) Soó and *Barlia robertiana* (Loisel.) Greuter (Matthews et al., 2009; M. L. Matthews, Institute of Systematic Botany, University of Zurich, Switzerland, pers. comm., 2010), is thus of great interest. According to Bell et al. (2009), in deceptive species, such trichomes or papillae can provide tactile cues for pollinators. As in many orchids (Stpiczynska, 1997, 2003; Stpiczynska and Matusiewicz, 2001; Davies and Stpiczynska, 2008; Bell et al., 2009; Matthews et al., 2009), the nectary trichomes of *Ascocentrum* species are unicellular and conical, whereas those of *Papilionanthe* are clavate and occasionally bicellular. Bicellular or multicellular trichomes also occur in the spurs of several species of Orchidinae (Matthews et al., 2009; M. L. Matthews, Institute of Systematic Botany, University of Zurich, Switzerland, pers. comm., 2010).

In *Ascocentrum* species, hairs are distributed as a wide band around the central region of the nectary spur. It is possible that their position deep within the spur prevents them from being destroyed during nectar probing. The arrangement of secretory hairs also appears to be correlated with that of the vascular bundles, the longer hairs usually occurring closer to the latter. The spurs of nearly all Aeridinae species studied here are supplied by two large and several smaller vascular strands. As a result, the distribution pattern, and thus density, of the hairs is relatively uniform. The only exception was *Papilionanthe vandarum*, where hairs occur mainly as a longitudinal band along the main vascular bundle that runs the length of the adaxial wall of the spur. This is similar to the arrangement found in the moth-pollinated *Angraecum germinyanum* Hook.f. (Davies and Stpiczynska, 2008).

Generally, nectar sugars are transported to the nectary, as pre-nectar, via the phloem. From here, they pass to the

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**Fig. 14.** Histology and ultrastructure of spur of *Stereochilus dalatensis*; (A–F) light micrographs; (G–I) transmission electron micrographs. (A) Section through bilocular nectary spur showing lumina lined with glabrous epidermis and vascular bundles embedded in parenchyma. Scale bar = 300 μm. (B) Detail of vascular bundle with phloem and xylem in equal proportions. Scale bar = 30 μm. (C–E) Epidermal cells with thick, outer walls and distended cuticle (arrows), large nuclei and plastids containing starch grains. Subepidermal cells are thin-walled. Scale bars = 20 μm. (F) Cuticle covering secretory epidermis following treatment with auramine O, but lacking fluorescence. Idioblast with raphides marked with asterisk. Scale bar = 25 μm. (G). Parietal cytoplasm of secretory epidermal cell, showing outer cell wall with distended cuticle. Scale bar = 2 μm. (H) Granular cytoplasm of secretory epidermal cell containing mitochondria, ER and plastids. Scale bar = 1 μm. (I) Detail of cytoplasm with ER and mitochondria. Scale bar = 1 μm. Abbreviations: see Fig. 1.
secretory cells, along either the symplost or the apoplast. In those members of the Aeridinae investigated, both forms of transport can coexist, as the cells are interconnected by numerous plasmodesmata, and the relatively thick cellulose cell walls, when tested histochemically, showed no evidence of barriers to apoplastic transport. Such an arrangement has been recorded for a number of other taxa (Fahn, 2000; Stpiczyńska et al., 2004; Nepi, 2007).

Anatomically, nectaries (nectary spurs) of Asiatic Ascocentrum species closely resemble the morphologically dissimilar nectaries of Neotropical species of Maxillarinae (Ornithochilum coccineum, O. sorophoritis), Laeliinae (Hexisca imbricata) and Oncidiinae [Symphygodorum sanguineum (Rchb.f.) Schltr.], especially in their possession of collenchyma (Stpiczyńska et al., 2004, 2005, 2009; Stpiczyńska and Davies, 2006). This tissue may not only function in protecting the delicate nectary tissue from the beaks of pollinating birds, but probably, simultaneously, provides an apoplastic route for nectar movement within the nectary.

Amyloplasts were absent from the nectary cells of many species, such as A. garayi. This is in contrast to the results obtained for most nectariferous species studied to date, where plastids differentiate to form amyloplasts and become implicated in nectar production. Typically, starch grains are abundant in amyloplasts at the presecretory stage, but as nectary secretory activity progresses, they disappear and plastids develop irregular profiles (Nepi, 2007). Starchless nectary plastids have also been observed in Gymnadenia conopsea (L.) R. Br. (Stpiczyńska and Matusiewicz, 2001), O. coccineum and O. sorophoritis (Stpiczyńska et al., 2004, 2009) and here sugars present in the nectar are probably delivered in the phloem. The phloem component of vascular bundles supplying the nectary spur was particularly well developed for those species investigated here. This agrees with a number of previous studies which showed that a relationship exists between phloem supply and nectar carbohydrate production (Nepi, 2007, and references therein). Although plastids within the secretory epidermal cells of those Aeridinae species studied here rarely contain starch, they frequently contain osmophilic material. This was particularly evident in P. vandarum, where such plastids might be involved in the synthesis of secondary metabolites. The nectary cells of all species investigated, regardless of pollinator, have numerous mitochondria, ER profiles, dictyosomes and small vesicles: cellular characters concomitant with granulocrine secretion (Nepi, 2007, and references therein).

The relatively thick cuticle overlying the secretory layer of all Aeridinae species studied lacks pores and cracks, and is probably permeable to nectar. In all species, it was stained with Sudan III, indicating the presence of lipids. However, the staining reaction with auramine O was more variable. For example, in hairless species, such as Schoenorchis gemmata and Stereochilus dalatensis, the cuticle did not stain at all with auramine O, whereas that of Sedirea japonica only stained slightly when compared with the staining reaction of Sudan III. The cuticle covering the secretory hairs also stained only slightly or, as in P. vandarum, did not stain at all with auramine O, whereas that overlying flat epidermal cells showed a greater uptake of stain. These results probably indicate that variation in the chemical composition of the cuticle results in localized differences in permeability to nectar, as has been recorded for Platanthera bifolia (L.) Rich. and P. chlorantha Custer ex Rchb. (Stpiczyńska, 1997, 2003). In Schoenorchis gemmata and Stereochilus dalatensis, however, nectar secretion is accompanied by distension of the cuticle. Although cuticular distension was not observed for other taxa, it is known to occur in certain ornithophilous Maxillarinae and Laeliinae (Davies and Stpiczyńska, 2008, and references therein; Stpiczyńska et al., 2009). On the basis of the results presented here, it would appear that distension of the cuticle is not confined to either ornithophilous or entomophilous species. Differences in the structure and thickness of the nectary cuticle were also observed for Aeridinae. For example, whereas the epidermal cuticle of Ascocentrum garayi is lamellate, that of Schoenorchis gemmata and Stereochilus dalatensis has a delicate, reticulate layer. Furthermore, fine cuticular striations, although present in nectariferous Sedirea japonica and Papilionanthe vandarum, are absent from Ascocentrum garayi, A. ampullaceum var. aurantiacum, A. curvifolium, Schoenorchis gemmata and Stereochilus dalatensis. Bell et al. (2009), however, observed cuticular striations on epidermal cells lining the spurs of nectarless Orchidaceae. Therefore, whether a given species is nectariferous or rewardless is not necessarily related to the presence, or otherwise, of cuticular striations.

Despite the small number of Aeridinae species investigated here, it would appear that the nectary spur of this subtribe varies considerably in its structure. Some show modifications characteristic of ornithophilous species, whereas others display characteristics of insect-pollinated taxa. The occurrence of identical nectary trichomes and similar spur vasculature in both ornithophilous and entomophilous species of Aeridinae indicates that these structures evolved independently of pollination syndrome. Comparison of the data presented here with previous results (Stpiczyńska and Davies, 2006; Davies and Stpiczyńska, 2008; Stpiczyńska et al., 2004, 2005, 2009) shows that certain, ornithophilous, Asiatic Aeridinae have a number of features in common with Neotropical, hummingbird-pollinated species assigned to subtribes Maxillarinae, Laeliinae and Oncidiinae, most notably the presence of a protective, subepidermal collenchyma. The occurrence of similar anatomical organization in orchid taxa found on other continents and assigned to other subtribes is indicative of convergence and thus appears to be related more to pollinator-driven selection than to phylogeny.

Given the enormity of Aeridinae and the relatively few species presented here, it is important that ultrastructural work now be extended to include other taxa selected according to their phylogenetic position, so as to improve upon our current knowledge and understanding of nectary diversity in this subtribe.

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LITERATURE CITED


