

## STUDIES IN *OBERONIA*, 9: LESSONS FROM EXCESS NAMES IN *OBERONIA* FOR ORCHIDACEAE SYSTEMATICS, INCLUDING A REVISION OF THE *OBERONIA* SECT. *SCYTOXIPHIMUM*

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**ABSTRACT.** The reasons for excess names in microfloral orchids such as *Oberonia* Lindl. can be traced to poor scholarship (e.g., failure to review the literature, ignoring expert advice), typological thinking, and erroneous assumption of microendemism. Some extraordinarily poor descriptions, including some from the 21<sup>st</sup> century, can be termed “taxonomic vandalism”. The outdated reliance on drawings as opposed to z-stacked photographs and scanning electron micrographs poses further problems due to an abundance of demonstrable problems with drawings. The *Oberonia* sect. *Scytoxiphium* Schltr. with eight described species is reduced to one species, *Oberonia heliophila* Rehb.f.; it is illustrated by original drawings, live photographs and scanning electron microscope images. The distribution is extended from Java through Micronesia and Samoa. The species occurs predominantly from 0–500 m, less frequently to 900 m, and possibly to even 1900 m. It flowers throughout the year.

**KEYWORDS/PALABRAS CLAVE:** *Oberonia*, *Oberonia* sect. *Scytoxiphium*, revision, revisión, synonymies, sinonimias, taxonomic vandalism, vandalismo taxonómico

**Introduction.** *Oberonia* Lindl. is a genus of malaxid orchids with some 470 published species, of which some 200–300 have been considered to be correct by various authors over the past 190+ years. That number has dropped recently by identifying at least 60 new synonyms (Geiger 2016, 2019, 2020a, Geiger *et al.* in press, Bunpha *et al.* 2019), but more excess names need to be formally removed (Geiger unpubl. data). It appears that the presumed diversity of *Oberonia* has been overestimated by approximately one third. The degree of overnaming in the genus is astounding. Previous work on minute organisms such as the marine microsnail family Scissurellidae *s.l.* (Geiger 2012 and references therein) with a similar number of species-level names had fewer unrecognized synonyms, but over 60 genuinely new species (and even genera). The question arises, why do microfloral orchids still contain so many unrecognized synonyms?

Here I try to provide some explanations, which may serve other orchid systematists as an incentive to critically assess already described orchid diversity, to encourage to formally synonymize excess names, and not to contribute to the problem by describing even more taxa that ultimately prove to be synonyms. The

last aspect, akin to the medical maxim of “do no harm” should apply to orchid systematics. The importance of alpha taxonomic assessment was recently stressed by Karremans *et al.* (2020) in their landmark contribution on *Vanilla* Plum. ex Mill.

*Taxonomic vandalism.*— It is remarkable that most of the names of *Oberonia* introduced in the later 20<sup>th</sup> and 21<sup>st</sup> century turn out to be synonyms of previously described species. In some cases, the descriptions are so poor in terms of lack of elementary scholarship that they qualify as “taxonomic vandalism” (see also Moore *et al.* 2014, Páll-Gergeley *et al.* 2020, Geiger 2020a). This term may sound like hyperbole, but examining the definition of vandalism as “willful or ignorant destruction of artistic or literary treasures” (Random House Dictionary of the English Language 1973) or “any activity that is considered to be damaging or destroying something that was good” (Cambridge Dictionary) shows that the term is appropriate in some cases.

In academic publishing, review of the existing literature is a key element of any contribution. If an author did not cite a single reference or missed a well-known

and key reference, and thereby introduced superfluous names instead of using a good, correct name, it fits the definition of vandalism. The first situation is found in Chen's (2003) description of *Hippeophyllum micrathum* S.C.Chen, a synonym of *Oberonia rhizomatosa* J.J.Sm. (Geiger 2020a), with not a single reference cited. The second case is exemplified by George *et al.* (2019) who did not cite Ansari and Balakrishnan (1990) as the key review of Indian *Oberonia* in their description of *O. saintberchmansii* Kad.V.George & J.Mathew, which would have immediately identified their specimen as *O. brunoniana* Wight (Geiger 2020a).

A second category is willful ignorance of expert advice. The present author communicated the identity of a specimen as *O. griffithiana* Lindl. That specimen was described anyway as *O. khuongii* Aver. & V.C.Nguyen (Averyanov *et al.* 2019). The new species was compared to a very dissimilar species (*O. cavaleriei* Finet), but *O. griffithiana* was only mentioned in passing and no differentiating characters were given. The protologue of *O. griffithiana* is easily available on-line from the Biodiversity Heritage Library, and the illustrations are exquisite. The two species were synonymized formally by Geiger (2020a).

*Explanatory taxonomy.*— If we subscribe to the premise that systematics as a branch of the natural sciences is explanatory (e.g., Popper 1983, Josephson & Josephson 1993), and the simplest explanation is the best (Ockham's Razor, Mach's Principle of Economy, parsimony), it follows that the goal of systematics should be to explain biodiversity by postulating the fewest species. The term parsimony is used here in the broad philosophical sense, as opposed to the more restricted meaning in phylogenetic systematics generally associated with Hennigian cladistics (Wiley *et al.* 1991). In cladistics, the phylogeny requiring the fewest character state changes, the most parsimonious tree, is the best evolutionary explanation for the characters observed. This is a special application of a general principle that the simplest explanation should be preferred, or the explanation requiring the fewest ad hoc assumptions. This guiding principle is so widely applied that it is known under several names including Ockham's razor and Mach's Principle of Economy.

van Steenis phrased the above: "It is not our task to find out how many species there are, but how few."

(P. Hovenkamp pers. comm.). This approach is also termed "lumping" as opposed to "splitting" in systematics. In a perfect world and as a normative goal, we strive to find out how many true species exist in the world. In the real world, though, there are borderline cases. Both, from the philosophical-economical explanation perspective, as well as following van Steenis' circumscription, it follows not to describe an additional species unless there is good evidence for it. From an explanatory and information criterion perspective, the postulation of a new species should be a measure of last resort. It should be viewed as an admission of failure to explain the observed specimen as an instance of already described species, applying the species-as-kind concept (Mahner & Bunge 1997).

Such a restrained approach to describing species may surprise some readers, particularly in the light of habitat loss and the sixth mass extinction of the Anthropocene. Taxonomy is a pure science, not a branch of practical conservation biology. As taxonomists we evaluate specimens and place them in order to the best of our abilities. If an endangered species turns out to be the same as a widespread one, a responsible taxonomist will synonymize the two. The reduced number of endangered species is of no concern to a taxonomist. Similarly, math does not change its rules. One plus one still is two, even if three would look better to some as it is a larger number.

There is always the possibility that novel characters may justify additional taxa. That possibility should not be abused by wantonly introducing new species and leaving it to subsequent workers to find the characters. Every new species must be justified in its protologue.

*Specimen/gathering vs. species.*— An overlooked source of unrecognized synonyms is the confusion between specimen/gathering and species. This point may seem to be ill founded as every biologist is well aware of the distinction. However, problems arise from implicit assumptions.

To set the stage, some elementary clarification is required. A single specimen and even multi-specimen gatherings are generally presumed to contain one genotype of one species. Accordingly, a specimen/gathering is one instance of a species. Species are composed of multiple specimens/genotypes, exhibiting some natural variability, which serves the raw material on which

selection acts; species may also show some variability due to phenotypic plasticity stemming from genotype  $\times$  environment interaction. The existence of variability is undoubted in biology, but it is hardly ever explicitly taken into account in species descriptions or comparisons. When natural variability is not considered, and the type specimen is taken as the immutable template of a species' morphology, this approach is referred to as typological thinking (Mayr 1994). Such practices are in stark contrast to the principles of systematics, where names are not given to specimens but to species, which are composed of multiple individuals making up at least one population. It is well understood that types are specimens by necessity, but they are only name bearers (semaphoronts) for the species, having some natural variability, that they define. Ideally, the name-bearing type (holotype, lectotype) exhibits near the average or typical form of the species, but there are many counter examples, while syntypes as well as isotypes and paratypes may document some variability of the species.

When comparing two species, the variability of both need to be considered. The assessment of variability is the domain of statistical analysis, both in terms of descriptive statistics (mean, mode, standard deviation, standard error, skewness, kurtosis) as well as in comparative statistics (t-test, chi square test, ANOVA, DFA). The principles of statistics can easily be applied to systematics.

The interrelation of sample size and smallest detectable significant difference is important to bear in mind. All other things being equal, with large samples, small significant differences can be detected. With small samples, the difference needs to be large to be demonstrably meaningful. Translated to systematics, for a species to be distinct, either a large sample size with consistent small differences is required, or the difference needs to be stark if only few specimens are available. Minute differences between few plants are more likely to represent intraspecific variability than species-level differences.

Assessments of single observations are a special case, because variance cannot be calculated from samples with  $n = 1$ . This case is treated in statistics as a t-test of a single value with a mean (Sokal & Rohlf 1981). The variance of the single specimen is presumed to be the same as the one from a sample with multiple observations. Translated to botanical systematics, the

variability of specimens from potentially new species, typically known from only limited material, needs to be presumed to be equal to that of a well-studied species with plenty of available material. When assessing species that both have only very limited material available (e.g., both only known from type), then the variance of both is presumed to be that of a reasonably close species. Species in the same genus, or section if well-defined, provide that information.

The practical application comes particularly when reading older diagnoses and comparisons, for instance by Schlechter, J. J. Smith, Ridley, or Gagnepain. Their descriptions were typically based on a single gathering,  $n = 1$ . If a comparison was provided, it was typically based on information from the protologue of other species also based on single gatherings,  $n = 1$ . However, a comparison of two *species* implicitly assumes and suggests that those differentiations are based on multiple specimens. That assumption is frequently mistaken, as the observational basis for all taxa is  $n = 1$ . Accordingly, a comparison of *specimens* is carried out, masquerading as comparison of *species*. Intraspecific variability could not be taken into account and adducing known variance from other species was never done. Such descriptions should all a priori be considered to be synonyms, unless the difference is very pronounced. Schlechter (1911), however, in his treatment of *Oberonia* species frequently noted that the differentiation is difficult, which is an immediate red flag.

In my experience as a practicing systematist over 30 years, in instances where few specimens are at hand, it is easy to focus on small differences and to separate them into putative new species. As more and more material is examined, small differences evaporate and are correctly recognized as intraspecific variability or ontogenetic stages. I am not opposed to describing species, and even genera, based on small number of specimens, but they need to be strikingly different. Examples include *Depressizona exorum* Geiger (2003) and *Severnsia strombiformis* Geiger (2016) both introduced as a new genus and a new species based on four and two specimens, respectively (Geiger 2003, 2016). They were both radically different from any other known genus or species.

*Appropriate comparisons.*— When describing a new species, comparison to the most similar taxa is criti-

cal. A lack of any comparison is a first indication of a problem (e.g., Chen 2003). It is more difficult to notice meaningless comparisons in little-known groups. Authors frequently compare their supposed new species to ones that bear very little resemblance to them, instead of those that are much more similar. That comparison then suggests erroneously a large difference despite limited material, although, if the appropriate comparison would have been made, the differences would be much smaller, or even non-existent.

In order to be able to identify the most similar species, it is necessary to understand the species concept of every even remotely similar species, at a global level. This typically requires an understanding of every name ever applied in the genus. This is no small undertaking. After at least seven years of immersing myself in one genus, I still do not fully understand many species names in *Oberonia*. Authors describing new species in many groups, particularly those that have not been globally revised, are highly likely to introduce synonyms.

The above-mentioned case of *O. khuongii* is a case in point. The specimen has a common vegetative habit with flattened leaves on a moderately caulescent shoot. The only comparison was made to the acaulescent *O. cavaleriei* Finet with terete leaves, a character only found in two other species. There are multiple more appropriate comparisons, such as to *O. rufilabris* Lindl., *O. insectifera* Hook.f., *O. jenkinsiana* Griff. ex Lindl., and lastly also *O. griffithiana* Lindl., which is the correct name for *O. khuongii*. Those species all have flattened leaves.

*Oberonia manipurensis* Chowlu *et al.* (2015) with moderately fleshy long falcate leaves and subquadrate lip without distinct lateral lobes was compared to *O. pachyphylla* King and Pantl. with short, stubby and very fleshy leaves, and *O. multidentata* Aver. (= *O. jenkinsiana* Griff. ex Lindl.?) with distinct serrated lateral lobes, but not to *O. mucronata* (D. Don) Ormerod & Seidenf. or any of its synonyms, which it represents (Geiger 2019).

*Oberonia saintberchmansii* mentioned above having flowers with distinct lateral lobes and reflexed remaining tepals was only compared to *O. falconeri* Hook.f. with a triangular lip without lateral lobes and spreading remaining tepals (section IV of Ansari & Balakrishnan 1990), but not to any species in Ansari

and Balakrishnan's (1990) section III with distinct lateral lobes and reflexed tepals (*O. bruononiana* Wight, *O. chandrasekharanii* V.J.Nair, V.S.Rachman & R.Ansari, *O. josephii* C.J.Saldanha, *O. nayarii* R.Ansari & N.P.Balacr., *O. balakrishnanii* R.Ansari, *O. platycaulon* Wight, *O. sebastiana* B.V.Schetty & Vivek., *O. seidenfadeniana* J.Joseph & Vajr., *O. wallichii* Hook.f. = *O. brunoniana*, and *O. wynadensis* Sivad. & R.T.Balacr.). This list of inappropriate comparisons could be extended significantly.

The number of instances of genuine disagreement on intraspecific variability vs. biodiversity is limited. One recent example is *O. janae* Aver. vs. *O. pachyphylla* King & Pantl. Averyanov *in* Averyanov *et al.* (2015), which erroneously cited shared characters as differences (e.g., serration of bracts), and did not adduce known variability in other species to estimate variability. The case is discussed in more detail elsewhere (Geiger 2020a).

The geographic dispersal ability of species needs to be considered, and the list of potential species that need comparison adjusted accordingly. In the case of *Oberonia*, microendemism is highly unlikely given that they have the smallest seeds in the family Orchidaceae, and by extension, angiosperms. Average seed size is on the order of  $150 \times 30 \mu\text{m}$  (Barthlott *et al.* 2014, Geiger 2014, 2020a, Geiger unpubl. data, Geiger *et al.* 2020), hence, wind-dispersal will likely be extensive in these epiphytic species.

The term endemism is not well-defined in biology, as it is relative the geographic area it is compared to. All species are endemic to the planet earth. In general, an "endemic" species has a narrower distribution than one would expect based on comparison to closely related species. Microendemic species are even more restricted than one would expect. There are no hard rules on the cut-offs. As an arbitrary number, one could use less than 10% and 1% of range of a widespread species as a first approximation for those two terms. Frequently they are used with respect to a country such as Vietnam, i.e., a political entity, which has no biological relevance.

Some orchid species seem to have narrow distributions despite their small seeds. Such occurrences require additional explanation, such as host association, special niche requirements, or a combination of biotic and abiotic factors. The null hypothesis, though, has to



include extensive dispersal abilities. Distribution is not a character of the species, and taxonomy only uses observations from the specimens. Any specimen should be able to be identified without knowing where it came from; two specimens that are indistinguishable except for provenance should be considered taxonomically identical, belonging to the same species.

A key problem in botany is the focus on geographically limited floras, rather than phylogenetic units. Species not known from a particular country are rarely considered in comparisons of new species descriptions. Imposing political boundaries on plants following phytogeographic patterns is untenable. With molecular techniques being applied, we can now confidently demonstrate extraordinarily wide distribution patterns in *Oberonia*, for instance from Malaya to French Polynesia (Geiger *et al.* 2020).

*The drawback of drawings.*— In botany, the standard for illustrations is usually still the line drawing. One perceived advantage is the easy juxtaposition of different parts of the plant at different scales. An overview of the entire plant is artfully integrated with an enlargement of the flower and other details. Such artistic composites can also be generated from images using current digital imaging technologies. In zoology, specimens <5 mm have customarily been illustrated with z-stacked photographs or SEM images for at least the past 20 years as will be apparent from a quick perusal of publications in *Zootaxa*. It is not clear why botany has lagged behind in adopting 20<sup>th</sup> and 21<sup>st</sup> century imaging technologies.

On the other hand, and this is a rather specific issue with small objects, the detail that can be observed and rendered with a stereomicroscope is limited. For one, the extremely limited depth of field makes it difficult to recognize the overall structure in three dimensions. Some stereomicroscopes are equipped with a diaphragm to increase depth of field but closing the aperture will also reduce resolution due to diffraction. For scanning electron microscopy, depth of field, or more accurately depth of focus, is usually sufficient and stacking is not necessary. It can be increased with longer working distance with a slight loss in resolution. At the relatively low magnifications used, that reduced resolution is inconsequential. In variable pressure, the longer mean free path leads to signal attenua-

tion, which can be compensated for with higher probe currents (100 vs. 500 pA). That large spot size is also inconsequential from the perspective of resolution at low magnification, but improves signal-to-noise ratio and, hence, image quality; see also Stokes (2008). Different detectors are more or less sensitive to changes in working distance in variable pressure. For instance, the Zeiss VPSE detector is more susceptible than the Zeiss C2DX detector (Geiger pers. obs.).

Light has intrinsic limitations with respect to potential resolution, which in practice is further limited by the optical system used. The theoretical resolution limit for light is half its wavelength ( $\lambda/2$  for  $\lambda = 400\text{--}700\text{ nm} \sim 1/4\text{ }\mu\text{m}$ , Ray 2002). The limited numerical aperture (NA) particularly of stereomicroscopes, typically less than 0.1, reduces the resolution limit to  $>7\text{ }\mu\text{m}$  ( $1.22 \times \lambda/\text{NA}$ : range 4.9–8.5  $\mu\text{m}$ ). The SEM is limited by the probe size, typically 2–5 nm (= 0.002–0.005  $\mu\text{m}$ ), i.e., approximately  $10^3\times$  better in linear dimension,  $10^6\times$  in 2D area. At low magnifications, the number of pixels of the digital capture is the limiting factor. For an image 3000 pixels wide with a field of view of 2 mm (an entire *Oberonia* flower), each individual pixel represents approximately 0.7  $\mu\text{m}$ . Even at low magnifications, the SEM has a better resolution by  $\sim 10^1\times$  in linear dimension, by  $\sim 10^2\times$  in 2D area. This level of detail is available to the investigator during the study, as the image can be displayed at full resolution on screen. Such an image shown in print approximately 3" wide with a 150 lpi line screen can reveal details at a scale of 14  $\mu\text{m}$ .

A further factor is the interrelationship of contrast and resolution, well-known from the modulation transfer function in light optics (Ray 2002). Low contrast features require lower spatial frequencies (= larger structures) than high contrast features to be discernable. Fine flower detail is typically low in contrast. It will increase the size of the smallest recognizable feature in light optical examination of flowers by an estimated factor of  $5\times \sim 35\text{ }\mu\text{m}$ . The SEM, on the other hand, accentuates edges due to effects of the electron beam and its interaction volume in the sample (Goldstein *et al.* 1992), hence, can display the full detail at the level of individual pixels. The SEM applies an unsharp mask filter to the image.

It is an open question of how much actual detail information is usually captured by a skilled scientific

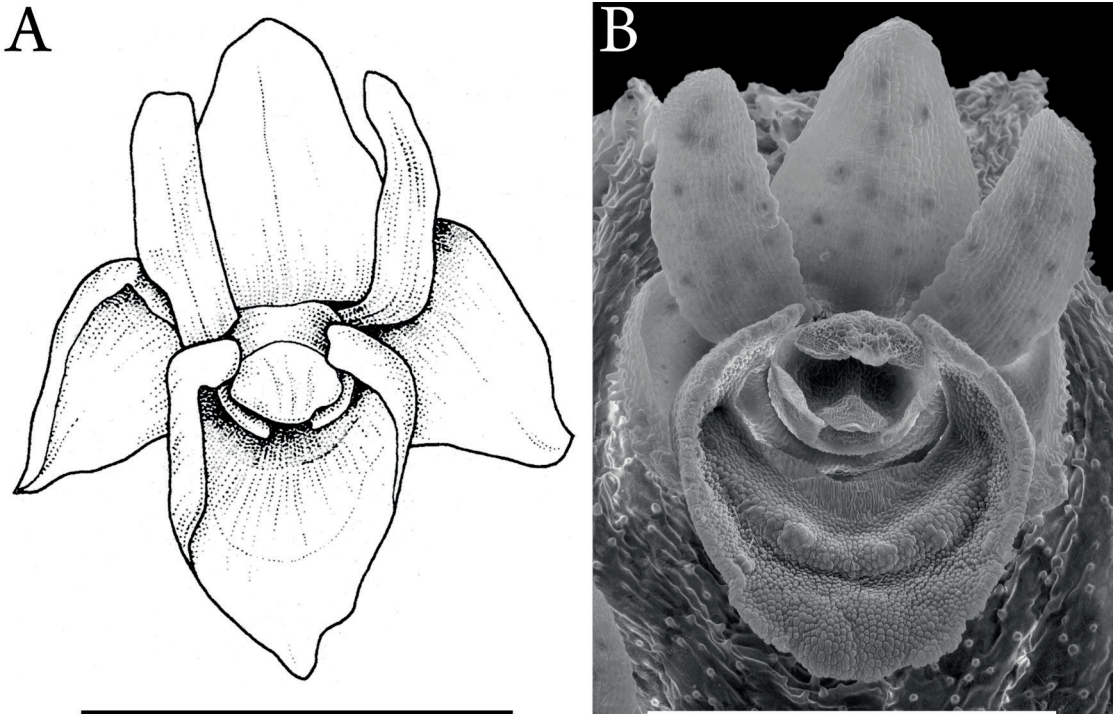


FIGURE 1. Comparison of botanical line drawing with SEM, *Oberonia pachyrachis*. **A.** Drawing from Seidenfaden (1978). **B.** SEM image. *Menzies & Dupuy 192 K 47170* from Thailand. Scale bars = 1 mm.

illustrator, but it is very doubtful that the theoretical limit of the observation optics is transferred to the drawing. The above sample calculations were carried out at the low magnification end, and thus were least favorable to the SEM. At higher magnifications, the SEM's advantages become much more pronounced.

A comparison of a standard line drawing and an SEM image serves here as a case in point (Fig. 1). I have chosen the best line drawing available for *O. pachyrachis* Rehb.f. ex Hook.f. from a key work on *Oberonia* by the highly respected botanist Gunnar Seidenfaden. Seidenfaden's (1978: Fig. 3C; Fig. 1A here) illustration is compared to an SEM image (Fig. 1B). The line drawing shows the shape and proportions of the tepals well, but many details are not recognizable. The concentric dotted rings on the mesochile and epichile seem to allude to the distinct folded ridges revealed by the SEM. However, without the SEM image, it is impossible to understand the 3D folds based on the drawing. The short hairs on the back of the lateral sepals, clearly visible in the SEM, are not indicated in the line drawing. Most drawings in the literature of *Oberonia* do not attain the quality of Seidenfaden's illustration. They may

be better referred to as sketches rather than drawings.

One argument in favor of line drawings is their inherent interpretative nature, and that specimen defects can be ameliorated to provide an idealized representation of the organism. However, this augmentation also has the potential for introducing unintended errors or suppressing details that may turn out to be significant in hindsight. Schlechter's (1923, 1934) drawings often show differences between the illustration of the entire flower and the individual floral elements. They may rather suggest intraspecific variability or show the appearance of the floral elements from different perspectives or may be drawing errors. For instance, the isolated lip of *O. crassilabris* Schltr. (Fig. 2H) shows distinct auricles, which are not shown in the drawing of the entire flower. Geiger (2019) discussed inconsistencies in the drawings of *O. nayarii* Ansari & N.P.Balacr. and *O. balakrishnani* Ansari in Ansari and Balakrishnan (1990). Inconsistencies in the drawings of *O. ensiformis* (Sm.) Lindl. flowers were discussed by Geiger (2020a). A particularly striking case is the drawing in the protologue of *O. caprina* Gilli (1983: fig. 29), which bears little resemblance to the holotype

(W 16722 *Gilli 164*); see Geiger (2019) for details. Unfortunately, a bad drawing is not immediately recognizable as such; it must be taken at face value. On the other hand, deficiencies of photographs or SEM images are immediately apparent. This uncertainty factor surrounding drawings is a significant drawback.

*Treatment in literature.*— The treatment of species in the literature can be a further indication regarding the validity of the species. While there are genuinely rare species, if highly similar species are only treated by reference to the protologue with no new material, it raises questions. In some cases, voucher material can be helpful in untangling questionable species names.

For most early 20<sup>th</sup> century synonyms, it is notable that they have hardly ever been cited after their introduction (e.g., *O. vulcanica* Schltr., *O. nitida* Seidenf.). Drawings are frequently copied and re-copied, while the discussions and comparisons are scant at best. It is assumed that every described species is correct. It may be better to treat every species epithet in a genus not clearly distinct from every species described before as a *nomen dubium* or *nomen inquirendum* (e.g., *O. zimmermanniana* J.J.Sm., *O. weneri* Schltr.).

The presumption of names to be correct may lead authors to focus on minute differences. Bunpha *et al.* (2019) distinguished *O. denticulata* Wight from *O. gammiei* King & Pantl. by the shape of the rostellum and the position of a slight depression in the lip slightly above or below the middle of the lip. Neither illustrations to support their claims nor any indication on the number of specimens examined was provided. In the same article, however, *O. nitida* Seidenf. was correctly synonymized under *O. denticulata*, and *O. falcata* King & Pantl. was correctly synonymized under *O. anthropophora* Lindl. The latter pair shows extensive variability in the shape of the lip, for which reason it is unclear why Bunpha *et al.* (2019) elected to consider minute differences as significant in one species pair, but meaningless in others. Last but not least, these authors missed the senior synonym of *O. denticulata*, namely *O. mucronata* (D. Don) Ormerod & Seidenf. as the correct name for both *O. denticulata* and *O. gammiei*.

The untenable distinction of *O. fungum-olens* Burkill from *O. padangensis* Schltr. by Bunpha *et al.* (2019) correctly synonymized by Geiger (2019) was discussed elsewhere (Geiger *et al.* 2020).

**Materials and methods.** Taxonomic assessments were made based on available information from primary and secondary literature, herbarium holdings (B, BM, F, K, MEL, MICH, MO, P, SING, US, W), and on-line databases (HBG, L).

Illustrations were processed in AffinityPhoto. Z-stack photography was carried out on a Zeiss Discovery V20 stereomicroscope with planapochromatic lenses and motorized focus. Images stacks were captured with a Zeiss AxioCam HRC camera and processed in ZereneStacker. For scanning electron microscopy (SEM), flowers were preserved in 95% ethanol, brought to 100% ethanol through three changes of 100% ethanol, critical point dried in a Tousimis 815A using standard settings, then mounted on double sided carbon tabs (Ted Pella), sputter coated with gold, and imaged in a Zeiss EVO 40 XVP using the VPSE detector in variable pressure (30 Pa) at 20 kV and probe currents ranging from 30–500 pA depending upon working distance and magnification.

#### Section *Scytoxiphium* Schltr.

TYPE: *Oberonia crassilabris* Schltr., subsequent designation by van Royen (1979: 711).

*Remarks.*— The section *Scytoxiphium* serves to illustrate some of the points made above. As the first sign of overnaming, Schlechter (1911: 176, translated from German) noted that “The establishment of boundaries of the species is difficult, because they are all closely related.” Schlechter (1911) described in his new section *O. pachyglossa* and *O. crassilabris* from New Guinea and included *O. dolichophylla* Schltr. and *O. rivularis* Schltr. both previously described from New Guinea, *O. heliophila* Rehb.f. from Fiji, and *O. betchei* Schltr. from Samoa, but not *O. asperula* J.J.Sm. described previously from New Guinea (Smith 1908: see below).

Although Schlechter (1911: 176, translated from German) referred to “rich material of *Scytoxiphium*”, the names were all introduced based on single gatherings ( $n = 1$ ) except for *O. dolichophylla* based on two gatherings. It might well be that he used the term rich in the sense of number of species, as opposed to a large number of specimens. As the Berlin herbarium with all Schlechter’s material was destroyed in World War II, it is impossible to verify either interpretation.

Schlechter (1911) compared his new species only to his own species from New Guinea. Although he



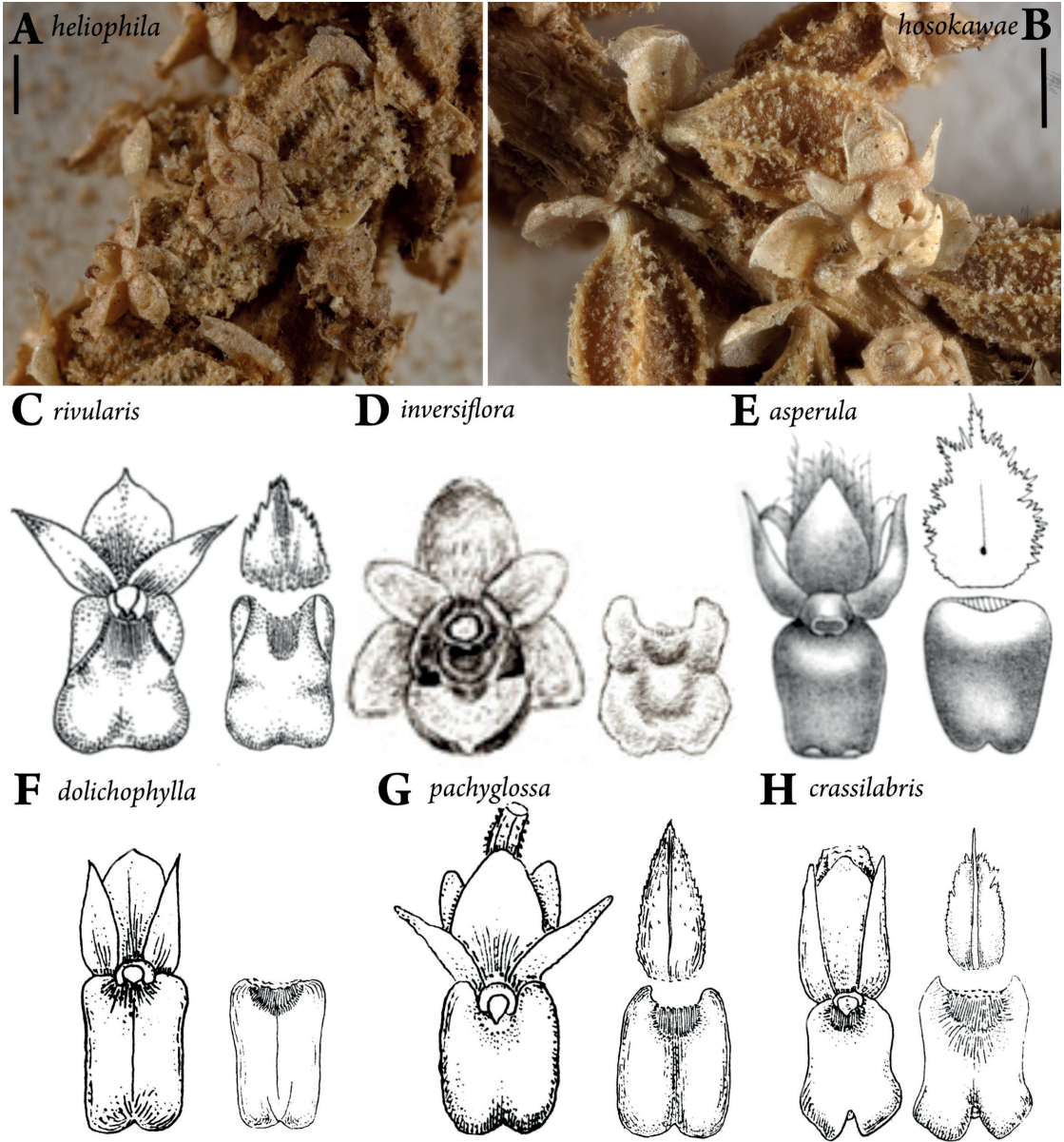


FIGURE 2. *Oberonia heliophila* and its synonyms. **A.** Flowers from lectotype of *O. heliophila* W37726. **B.** Flowers on MICH syntype of *O. hosokawae*. **C–H.** Drawings of flower, isolated lip, and bract. **C.** Original figures of *O. rivularis* from Schlechter (1911). **D.** Drawing of flower of *O. inversiflora* by J.J. Smith (from Schuiteman & deVogel 2006). **E.** Illustration of flower of *O. asperula* from Smith (1909). **F.** Illustration of *O. dolichophylla* from Schlechter (1923); the bract was not figured. **G.** Illustration of *O. pachyglossa* from Schlechter (1923). **H.** Illustration of *O. crassilabris* from Schlechter (1923). Scale bars A, B = 1 mm.

included *O. heliophila* from Fiji and *O. betchei* from Samoa in his section *Scytoxiphium*, those species were not mentioned again. Schlechter (1911: 177, translated

from German) noted under *O. pachyglossa* that "... in *Scytoxiphium* the individual species are always closely related amongst each other and not only in the same

areas, but also in widely separated regions, whose floras tend to have little in common.” Despite morphological similarities, he discounted those similarities a priori because species distributed over larger areas was considered highly improbable. In Schlechter’s opinion, there was no need of comparing the New Guinea species to those from areas further away.

Smith (1908, 1912) described *O. asperula* and *O. inversiflora* J.J.Sm., the latter explicitly in the section *Scytoxiphium*, introduced one year earlier. It demonstrates that Smith was aware of Schlechter’s (1911) work on the orchids of New Guinea. However, neither of Smith’s species were compared to any other. Smith is known to have been an extreme splitter, even in cases where he wondered whether his species had already been described, such as *O. salakana* J.J.Sm. [= *O. merapiensis* Schltr. as discussed in Geiger (2019)]. The lack of comparisons, the  $n = 1$  problem, and Smith’s demonstrated tendency to excess splitting make his taxa highly suspicious.

*Oberonia hosokawae* Fukuy. (in Hosokawa 1941) was also described in section *Scytoxiphium*. While the membership in the section appears to have been discussed in the remarks in Japanese, no other species assigned to the section seems to have been discussed. The lack of comparison and the  $n = 1$  problem make this species equally suspicious. The specific epithet was erroneously corrected to *hosokawai* by Geiger (2020a), an error noted by Geiger (2020b).

The synonymy of several of the species under *O. heliophila* has been discussed elsewhere (Geiger 2020a): *O. asperula*, *O. rivularis*, *O. inversiflora*, *O. hosokawae*. *Oberonia pachyglossa* Schltr. has been treated as a synonym of *O. heliophila* by WCSP (2020); I have not been able to trace a literature reference for the establishment of this synonymy. *Oberonia crassilabris* and *O. dolichophylla* are still recognized species. These three species are formally synonymized here.

#### TAXONOMIC TREATMENT

The type concepts used are in strict accordance with the ICN (McNeill 2014, 2015). Isotypes are duplicate specimens of the same gathering as the holotype explicitly specified with a repository in the protologue. If no holotype was specified, then all specimens are referred to as syntypes, even if from a single gathering.

*Oberonia heliophila* Rchb.f. (1878: 56). (Fig. 2–7).

TYPE: *U.S. Exploring Expedition s.n.* (lectotype W37726; designated by Kores 1989), Mountains of Mathuata Province, Vanua Levu, Fiji. The rather convoluted assessment of other type material has been discussed elsewhere (Kores 1989, Geiger 2020a). Thanks to Paul Omerod (pers. comm.) the *Gräffe s.n.* [1257] syntypes from Upolu, Samoa, were recently found at HBG 501809 and HBG 500445; an additional duplicate is at MO 4338405. While the collector was spelled “Graeffe” in Reichenbach’s (1878: 56) Latin protologue, the label spells the name with an umlaut.

Syn.: *Oberonia dolichophylla* Schltr. in Schumann and Lauterbach (1905: 114). Schlechter 1923: pl. 69, fig. 248.

TYPE: *Schlechter 14579* (syntype: B, lost). At the river board of the Garup, at the foot of the Toricelli mountains [Papua New Guinea], ~100 m, **syn. nov.**

Syn.: *Oberonia pachyglossa* Schltr. 1911: 177. Schlechter 1923: pl. 69, fig. 249.

TYPE: *Schlechter 16756* (syntype: B, lost). Forests at the Kaulo [River, Madang, Papua New Guinea] 400 m, **syn. nov.**

Syn.: *Oberonia crassilabris* Schltr. 1911: 177; Schlechter 1923: pl. 69, fig. 250.

TYPE: *Schlechter 17948* (syntype: B, lost). Forest of the Finisterre mountains, Papua New Guinea, 700 m, **syn. nov.**

MATERIAL EXAMINED: **Country unknown.** *Wight 441* (MEL s.n.), Arbor dolan. **Fiji.** *Degener 15465* (F 1473815, K s.n., MO 1256922, MICH s.n., P 00310689), Ra, vicinity of Rewasa, near Vaileka. *Degener & Ordonez 14166* (K s.n.), Thakaundrove, Marvu, near salt lake. *Gillespie 3877* (K s.n.), Valley of Kalindina near Nambai, Namosi Province. *Greenwood 603* (K s.n.), Mount Mamata coast. *Greenwood 709* (K s.n.), Haulikno, N side. *Greenwood 1113* (K s.n.), Naitasiri, near Nasinu. *Praham 30* (BM 000088476), Vanua Levu. *Smith 7075* (K s.n., P 00310688), Tailevu, Hills E of Wainimbuka River vicinity of Ndakui-vuna. *Parks 20174* (K s.n.). **Indonesia.** *Comber 1373* (K s.n.), Rannpane, N of Semeru. **New Guinea.** *Carr 10045* (BM s.n., CANB 46410, CANB 46321, SING 0141494, SING 0141462 K s.n.), Koitaki. *Carr 10667* (B s.n., BM 000088448, CANB 46443, F 1497506, K s.n., P 00364412, SING 0141482, SING 0141483), Kokoda. *Conn 544* (MEL 1528545), Gulf Province, N side of Lake Tebera. *Reeve 923* (CANB 8500636), Lagaip District. *O’Byrne O.020* (SING



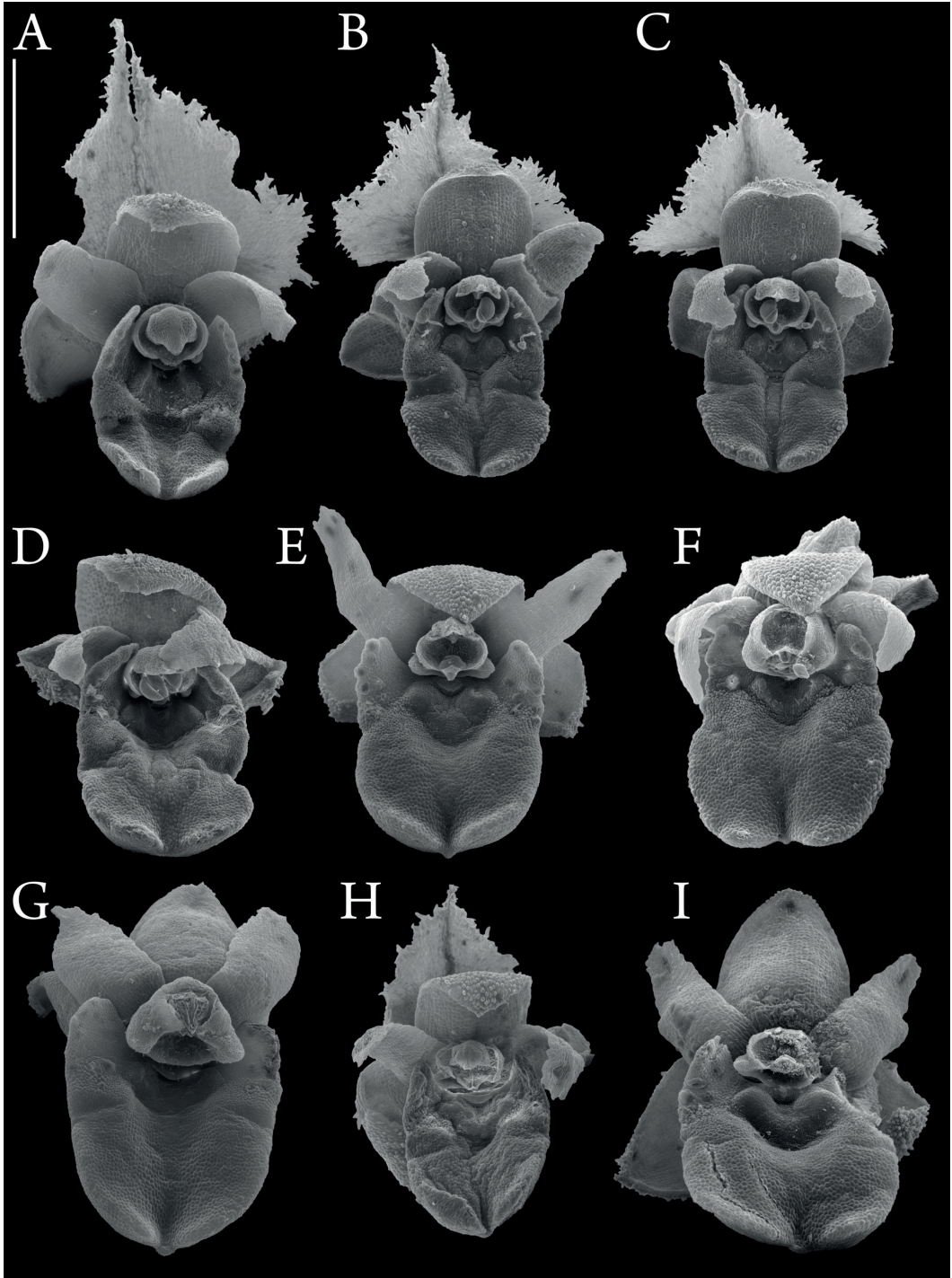


FIGURE 3. *Oberonia heliophila* SEM images of face of flowers. **A.** *Schuiteman 90/555* L 23197 from Papua New Guinea. **B–C.** *Harris 1514* L 19688 from Papua New Guinea. **D.** *deVogel s.n.* L 20059 from New Guinea. **E.** *Mulder s.n.* L22448 from Fiji. **F.** *Hunt 2226* K28510 from the Solomon Islands. **G.** *M.A. Clements 5603* CANB 8916245, from Vanuatu, live image in Figure 6H. **H.** *Mason 1645* K 7169, from Fiji. **I.** *Mulder s.n.* L 24117 Fiji. Scale bar = 1 mm.

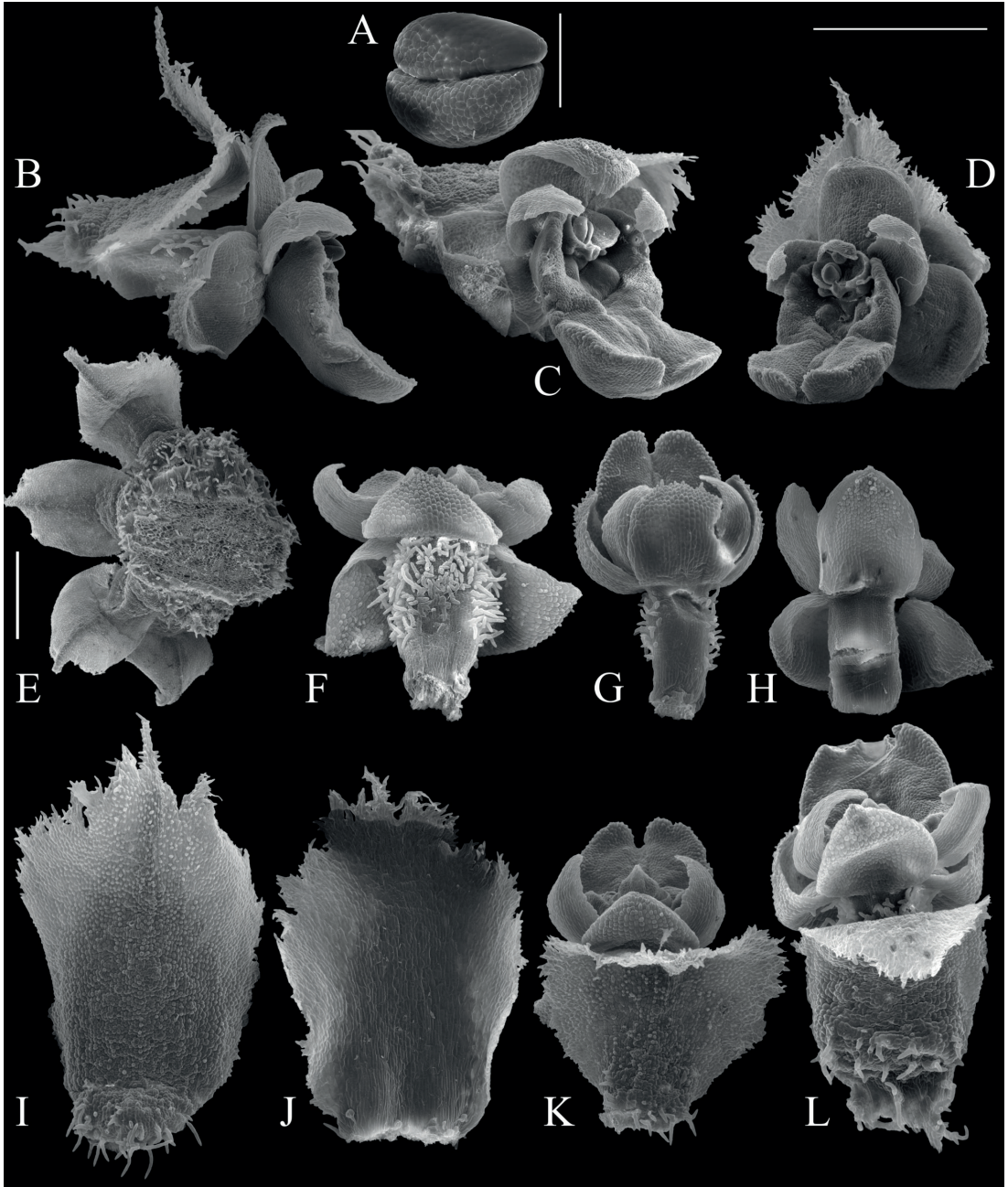


FIGURE 4. *Oberonia heliophila* SEM images. **A.** Pollinaria. Scale bar = 100  $\mu$ m. **B.** Flower lateral with curved up bract and sparse hairs on pedicelled ovary. **C–D.** Oblique view of flower showing thick lip. **E.** Cross section through rachis with several attached bracts. Scale bar = 1 mm. **F–H.** Top of view of flower without bracts showing variable degree of hairs on pedicelled ovary. **I–J.** Isolated floral bract. **I.** External view. **J.** Internal view. **K.** Top view of flower with spreading bract. **L.** Top view of flower with clasping bract. Scale bar except A, E = 1 mm. A, I, J. *deVogel s.n.* L20059 from New Guinea. B, D, G, K. *Harris 1514* L19688 from Papua New Guinea. C. *deVogel s.n.* L20059 from New Guinea. E, F. *Hunt 2226* K28510 from the Solomon Islands. H. *Schuiteman 90/555* L23197 from Papua New Guinea. L. *Leg. ign. s.n.* K 21001, *loc. ign.*



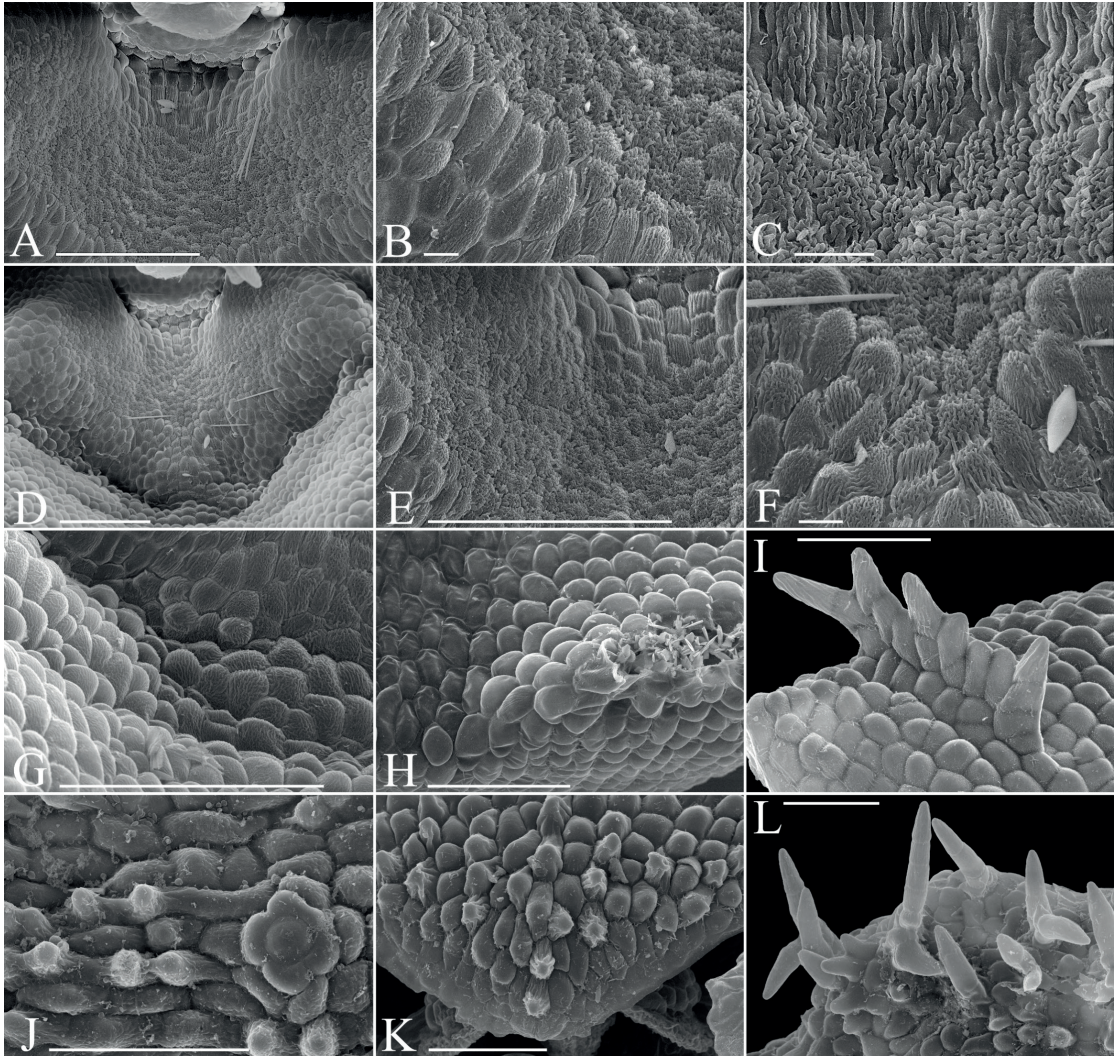


FIGURE 5. Floral details of *Oberonia heliophila*. **A, D.** Disc area with sac. Scale bar = 100  $\mu$ m. **B.** Transition between mesochile with elongated moderately rugulate cells to disc with round strongly rugulate cells. Scale bar = 10  $\mu$ m. **C.** Transition between disc with round strongly rugulate cells to sac with rectangular striate cells. Scale bar = 10  $\mu$ m. **E.** Transition from mesochile (lower left) through disc (center) to sac (upper right). Scale bar = 100  $\mu$ m. **F.** Median tip of disc (round strongly rugulate cells) and mesochile (elongated moderately rugulate cells). Scale bar = 10  $\mu$ m. **G.** Transition between mesochile and epichile with crease and reduction in rugulosity. Scale bar = 100  $\mu$ m. **H.** Edge of thick epichile. Ruptured cells with mineral deposits. Scale bar = 100  $\mu$ m. **I, L.** Trichome ridge on lateral sepals. Scale bar = 100  $\mu$ m. **J.** Top surface of bract with short trichomes and cross cells. Scale bar = 100  $\mu$ m. **K.** Tip of median sepal with rugulate-mammilate cells. Scale bar = 100  $\mu$ m. **L.** Trichomes at base of bract. Scale bar = 100  $\mu$ m. (A–I: *deVogel s.n.* L 20059 from New Guinea. J–L: *Harris 1514* L 19688 from Papua New Guinea).

0141441), Papua New Guinea. *Reeve 5397* (K s.n.), Seargu [Beneni] SW side of Lak Kutubu, Nipa District, Southern Highlands. **Samoa.** *Mansfeld 146* (P 00310612), near Malololelei. *Mansfeld 1882* (K s.n.), Matavanu. *Vaupel 234* (K s.n., MO 1614357), Matautu. *Walter s.n.* (MEL 2394315), Samoa. *Whistler W2790* (K s.n.), W of Aoloaufou. *Whit-*  
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*mee 43* (MEL 2394729), Samoa. *Whitmee 45* (K s.n.), Samoa. *Whitmee 48* MEL 2394730), Samoa. *Whitmee 165* (K s.n.), Samoa. *Whistler W1702* (K s.n.), N of Potlacthe Lumber Mill at Asau. **Solomon Islands.** *Hunt 2226* (K s.n.), 1/4 mile below confluence of Warahito and Pegato rivers. *Wickison 87* (K s.n.), Chariveghu Drainage system.

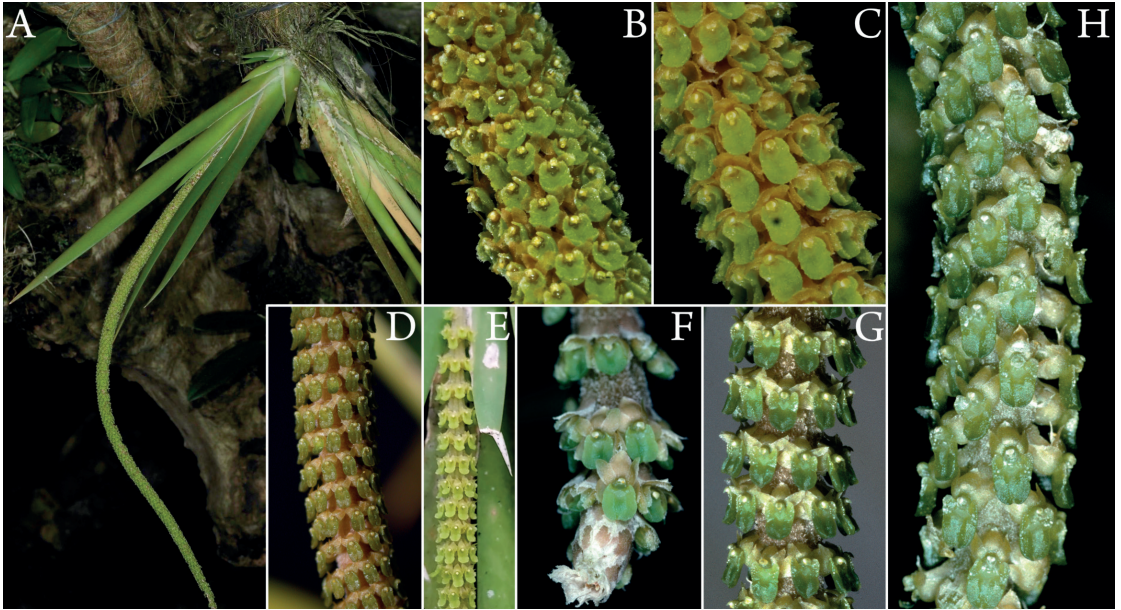


FIGURE 6. Live *Oberonia heliophila*. A–C. Cultivated plant from Papua New Guinea, collection and photographs Jeffrey Champion. A. Habit. B. Flowers of plant shown in A, first flowering with flowers scattered on inflorescence. C. Flowers of plant shown in A, second flowering, with flowers in whorls. D. Flowers of plant in cultivation from Papua New Guinea at Leiden Botanical Garden 20031565, photograph by Eduard de Vogel. E. Almost entirely green flower in plant in cultivation from Papua New Guinea, collection and photograph of Maryse Devaevae. F. Flowers of *Clements KK6916* from Simbai, New Guinea. G. Flowers of *Clements 5392* from Efate, Vanuatu. SEM image in Figure 3G. F–H. Photographs by Mark Clements.

**Vanuatu.** *Clements 5603* (CANB 8916245), 30 km W of Luganville, Santo, Wounauss Village area. *Morrison s.n.* (K s.n.), Gulley near Amlganhat, Aneityum mountain. **Wallis and Futuna.** *Pillon 895* (P 02102968), Wallis, Mout Lulu Fakahega. *MacKee 39257* (P 00310691, P 00310690), Wallis, Mount Lulu.

**NOTES:** *Oberonia dolichophylla* is a synonym of *O. heliophila*. There are no known surviving syntypes of *Schlechter 14579*. *Schlechter 19997* of *O. dolichophylla* at BO (Cribb & Robbins 1990: a duplicate of the now destroyed B specimen) is not a syntype but a voucher for the species' listing in *Schlechter* (1911). The species was described as rather large for an aculescent *Oberonia* (45 cm), with pointed and hairy bracts, green flowers, oblique lanceolate petals, lip oblong with truncated end [= subquadrate], and the epichile with a shallow notch. All these characters agree with *O. heliophila*. *Schlechter* in *Schumann & Lauterbach* (1905) compared his *O. dolichophylla* only known from the type *Schlechter 14579* with *O. rivularis* only known from the type *Schlechter 13801*. It is an example of comparing specimens instead of species

and the  $n = 1$  problem discussed above. He noted the longer inflorescence, bracts and lip, as well as thicker leaves in *O. dolichophylla*. The length of the inflorescence is meaningless in *Oberonia* as has been shown from plants grown in cultivation (cf. Geiger 2018, 2019). The diameter of the inflorescence is given as 1 mm in *O. rivularis* and 2 mm in *O. dolichophylla*, the bracts are approximately 1.5 mm long in *O. dolichophylla* and 1 mm in *O. rivularis*, and the lip is given as 2 mm long in *O. dolichophylla* and slightly  $>1$  mm in *O. rivularis*. Whether those measurements are accurate is unknown and unverifiable as the material at B was destroyed. Inaccuracies in measurements and scale bars have been documented previously (Geiger 2019). Given that the comparisons are based on single specimens ( $n = 1$  each), those small differences are not sufficient to justify species-level differentiation.

*Oberonia pachyglissa* is a further synonym of *O. heliophila*. The name has rarely been mentioned in the literature since its initial publication. O'Byrne's (1992, 1994) voucher (SING 0141441) can be referred to *O. heliophila*, while Schuiteman and de Vogel (2006)



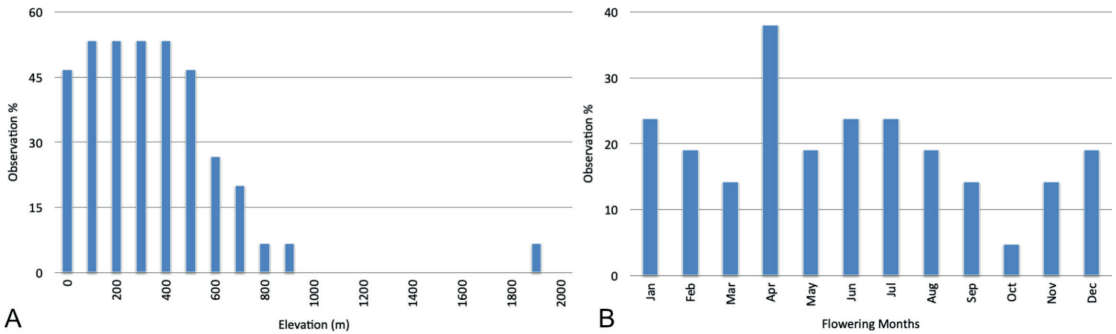


FIGURE 7. Ecological data for *Oberonia heliophila*. **A.** Elevation,  $n = 16$ . **B.** Phenology,  $n = 22$ . (references under any known synonym. Elevation: Christophersen 1935, Cribb & Whistler 1996, Fukuyama in Hosokawa 1941, Hawkes 1952, Kores 1991, Lewis & Cribb 1991, Millar 1999, O'Byrne 1994, Schlechter 1911, Schlechter in Schumann & Lauterbach 1905, Schuiteman & de Vogel 2006, herbarium records. Phenology: Christophersen 1935, O'Byrne 1994, Reichenbach 1878, Schlechter 1910, 1911, Schlechter in Schumann & Lauterbach 1905, Schuiteman & de Vogel 2006, Smith 1909, 1915, herbarium records).

reproduced Schlechter's (1923) figure. The species was described as being of large habit (55 cm), having dentate bracts, green flowers, linear-lingulate petals, with a thick, subquadrate lip with small auricles and somewhat truncated epichile. All these characters agree with *O. heliophila*. Schlechter considered *O. pachyglissa* only known from type Schlechter 16756 and *O. dolichophylla* only known from type Schlechter 14579 as distinct based on the slenderer inflorescence (3 mm in *O. dolichophylla*, no measurements given for *O. pachyglissa*) and the linear petals. The claim about the width of the inflorescence cannot be evaluated due to lack of data. The shape of the petals is continuously variable in *O. heliophila* (Fig. 3); bimodal distribution, pattern, or covariance cannot be identified. The simpler explanation is variability in this character, rather than postulating multiple, almost cryptic species with additional intermediates that cannot be clearly assigned to any of the postulated species.

*Oberonia crassilabris* is a synonym of *O. heliophila*. The taxon has been designated as the type species of the section *Scytoxiphium* by van Royen (1979) and a second time by Schuiteman and de Vogel (2006), who copied Schlechter's figure. The type specimen was described as large (60 cm), with deeply incised bracts, tepals with a green lip with remaining tepals brownish, thick lip, subquadrate with small auricles and bilobed epichile over one quarter of lip length. All these characters agree with *O. heliophila*. Schlechter considered *O. crassilabris* only known from type Schlechter 17948

and *O. dolichophylla* only known from type Schlechter 14579 as distinct based on the oblique lanceolate vs. oval lanceolate petals, and the depth of the incision of the epichile of the lip. As indicated above, the shape of the petals is variable, and the alleged differences are at best slight. The depth of incision of the epichile is equally variable with no pattern discernible (Fig. 3).

A syntype of *O. betchei* Schltr. was found in MEL and confirms the synonymy with *O. heliophila* (Geiger pers. obs.).

*Plant* large, acaulescent, fan-shaped, leaves to 75 × 1.6 cm, acuminate, laterally compressed, fleshy; flowering plants as small as 25 cm. *Inflorescence* terminal, to 64 cm long, flowers typically in clearly separated whorls (Fig. 6C–F), occasionally scattered (Fig. 6B, H). *Bract* as long as flower to somewhat longer, typically curved upwards (Fig. 4A, D, K, L), typically wider than flower, occasionally as wide as flower, triangular acuminate, at base hirsute, at tip strongly dentate, occasionally lacinate (Fig. 3B–C, 4D, I–L), frequently with 1–2 terminal awns (Fig. 3A). Pedicelled *ovary* rather short for genus, strongly pubescent towards flower to glabrous (Fig. 4B, F–H). *Flower* with lip in green tones, remainder of tepals typically in shades of tan (Fig. 4E–H), occasionally yellowish (Fig. 4B–C), occasionally entire flower light green (Fig. 4E). Considerable variation in flower color on same inflorescence (O'Byrne 1994: as *O. pachyglissa*). *Sepals* triangular, sometimes with sparse short hairs abaxially (Fig. 4C,



F). *Petals* narrowly ovoid to lanceolate, about as long as sepals. *Lip* thickened (Fig. 4B–C), auricles extending to middle of gynostemium; indistinct lateral lobes with irregular margins (Fig. 3, 4C–D); disc with shallow sac (specific shape affected by preservation: Figs 3, 4C–D, 5A, D) flanked by thickened pads of variable shape and distinction; indistinct constriction in middle of lip (Figs. 3, 4C–D, 5D); epichile broad with more or less distinct apical notch separating two lobes (Fig. 3). *Gynostemium* short and thick for genus. *Pollinaria* composed of two pollinia of unequal size, kidney shaped, without caudicle or viscida (Fig. 4A).

Cell surface morphologies consisting of striate around disc (Fig. 5C, E), coarsely rugulate around mesochile end, central portion of epichile (Fig. 5C, F), finely rugulate on main portion of mesochile and epichile (Fig. 5A, B, G), glabrous pneumatophore at tips of epichile (Fig. 5H). Short rugulate trichomes on the backside of the sepals (Fig. 5J, K), longer uniserial trichomes at base of bract (Fig. 5L).

The arrangement of flowers in whorls and scattered is variable as demonstrated by repeat flowering of same plant in cultivation, based on photographs taken by J. Champion (Fig. 6A–C). The presence and density of the hairs on the pedicelled ovary varies continuously with no patterns discernible. A few conditions are illustrated here: dense hairs (Fig. 4F), scattered hairs (Fig. 6B), sections variously hairy and glabrous (Fig. 4G), glabrous (Fig. 4H).

**DISTRIBUTION:** Indonesia (Java: *Comber 1373* K s.n.), Fiji, Micronesia, New Guinea, Samoa, Solomon Islands, Vanuatu, Wallis and Futuna (citation under any of the known synonyms: Cribb & Whistler 1996, Hawkes 1952, Kores 1989, 1991, Lewis & Cribb 1989, 1991, Millar 1999, O’Byrne 1994, Parham 1972, Reichenbach 1878, Schlechter 1910, 1911, Schuiteman & de Vogel 2006, Smith 1908, 1912, 1915, Williams 1938, 55 herbarium records: B, BM, CANB, F, K, MEL, MICH, MO, NSW, P, SING, US, W).

**ECOLOGY:** Mostly pendulent, occasionally erect, branch and trunk epiphyte on mangrove trees, bread fruit (*Artocarpus altilis*: Moraceae), *Hevea* (Euphorbiaceae), *Terminalia catappa* (Combretaceae), *Pterocarpus* (Fabaceae), *Pometia* (Sapindaceae), mahogany (*Swietenia humilis*: Meliaceae), *Melaleuca* sp. (Myrtaceae) and palms (*Cocos nucifera*, *Metroxylon* sp.). In coastal

to montane forest, sun exposed. The species flowers throughout the year (Fig. 7B).

*Oberonia heliophila* is found regularly from sea level to about 500 m, with decreasing frequency to 900 m (Fig. 7A). The single record from 1900 m (*Comber 1373* K s.n. from Java) needs verification. The elevation at the named location matches the indicated elevation based on GoogleEarth. That record could be mislocalized. However, a photograph by Mark Clements of the species was taken around Simbai, New Guinea, which is at approximately 1900 m elevation, including its surroundings (Fig. 4E).

**CULTIVATION:** This warm growing species can be cultivated on tree fern slabs, possibly with some coco fiber pad, ideally in bright shade (lath house), but tolerating wide range of light, with high humidity, daily watering, and excellent ventilation (O’Byrne 1994, Schuiteman & de Vogel 2006, J. Champion pers. comm.).

**Discussion.** One reviewer criticized that the findings here are not supported by molecular data, therefore, are only an unsubstantiated opinion. A similar position was advanced by Jones (2021) with respect to the Australian *Oberonia* names synonymized by Geiger (2019). Such a point of view is ill founded for the following reasons.

1) Taxonomic assessments are based on types. Type specimens cannot be destructively sampled. Taxa with lost types have no chance of being evaluated based on genetic data, which applies to all three names synonymized here. Taxonomic assessment is based by necessity on the available information, such as herbarium sheets, illustrations and the protologue.

2) If species can be described based on morphology, then they can also be synonymized based on morphology. An asymmetry of evidentiary burden is unjustifiable. Otherwise, one would have to require all new species descriptions to be supported by molecular data as well. The vast majority of species descriptions even today are based exclusively on morphology, some notable exceptions notwithstanding.

3) The inclusion of molecular data does not guarantee a better outcome. Questions such as selection of marker(s), choice of analytical settings, choice of ingroup and outgroup taxa, mistakes in the laboratory (contamination), and identification of material all har-

bor potential for erroneous results. There is no question that molecular data have produced tremendous advances in systematic biology, and I have used those techniques myself. However, there have also been high-profile blunders.

4) The insistence on using molecular data would invalidate all systematic paleontology.

Additionally, the main point made here of incorporating intraspecific variability into taxonomic assessments is supported by molecular data from better known species, such the molecular phylogeny of *O. equitans* (Forster) Mutel by Geiger *et al.* (2020a) including fine morphological examination of flowers. Extensive morphological variability is also suggested by the molecular phylogeny of Li *et al.* (2016), which showed six samples of *O. austro-yunnannensis* S.C.Chen & Z.H.Tsi and ten samples of *O. jenkinsiana* Griff. ex Lindl. in a completely unresolved polytomy with extremely short terminal branches. This is a strong indication of synonymy.

Last but not least, scientific progress is incremental, and all taxonomic assessments are opinions, whether supported by morphology or molecular data. Additional information can either lead to confirmation of an earlier result, or it can lead to a new hypothesis being advanced. I welcome a re-examination of this proposal.

**Conclusion.** Within *Oberonia* sect. *Scytoxiphium*, taxonomic ranks have typically been employed at one

level higher than is appropriate for species recognition. Specimens were named as species, while the single species was treated as a section. The underlying reason is typological thinking, not considering intraspecific variability, assuming limited distribution of species, and failing to review literature. The section is reduced from eight species to one. Distribution of the section ranges from Java through Micronesia to Samoa.

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