Tansley Review No. 110 Numerical and physical properties of orchid seeds and their biological implications

JOSEPH ARDITTI¹* AND ABDUL KARIM ABDUL GHANI²

¹Department of Developmental and Cell Biology, University of California, Irvine, CA 92697–2300, USA ²Department of Botany, University Kebangsaan Malaysia, 43600 Bangi, Selangor, Malaysia

Received 1 June 1999; accepted 23 September 1999

CONTR	ENTS			
	Summary	367	(b) Dispersal	387
Ι.	INTRODUCTION	367	(c) Birds	415
II.	NUMBER	368	2. Water	416
III.	Size	379	(a) Physical considerations	416
IV.	AIR SPACE IN THE SEEDS	381	(b) Dispersal	416
V.	FLOATATION AND DISPERSAL	383	VI. CONCLUSIONS	417
	1. Air	383	Acknowledgements	417
	(a) Physical considerations	383	References	418

SUMMARY

Orchid seeds are very small, extremely light and produced in great numbers. Most range in length from c. 0.05 to 6.0 mm, with the difference between the longest and shortest known seeds in the family being 120-fold. The 'widest' seed at 0.9 mm is 90-fold wider than the 'thinnest' one, which measures 0.01 mm (because orchid seeds are tubular or balloon-like, 'wide' and 'thin' actually refer to diameter). Known seed weights extend from 0.31 μ g to 24 μ g (a 78-fold difference). Recorded numbers of seeds per fruit are as high as 4000000 and as low as 20–50 (80000–200000-fold difference). Testae are usually transparent, with outer cell walls that may be smooth or reticulated. Ultrasonic treatments enhance germination, which suggests that the testae can be restrictive. Embryos are even smaller: their volume is substantially smaller than that of the testa. As a result, orchid seeds have large internal air spaces that render them balloon-like. They can float in the air for long periods, a property that facilitates long-distance dispersal. The difficult-to-wet outer surfaces of the testa and large internal air spaces enable the seeds to float on water for prolonged periods. This facilitates distribution through tree effluates and/or small run-off rivulets that may follow rains. Due to their size and characteristics, orchid seeds may also be transported in and on land animals and birds (in fur, feathers or hair, mud on feet, and perhaps also following ingestion).

Key words: orchids, orchid seeds, orchid dispersal, dust seeds, seed floatation, seed production, seed size, seed numbers.

I. INTRODUCTION

Orchids seeds are unique. They differ from those of most angiosperms and resemble the so-called 'dust

*Author for correspondence (tel +1 949 824 5221/+1 949 559 4656; fax +1 949 824 4709/+1 949 559 1231; e-mail jarditti@uci.edu).

seeds' of other plants (Fleischer, 1929, 1930; Ziegenspeck, 1936; Hoene, 1945; Rauh *et al.*, 1975; Rasmussen, 1995). This term probably has its origins as a direct or literal, but not a contextual, translation of the German 'staubförmiger Samen' or 'Staub-Samen. 'Dust seeds' as a term sounds odd in English and may not be completely descriptive, but it is generally accepted, defined to some extent, and in common use – it simply means that the seeds are small. For a more extensive discussion of terminology see Rauh *et al.* (1975) and Barthlott (1976). The seeds were first drawn by the Swiss naturalist Conrad Gesner sometime between 1540 and his death in 1565 (Jacquet, 1994). Since that time, descriptive studies of the seeds form a colourful history (Box 1). Developmental, structural, anatomical, morphological, physiological, ecological, symbiotic and horticultural aspects are discussed elsewhere (Mutsuura *et al.*, 1962; Vij *et al.*, 1981; Weatherhead *et al.*, 1986; Arditti, 1992). The present review will deal only with the purely numerical and physical characteristics of orchid seeds and their biological implications (Figs 1–9; Tables 1–6).

II. NUMBER

Among the best known and most frequently mentioned characteristics of orchids are their small (Figs 1-4; Table 1) and very light (Tables 1-3) seeds, which are usually produced in large numbers (Table 2; Darwin, 1904; Burgeff, 1936; Arditti, 1967, 1979, 1992; Arditti & Ernst, 1984; Rasmussen, 1995). To this day one of the most interesting and perceptive discussions of orchid seed number is by Charles Darwin (Box 2). It is important here as a classic backdrop for modern considerations.

Calculations like Darwin's can result in astronomical numbers (Box 2). Production of large numbers of seeds has been described as a common characteristic in plants that have very specific germination requirements, such as mycotrophy (Rauh et al., 1975; Rasmussen, 1995). These numbers ensure that at least a few seeds will germinate and perpetuate the species - even if Darwin took a dim view of the 'contrivance'. Both the large number and the physical characteristics of orchids facilitate extensive coverage of areas around the seed parent and wider dispersion further away. A consequence of such dispersal is colonization of new and existing favourable sites (Nakamura & Hamada, 1978; Rasmussen, 1995; Murren & Ellison, 1996, 1998; Tremblay, 1997; Carey, 1998; Kull, 1998, Light & MacConaill, 1998; Neiland & Wilcock, 1998; for reviews of evolutionary implications of the nature of orchid seeds see Benzing, 1981, 1987).

It is possible to argue that the production of so many seeds is wasteful of resources and risky, because evolution is unforgiving of waste. Such arguments must be posited against the fact that the orchids are evolutionarily very successful. This success indicates that the survival benefits of begetting many small seeds outweigh the cost of producing them. According to one estimate, the fruit-set in *Cypripedium calceolus* (Figs 3, 7, 8; Tables 1, 2) on the clone level is essentially cost free because ramets that bear fruits 'stay in leaf longer

[and this brings about] compensation on the annual ramet level [because of] its extended growing period (Kull, 1998). Altogether it seems that regardless of the input by individual plants, the resources devoted to the production of many small seeds by orchid species are a good investment. Moreover, orchid embryos are small (Table 1), some consisting of only a few cells; most have no endosperm (Weiss, 1916; Burgeff, 1936; Arditti & Ernst, 1984; Arditti, 1967, 1979, 1992). Their food reserves consist of cellular inclusions like oil droplets and starch grains at levels that are not high in absolute terms. This means that the resources invested by orchids in the production of numerous seeds may not be as large and wasteful as they seem. Even if not cost-free as in Cypripedium calceolus (Fig. 3; Tables 1, 2) the resources expended by orchids may be the same or perhaps even smaller than those invested by other plants in fewer but larger seeds with more reserves.

A comparison between orchid seeds and coconuts (i.e. the seeds of coconut trees) may be instructive. Coconuts are an excellent contrast for orchids because the trees produce relatively few large seeds, which also have lipid reserves. A comparison with medium-sized lipid-containing seeds produced in larger numbers than coconuts (sunflowers or peanuts, for example) would provide less of a contrast. Also, the lipid content of orchids and coconuts is similar. As far as we can determine, comparisons of this type for orchids have not been reported previously. Such comparisons would gain much if data on resources committed to seed production could be related to annual productivity, the locales of seed germination and the likelihood of biophagy. Unfortunately there are not enough published data for more advanced comparisons.

Capsules of Cycnoches ventricosum var. chlorochilon contain 4000000 seeds (Tables 2, 3) each weighing 3.6 µg. The total weight is 14.4 g seeds fruit⁻¹. Assuming that these seeds contain as much lipid energy as fresh coconut solid endosperm (1470 kjoules 100 g⁻¹) the total is still a minuscule 212 kjoules 14.4 g⁻¹ or 0.000419618 kjoules seed⁻¹. This assumption is justified by findings that show that orchid seeds are fatty in nature (as are coconuts) and may contain as much as 32 % lipids (Knudson, 1929; Arditti, 1967, 1979, 1992; Harrison, 1977; Harrison & Arditti, 1978; Arditti & Ernst, 1984).

Fresh solid coconut endosperm ('meat') contains 34% fat and 212 kjoules 14.4 g^{-1} (Diem, 1962). A nut purchased at random in a food store had 380 g 'meat' (5586 kjoules; 26 times as many as a *Cycnoches* seed). It also contained 120 g of liquid endosperm ('coconut water'). Coconuts fresh off a tree contain more 'water.' This liquid is also energy rich and contains a variety of substances such as vitamins, hormones, amino acids and lipids. Thus the total energy content of a coconut, excluding the shell and outer husk, may be as high as 6000 kjoules,

Box 1. A historical perspective

After first being drawn by Gesner (also referred to as Gessner, Gesnerus and Gesneri, 1516–1565; for a likeness see Jaquet, 1994) in the middle of the 16th century, there have been many descriptive studies of orchids undertaken. These first illustrations were intended to be part of *Historia Plantarum*, a book that Gesner planned as a companion for his remarkable *Historia Animalum*. Gesner's early death from the plague prevented completion and publication of *Historia Plantarum*. It also started some of the plates and the manuscript on a two centuries-long odyssey that ended with publication between 1751 and 1759. The tribulations of another set of plates were even longer. They were lost, found in the library of the University of Erlangen (Germany) only after World War II and published between 1972 and 1980 (i.e. about 450 years after they were painted).

Until very recently the existence of drawings of orchid flowers, fruits and seeds (some of them magnified) in Gesner's *Historia Plantarum* was not widely known. Awareness of the seed drawings in the University of Erlangen lost plates was more limited even after they were published. A recent survey (J. Arditti, unpublished) found several plates that contain paintings of seeds.

Georgius Everhardus Rumphius (c. 1627-1702, for a portrait see Arditti, 1984), the so-called Blind Seer of Ambon (an island in the Malukku, formerly Moluccas, Archipelago or Spice Islands, Indonesia), described orchid seeds in his Herbarium Amboinense as a 'yellow flour ... largely shed and blown away on the wind' (de Wit, 1977). Herbarium Amboinense also had a long odyssey (de Wit, 1977). The original was lost in a fire that nearly destroyed the entire city of Ambon. It was reconstituted by assistants because by then Rumphius was blind. The new set was sent to Batavia (now Jakarta) in 1690, copied there for the personal use of the Dutch Governor General and shipped to Holland in 1692. It was lost at sea when the ship that carried it, the Waterland, was sunk by the French. Another copy was made from the one owned by the Governor General. It was shipped to Holland in parts. The complete manuscript reached Amsterdam in 1696-1697. Permission to publish was granted c. 1736 to Johannes Burman (1706–1779), Professor of Botany at the University of Amsterdam for 51 years starting in 1728 (his son Nikolas Laurens Burman, 1734-1793, was also a professor in the same university from 1769 until his death). Burman published six volumes between 1741 and 1750. A companion volume, the Actuarium, was publisued in 1755. Thus, the first description of orchid seeds by Gesner was published 20 years after the second one by Rumphius. However, Gesner's drawings remain the first illustrations of orchid seeds that are known to have been published. Rumphius drew orchid fruits, but never very small flowers and seeds. One reason for this may have been his failing eyesight.

A third set of orchid paintings that includes seeds was also published after a long delay (150 years) in a limited edition of very large and heavy books which are not easily accessible. These books report some of the findings of a Spanish scientific expedition to New Grenada led by José Celestino Bruno Mutis y Bogio (1732–1808). The expedition surveyed the flora of what is now Colombia. Its artists produced more than 200 plates of orchids between 1783 and 1816. The plates languished until the first (introductory) volume in a projected multi-volume series dealing with the Mutis expedition was published (Perez-Arbelaez *et al.*, 1954). Three volumes containing orchid plates were published after that (Schweinfurth *et al.*, 1963; Schweinfurth *et al.*, 1969; Fernandez Perez, 1985). More are to be published. Some of the plates contain excellent paintings of different stages of fruit development, maturation and dehiscence. There are also cross- and longitudinal-sections of fruits that show seeds as white fluffy masses, brown lines and/or dark dots. A number of plates contain drawings of seeds, some small and described as being natural size, others are magnified. Several of the 'natural size' line drawings of fusiform seeds have dots in their middle (and widest) part, which obviously represent embryos. A number of the greatly magnified illustrations of seeds are inaccurate because they look like woolly or fluffy, narrow and pointed ellipses. Small line drawings which show seeds that look like drops or apostrophes also seem unrealistic.

All illustrations in the Flora de la Real expedicion Botanica del Nuevo Reino de Granada were made after the publication of *Herbarium Amboinense* and *Historia plantarum*, but they are still among the earliest representations of orchid seeds (second drawings after Gesner's and third descriptions following *Herbarium Amboinense* and *Historia Plantarum*). They are probably the first clearly to indicate size. The third, as well as most accurate in terms of size and appearance are attractive colour paintings of orchid seeds (Fig. 1a–c), made and published in Austria (Beer, 1863). Strangely, relatively few colour paintings of orchids were published during the last century, but there is an abundance of line drawings with proper size indications (Bernard 1909; Burgeff, 1936; Arditti, 1992; Rasmussen, 1995). Hans Burgeff (1883–1976) was the first to provide weights determined with relatively modern equipment and to measure floatation times (Burgeff, 1936). His measurements were extended through determination of seed volume and air space (Arditti *et al.*, 1979, 1980; Healey *et al.*, 1980).

Photographs (standard, macro and micro) started to appear in the literature as appropriate photographic equipment became available. Due to their dimensions and shape, orchid seeds are difficult to photograph because of greatly reduced depth of field at appropriate magnifications. The first SEM photographs were published in the mid-1970s by W. Barthlott and his associates in Germany. Other SEM studies, structural, morphometric and taxonomic, followed (e.g. Arditti *et al.*, 1979, 1980; Healey *et al.*, 1980; Chase & Pippen, 1988, 1990). The first descriptions and drawings of orchid seedlings were published in 1804 by the British botanist R. A. Salisbury (for a photograph see Arditti, 1984).

Horticultural attempts to germinate orchid seeds date back to the mid-1800s. The first success was reported in 1849 by David Moore (1807–1879, for a portrait see Arditti, 1984), Director of the Glasnevin Botanic Gardens in Ireland (he wrote religion-based anti-Darwin tracts during his later years) and by two British gardeners, J. Cole and R. Gallier, later in the same year. In 1899 Noël Bernard (1874–1911, for photographs and a short biography see Arditti, 1984, 1990), Professor of Botany at the University of Caen, France became the first to describe the role of mycorrhiza in orchid seed germination. Lewis Knudson (1884–1958, for photographs and a biography see Arditti, 1984. 1990), Professor of Botany at Cornell University, formulated methods for asymbiotic germination. He first published his method in 1921 in Spain, in the Spanish language in an (at least presently) obscure journal (Arditti, 1984, 1990, 1992; Jacquet, 1994).

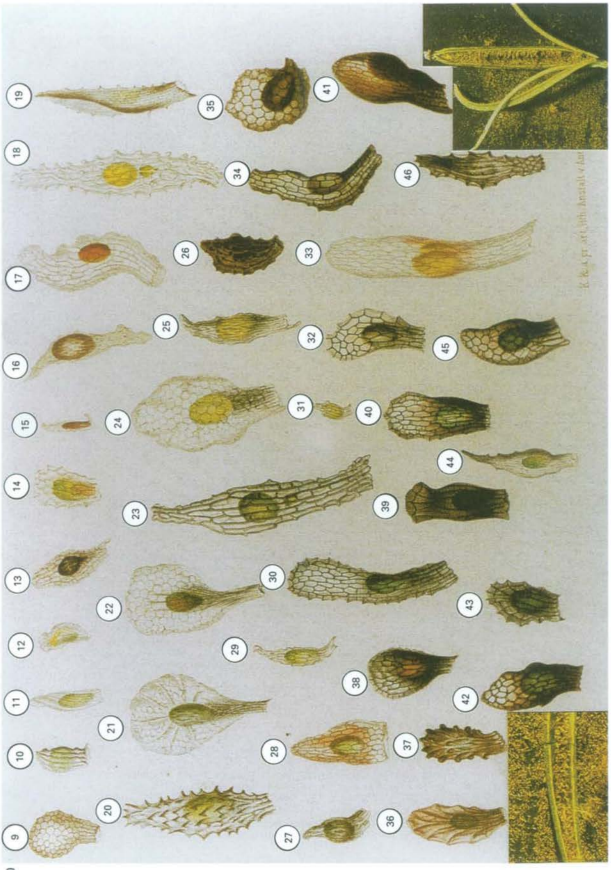


Fig. 1a. For legend see page 372.

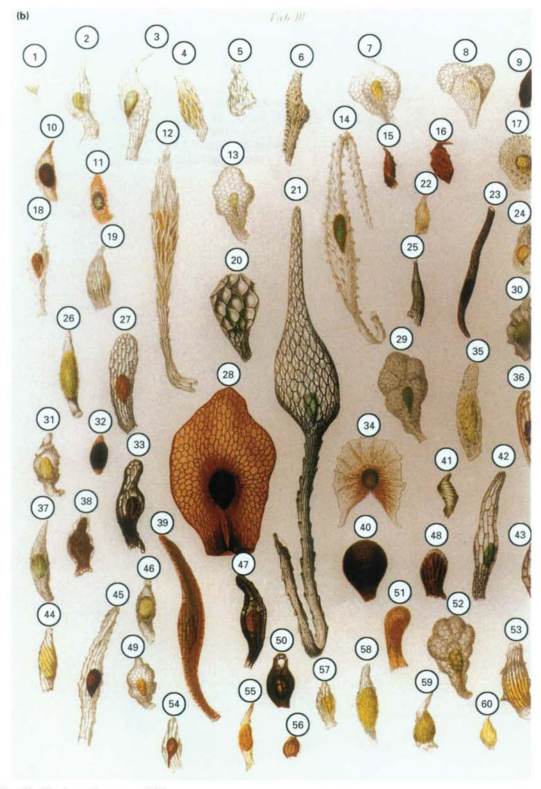


Fig. 1b. For legend see page 372.

which would be the content of 113 207 547 *Cycnoches* seeds (or 2.83 times the number of seeds found in a single capsule; this orchid probably produces more than one capsule per plant only rarely, if at all). Coconut trees may produce up to 75 fruits per plant,

but a reasonable average is 25. If so, 160000 trees would be needed to produce 4000000 seeds, which will contain 240000000 kjoules in the 'water' and 'meat' alone. Not all orchid seeds produce mature plants, but the same is true for coconuts. Therefore

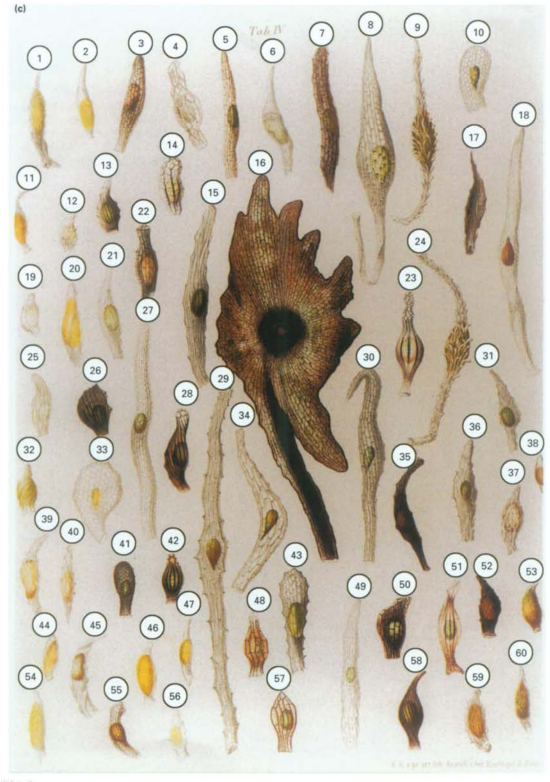


Fig. 1c.

Fig. 1. (Legends for (a-c)). Paintings and photographs of orchid seeds. Note that the first name used in each case is the name in the original source; some current names are also given to ensure accuracy (it is not uncommon to have several taxa lumped into a single concept or one species split into more than one taxon), facilitate identifications and make possible correlations between seed characteristics and different classification approaches. (a) Circled nos. 1-38: 1, Sturmia loesseli (now Liparis loesseli); 3, Octomeria lancifolia; 4, Maxillaria crocea; 5, variety of Dactylorhiza (Orchis) maculata; 6, Gymnadenia conopsea; 7, Angraecum bilobum (now Aerangis biloba, or Aerangis kirkii); 8, Phajus albus (now known as Thunia alba); 9, Stanhopea tigrina var. superba; 9, Listera ovata; 10, Mormodes buccinator; 11, Barkeria melanocaulon; 12, Mormodes viridiflora; 13,

the energy cost per mature offspring is much lower for the orchid. Clearly energy investment in the future of the species by *C. ventricosum* var. *chlorochilon* is more efficient than that of coconuts. Similar comparisons with other fatty seeds should prove interesting.

Another benefit of the relatively small energy investment in numerous seeds by the orchid is the

Stanhopea warczewitzii; 14, Stanhopea tigrina; 15, Mormodes pardina var. unicolor (now Mormodes maculatum); 16, Stanhopea aurea (now Stanhopea wardii); 17, Listera ovata 18, Himantoglossum hircinum; 19, Tetragamestus modestus (genus now known as Scaphyglottis). 20, Orchis longicornu; 21, Ornithocephallus; 22, Corallorhiza innata; 23, Scaphyglottis vestita; 24, Orchis maculata (Dactylorhiza maculata); 25, Epipactis latifolia; 26, Pyrola rotundifolia (this is not an orchid - it belongs to the Pyrolaceae); 27, Orchis coriophora; 28, Orchis secundiflora; 29, Odontoglossum sphacelatum (now Oncidium sphacelatum); 30, Orchis fragrans; 31, Orchis brevicornu (correct specific epithet is brevicornis; species now known as Orchis patens); 32, Orchis latifolia (now Dactylorhiza majalis); 33, Neottia nidus-avis; 34, Nigritella angustifolia (now Nigritella nigra); 35, Orchis intacta (now Neotinea); 36, Epidendrum stamfordianum; 37, Gymnadenia longifolia; 38, Cypripedium barbatum (now Paphiopedilum barbatum). The insert at bottom left shows magnified seeds inside an opened Sarcochilus capsule; the insert at bottom right shows a dehisced fruit of Sarcochilus at about natural size. (b) Circled nos. 1-60: 2, Calanthe veratrifolia; 3, Phajus grandifolius (now Phaius tankervilleae); 4, Zygopetalum mackayi; 5, Zygopetallum intermedium; 6, Bletia sheperdii (now Bletia verecunda); 7, Stanhopea insignis; 8, Stanhopea oculata; 9, Thelymitra ixioides; 10, Leptotes bicolor; 11, Promenaea stapelioides; 12, Govenia lilacina (correct name is Govenia liliacea); 13, Gongora maculata var pallida; 14, Sobralia liliastrum; 15, Aerides odoratum; 18, Cattleya amethystina (now Cattleya intermedia); 19, Dicrypta glaucescens var. Hort. (genus now known as Maxillaria); 20, Eulophia streptopetala; 21, Epidendrum cinnabarinum; 22, Miltonia morelliana (now Miltonia spectabilis var. morelliana); 23, Goodyera procera; 24, Acropera loddigesii (now Galeola galeata); 25, Brassia cowanii (now Brassia verrucosa); 26, Cattleya bicolor; 27, Cymbidium odontorhizon (several reference works do not list such an orchid); 28, Epistephium parviflorum; 29, Gongora bufonia; 30, Acropera (Gongora) luteola; 31, Acropera maculata (now Gongora quinquenervis); 32, Luisia teretifolia; 33, Orchis acuminata (at one time this orchid was classified as Orchis tridentata var. lactea; more recently it was listed as Orchis lactea); 34, Haematorchis altissima (at one time Galeola altissima and now Erythorchis altissima); 35, Odontoglossum bictoniense; 36, Habenaria tridentata; 37, Epidendrum cochleatum; 38, Coricium crispum (genus has been/is also known as Pterygodium, Ommatodium and Anochilus); 38, Orobanche, Orobanchaceae (broomrape, a nonorchidaceous plant); 39, Goodyera discolor (now Ludisia discolor); 40, Vanilla planifolia; 41, Odontoglossum pulvinatum (actually Oncidium pulvinatum); 42, Phajus maculatus (now Phaius flavus); 43, Phajus bicolor (now Phaius tankervilleae); 44, Trichopilia albida (now known as Trichopilia fragrans); 45, Neottia speciosa (now Stenorhyncus speciosa); 46, Epidendrum lancifolium (now Epidendrum cochleatum); 47, no caption in original; 48, Habenaria hispidula; 49, Triphora pendula (several sources treat Triphora as a synonym of Pogonia); 50, Corycium orobanchoides (also known as Pterigodium orobanchoides); 51, Neottia aestivalis (now Spiranthes aestivalis); 52, Cirrhaea viridi purpurea (may well be a hybrid between Cirrhaea viridifusca and Cirrhaea purpurascens; the other names for this genus, Scleropteris and Sarcoglossum, go in and out of fashion); 53, Acropera citrina (now Gongora galeata); 54, Disperis villosa; 55, Sarcanthus rostratus (genus now part of Cleisostoma, the species was also known as Vanda recurva and Vanda rostrata); 56, Dicrypta tenella (genus now known as Maxillaria); 57, Promenaea rollinsonni; 58, Cattleya tigrina (now Cattleya guttata); 59, Dicrypta glaucescens (genus now known as Maxillaria); 60, Sarcanthus violacea (one possibility is that this is now Rhyncostylis violacea, which was variously known as Anota violacea, Saccolabium violaceum and Vanda violacea - there is a very large number of transfers between these genera). (c) Circled nos. 1-60: 1, Pholidota rubra (current name unknown); 2, Epidendrum ramosum; 3, Goodyera repens; 4, Goodyera semipelucida; 5, Neottia pubescens (now a synonym of Goodyera pubescens); 6, Laelia galeottiana (current name unknown); 7, Gamoplexis orobanchoides (Gamoplexis is a synonym of Gastrodia); 8, Epidendrum crassifolium (but also Epidendrum crassifolium and Epidendrum ellipticum); 9, Sobralia decora; 10, Habenaria dilatata (now Platanthera dilatata); 11, Otochilus fusca; 12, Dendrobium plicatile (now Flickingeria fimbriata); 13, Cerathandra chloroleuca (genus name is spelled Ceratandra; present name Ceratandra atrata); 14, Disa cernua; 16, Cyrtosia lindleyana (current genus name unknown); 17, Cattleya crispa (now Laelia crispa); 19, Xylobium squalens; 20, Laelia anceps; 21, Epidendrum ciliare; 22, Satyrium nepalense; 23, Disa cornuta; 24, Sobralia macrantha; 25, Huntleya violacea (also known as Bollea violacea); 26, Ophrys funerea (a synonym of Ophrys fusca); 27, Pleurothalis sessiliflorum (current name unknown); 28, Satyrium carneum; 29, Acanthophippium bicolor; 30, Cymbidium sinense; 31, Cattleya harrisonii (now Cattleya harrisoniana); 32, Agraphyllum sp.; 33, Stanhopea violacea; 34, Phajus wallichii (synonym of Phaius tankervilliae); 35, Selenipedium schlimii (now a synonym of Phragmipedium schlimii); 36, Laelia perinmii; 37, Epidendrum papillosum (or Encyclia alata); 38, Isochilus lineraris; 39, Cattleya loddigesii; 40, Cattleya forbesii; 41, Malaxis liliifolia (now a synonym of Liparis liliifolia); 42, Disa pulchella; 43, Neottia vitalis (current name unknown); 44, Trichocentrum fuscum; 45, Epidendrum stamfordianum var. carnea; 46, Dendrobium cretaceum; 47, no caption in original; 48, Pterygodium inversum; 49, Pelexia adnata (current name unknown); 50, Pterygodium volucre (correct name Pterygodium volucris); 51, Pterygodium catholicum; 52, Acroclaene punctata; 53, Vanda coerulea; 54, Brassavola cordata; 55, Apaturia senilis (correct spelling Apatura; since at present this genus is considered to be a synonym of Pachystoma, the species name becomes Pachystoma senile, which in turn is reduced to synonymy under Pachystoma pubescens); 56, Dicrypta bauerii (now Maxillaria crassifolia); 57, Satyrium bicallosum (current name unknown); 58, Lycaste harrissonii (correct name Lycaste harrisoniae; now Bifrenaria harrisoniae); 59, Otochilus porrecta; 60, Aerides sp. Seed paintings ($\times 100$ in the original) are from Beer, 1863. Magnifications here are: (a) $\times 109$; 1(b) $\times 84$; 1(c) $\times 81$. Original colour prints, reprinted, with permission, from Arditti (1993). Photographic inserts in (a) are from Upton (1992), which does not provide a magnification factor.

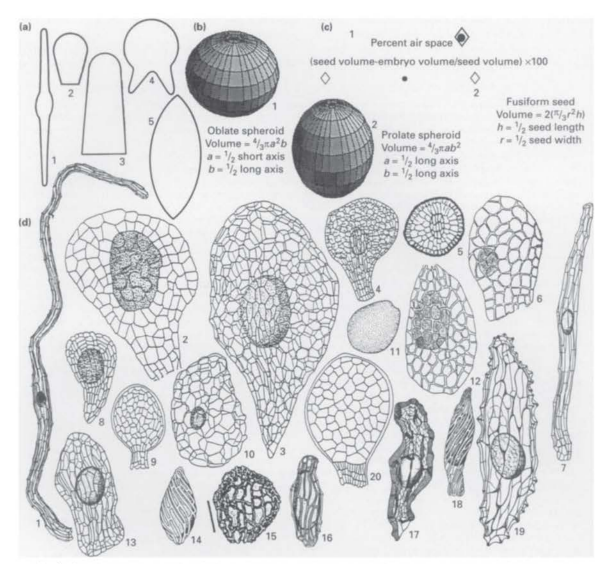


Fig. 2. Orchid seeds and embryos: general shapes, geometry and line drawings. (a) Five general shapes into which orchid seeds have been classified (Clifford & Smith, 1969). The most common are shapes 1 and 5. They are followed by forms 2 and 3. Form 4 is relatively rare. Form 5, unidentified. (b) Orchid embryos are prolate (1) not oblate (2) spheroids. (c) Fusiform orchid seeds: formula and diagram of percent free air space (1), together with formula used to calculate the volume of this form (2). (d) Line drawings of orchid seeds: 1, Acanthephippium sylhetense, ×75; 2, Stanhopea tigrina, magnification unknown; 3, Stanhopea oculata, ×150; 4, Stanhopea oculata, magnification unknown (two drawings from different sources are presented here to show that there is variability in seed form and size); 5, Vanilla planifolia, cross section diameter 0.367 mm; 6, Listera australis, ×97; 7, Haemaria discolor (now Ludisia discolor), ×89; 8, Pleione formosana (now Pleione bulbocodioides), magnification unknown; 9, Zygopetalum intermedium, magnification unknown; 10, Epipogium nutans (now Epipogium roseum), ×166; 11, Vanilla planifolia, magnification unknown; 12, Listera convallarioides, ×97; 13, Liparis loeselii, ×113; 14, Vanda, ×80; 15, Schizochilus zeyheri, bar, 0.1 mm; 16, Lycaste skinneri, $\times 111.5$; 17, Serapias cordigera, $\times 83$; 18, Dendrobium, $\times 108$; 19, Catasetum macrocarpum, × 128; 20, Stanhopea saccata, magnification unknown. Sources: 1, 3, 5, 7, 10, 12, 13, 14, 17, 16, 18, 19 - Burgeff, 1936; 4, 11 - Poudubnaya-Arnoldi & Selezneva, 1957b; 2, 8, 9, 20 - Cherevchenko & Kushnir, 1986; 6, 11 -Curtiss, 1893; 15 - Kurzweil, 1993.

minute size and unique structure of these seeds. Coconuts, being large, heavy, with a hard shell and spongy husk, can only be dispersed through transport on land (which is not always easy for a fruit of this nature) or floatation (sometimes for prolonged periods in oceans with no certainty of reaching suitable land). However, orchid seeds can be transported on, and perhaps in, the bodies of land animals (fur, hair, mud on parts of the body) and birds (feathers or mud on their feet) as well as by floatation in water or air (Went, 1957; Gandawijaja & Arditti, 1983; Arditti, 1992).

Altogether, orchids derive important survival benefits from a penurious approach to seed production. Individual capsules may contain many seeds (Table 2), but some species and populations set fruit

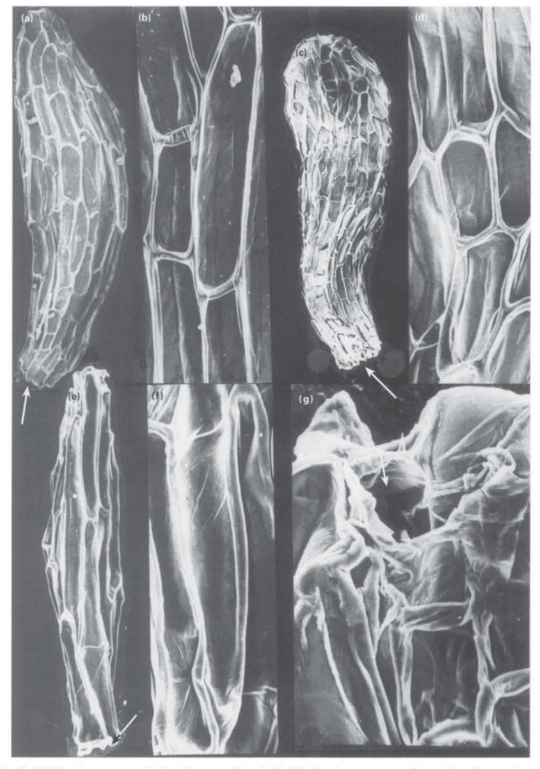


Fig. 3. SEMs of mature seeds (a,c,e), testa cell walls (b,d,f,g) and suspensor end openings (arrows in a,c, e and g) of Cypripedium calceolus var. parviflorum (a, $\times 110$; b, $\times 465$), Cypripedium reginae (c, $\times 55$; d, $\times 4600$) and Paphiopedilum cv. Susan Tucker \times Paphiopedilum parishii (e, $\times 327$; f, $\times 780$; g, $\times 1200$). Testa cell walls are smooth or nearly so. In (g), the opening in the suspensor end of the seed that is shown is wide enough to allow entry of water and fungal hyphae. Reproduced, with permission, from Arditti et al. (1979).

sparingly, one example being *Lepanthes caritensis* (Tremblay, 1997). Recent studies have shown that there are several factors that limit fruit set and seed

production in orchids (Willems, 1992). Pollinator availability is one such factor in the following: Aspasia principissa (Zimmerman & Aide, 1989);

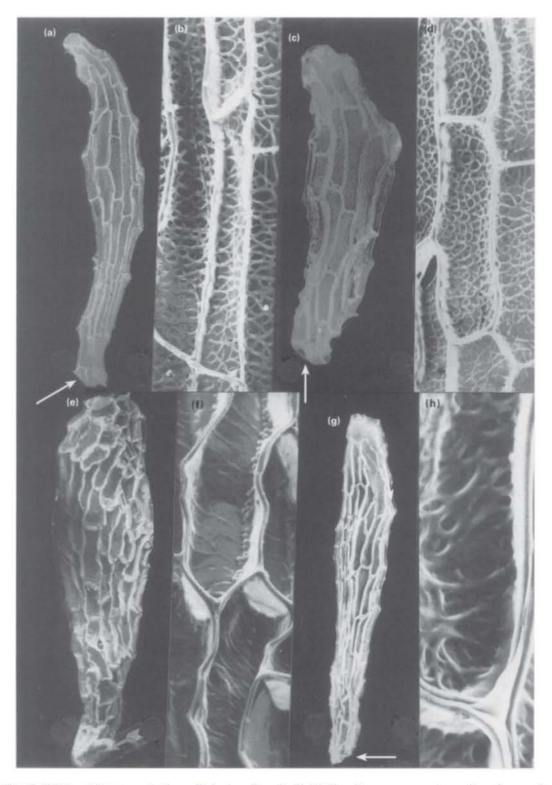


Fig. 4. SEMs of intact seeds (a,c,e,g), testa cell walls (b,d,f,h) and suspensor end openings (arrows in a,c,e,g) of California (a, $\times 135$; b, $\times 862$) and Colorado (c, $\times 260$; d, $\times 673$) plants of *Calypso bulbosa*. Testa cell walls (b,c) are reticulated, but reticulations as well as seed shape and size from the two populations are different. The California seeds are mature, those from Colorado are not. Seeds of *Corallorhiza maculata* (e-h: e, $\times 100$; f, $\times 47$; g, $\times 90$; h, $\times 1200$) differ in shape and reticulations *Reproduced, with permission, from Arditti* et al. (1980).

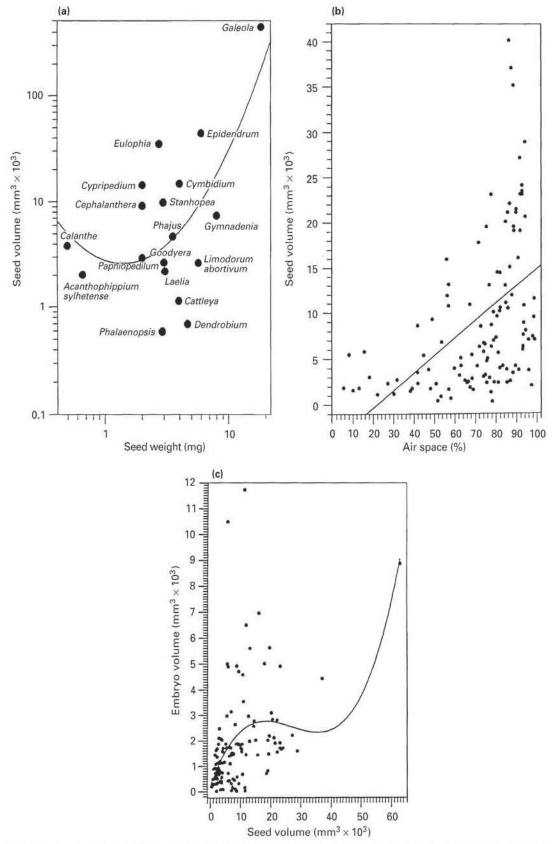


Fig. 5. Volume, percent free air space and weight of orchid seeds. (a) Volume is calculated as $2(\pi/3r^2h)$. Except in very light seeds volume increases with weight. (b) Percent air space is given as ((seed volume–embryo volume/seed volume) × 100). This increases with seed volume. There are 119 data points in this plot. (c) Seed volume vs embryo volume. There are 123 data points in this plot. The lines were generated with SigmaPlot* (SPSS Inc., Chicago, IL, USA).

Dactylorhiza (Fig. 1a; Tables 2, 3); Goodyera (Fig. 1b,c; Tables 1–3); Ionopsis utricularioides (Montalvo & Ackerman, 1987); Ophrys (Fig. 1c; Table 3); Orchis (Fig. 1a,c; Tables 1–3); Platanthera (Fig. 1c;

Tables 1–3); *Serapias* (Fig. 2) in Europe (Neiland & Wilcock, 1995); and other orchids (Ackerman & Zimmerman, 1994). Pollen load may be a determinant in the case of *Calypso bulbosa* var. *americana*,

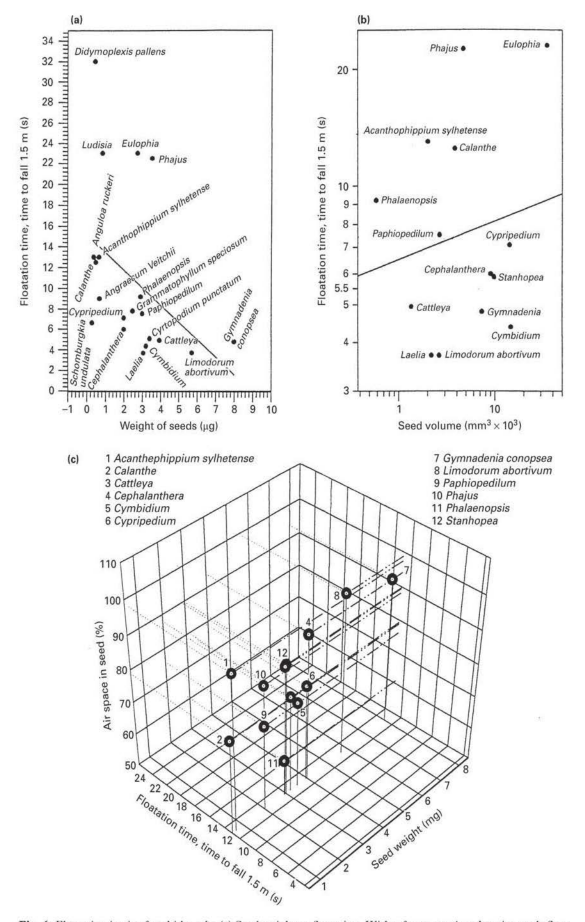


Fig. 6. Floatation in air of orchid seeds. (a) Seed weight vs floatation. With a few exceptions heavier seeds float for shorter periods. Mean weight of seeds 2.873 μ g (SD 2.17); mean floatation time 9.973 s (SD 7.64). (b) Seed volume vs. floatation. Larger volumes increase floatation time. (c) Floatation vs seed weight and percent air space. Larger seed volumes and smaller weights increase floatation time. Lines generated with SigmaPlot[®].

but not *Cypripedium* (Figs 3, 7; Tables 1, 2), and *Amerorchis* (Orchis) rotundifolia (Table 3; Proctor & Harder, 1994). Unknown and/or compatibility factors may also be involved because one third of handpollinated flowers of *Ionopsis utricularis* (utricaroides) failed to set fruits in field experiments in Puerto Rico (Montalvo & Ackerman, 1987), and additional pollination could not augment fruit production in *Oeceoclades maculata* (Table 2; Gonzales-Diaz & Ackerman, 1988).

Resource limitations and prior fruiting play a major role in reduced fruit and seed set in orchids. Fruit and seed production during one year can reduce growth, number of flowers, fruit set and seed numbers during the following season (Montalvo & Ackerman, 1987; Snow & Whigham, 1989; Zimmerman & Aide, 1989; Ackerman & Zimmerman, 1994). This occurs in Aspasia principissa (Zimmerman & Aide, 1989). Fruit set itself may be a limiting factor in Cypripedium (Figs 3, 7; Tables 1, 2; Nilsson, 1979). For example, fruiting by 3500 flowers of Cypripedium calceolus (Figs 3, 7; Tables 1, 2) in eight Estonian populations over 11 yr was 10.5% (Kull, 1998). Fruit set by the neotropical Brassavola nodosa (Table 3) was 30 % in the summer, but less during other seasons for a total of 13 % over a seven month period (Murren & Ellison, 1996). On the average fruit set by nectarless and nectariferous orchids in North America is 19.5% and 49.3%, respectively. Equivalent figures are 27.7% and 63.1% for Europe, 41.4% and 74.4% in the southern temperate hemisphere, and 11.5% and 24.9% for the tropics (Neiland & Wilcock, 1998). Darwin also devoted some thought to this subject (Box 3).

III. SIZE

As already mentioned, orchid seeds are very small (Beer, 1863; Burgeff, 1936; Ziegenspeck, 1936; Poddubnaya-Arnoldi & Selezneva, 1953, 1957a,b; Arditti, 1967, 1979, 1992; Rauh et al., 1975; Arditti et al., 1982; Arditti & Ernst, 1984; Rasmussen, 1995; Figs 1-4; Tables 1, 2). Variability in size is considerable in the family, in genera and even within species. The longest seed (6.0 mm), that of the neotropic Epidendrum secundum type is 120 times longer than the shortest (0.05), which is produced by the New Caledonian Anoectochilus imitans (Hallé, 1977). Note that Epidendrum secundum was or could be assigned to three different genera - such taxonomic convolutions are common in orchids. Hence this orchid was known as Stenocorvne secunda, but could also be Bifrenaria secunda because Rudolph Schlechter (1872-1925) as well as more recent taxonomists consider Stenocorvne to be a synonym of Bifrenaria.

Other long seeds are those of: *Lecanorchis japonica* (4.28 mm; Table 1; Kim *et al.*, 1979); *Epidendrum*

Very short seeds are produced by: Anoectochilus imitans (0.050 mm; Hallé, 1977); Aplectrum hyemale (0.110 mm; Table 3; Stoutamire, 1981); Dichaea (0.120 mm; Barthlott & Ziegler, 1981); Earina valida (0.150–0.200 mm; Hallé, 1977); Liparis elliptica (0.150 mm; Hallé, 1977); several Phraetia (Table 1) spp. (0.120–0.170 mm; Hallé, 1977); Schoenorchis micrantha (0.150 mm; Hallé, 1977); and Vanilla planifolia (Figs 1b, 2; Table 1), although seed length in this species may vary between 0.133 and 0.300 mm (Bouriquet, 1947).

Among the 'widest' seeds are those of *Dendrobium insigne* (0.9 mm; Benzing & Clements, 1991) and *Galeola nudifolia* (0.93 mm; Table 1; Barthlott & Ziegler, 1981). They are 90 times 'wider' than the 'thinnest' ones, those of the *Gastrodia* type (Fig. 1c; Dressler, 1993), which measure 0.01–0.1 mm. (Since orchid seeds are actually tubular, 'wide' and 'thin' are terms of convenience that refer to diameter, which is technically very difficult to measure.)

Other orchids which produce slender seeds include terrestrial species (0.07-0.4 mm; Tables 1, 2; Rasmussen, 1995; the assertion by Rasmussen that seeds of epiphytic species are 'smaller' than those of terrestrial ones is incorrect). These include: Acanthephippium (sometimes spelled Acanthophippium) sylhetense (Fig. 2, Tables 1, 4; 0.07 mm; Burgeff, 1936); Cattleya aurantiaca (Table 3; 0.08 mm; Arditti, 1992); Chiloschista lunifera (0.05 mm; Barthlott & Ziegler, 1980); Dendrobium nobile (Tables 1, 3; 0.08 mm; Ito, 1955); Goodyera pubescens (Fig. 1c, Tables 1,3; 0.08-0.1 mm; Curtiss, 1893; Arditti, 1992); Haemaria discolor (now Ludisia discolor); 0.09 mm; Figs 1b, 2, Table 3; Burgeff, 1936); Hormidium coriaceum (0.09 mm; Jordão et al., 1988); Leochilus scriptus (0.05 mm; Chase & Pippen, 1988); Lockhartia oerstedii (0.075 mm; Chase & Pippen, 1988); Maxillaria rufescens (0.075 mm; Chase & Pippen, 1988); Miltonia spectabilis (0.062 mm; Chase & Pippen, 1988); Mormolyca ringens (0.034 mm; Chase & Pippen, 1988); Orchis anatolica (0.027 mm; Tables 1,4; Wildhaber, 1972); Pachyphyllum muscoides (0.075 mm; Chase & Pippen, 1988); Phalaenopsis (0.075-0.96 mm; Tables 1, 3; Linder, 1930; Bernard, 1937); Thunia marshalliana (0.091 mm; Table 3; Henriksson, 1951); Trichocentrum panduratum (0.061 mm; Henriksson, 1951); Zeuxine elongata (0.074 mm; Table 1; Molvray & Kores, 1995).

Orchid embryos are also very small (Table 1). Some of the smallest embryos are produced by the following: *Acanthephippium sylhetense* (0.058×0.042 mm; Fig. 2, Tables 1, 3; Burgeff, 1936); *Pogonia triantophorus* (now *trianthophora*; 0.08×0.06 mm; Table 1; Curtiss, 1893); and *Vanilla planifolia*

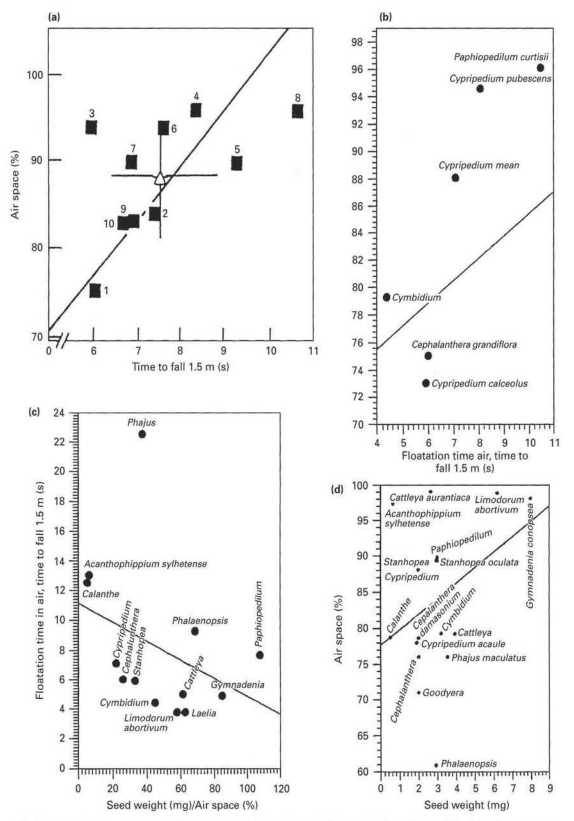


Fig. 7. Floatation of orchid seeds in air vs percent air space and weight. (a) Floatation time vs air space in seeds of California orchids (graph and statistics – Gandawijaja & Arditti (1983); floatation time – Burgeff (1936); air space – Arditti *et al.* (1979, 1980), Healey *et al.* (1980)). Nos. 1–10: 1, *Cephalanthera grandiflora* (now *Cephalanthera damasonium*: percent air space, 75; seconds to fall 1.5 m, 6.0); 2, *Corallorhiza innata* (83 %, 7.2 s); 3, *Cypripedium calceolus* var. *parviflorum* (93 %, 5.9 s); 4, *Cypripedium calceolus* var. *pubescens* (96 %, 8.1 s); 5, *Epipactis latifolia* (now *Epipactis helleborine*: 90 %, 9.1 s); 6, *Epipactis palustris* (93 %, 7.5 s); 7, *Epipactis rubiginosa* (now *Epipactis atrorubens*: 90 %, 6.8 s); 8, *Paphiopedilum curtisii* (now *Paphiopedilum superbiens*: 96 %, 10.5 s); 9, *Platanthera bifolia* (82 %, 6.8 s); 10, *Platanthera chlorantha* (82 %, 6.6 s). Mean percent air space for these orchids is 88 ± 6.7 . The mean time to drop 1.5 m is 7.5 ± 1.4 s. The correlation coefficient is 0.5%. The open triangle indicates the point at which the means intersect; vertical bar is SD for percent air space; horizontal bar is SD for floatation time. (b) Floatation time of orchid seeds of similar weight with different percentages of free air space. Time of floatation increases with percent air space. Mean weights: *Paphiopedilum*

 $(0.1 \times 0.078 \text{ mm}; \text{Fig. 1b}, 2, \text{Table 1}; \text{Bouriquet}, 1947)$. Large or small, the great majority of orchid embryos do not have a cotyledon or an endosperm (Arditti, 1967, 1979, 1992; Arditti & Ernst, 1984).

IV. AIR SPACE IN THE SEEDS

Most orchid seeds are fusiform (Figs 1–4) and approximate two cones joined at their bases. Therefore their internal volume can be estimated with the formula:

$$2[(w/2)^2 \cdot (0.5l) \cdot (1.047)] = 2\left(\frac{\pi}{3}r^2h\right)$$
$$= 2(1.047r^2h) \qquad \text{Eqn 1}$$

(w, width; $(w/2)^2$, radius²; *l*, seed length, 0.5*l*, *h*(eight of each cone); $\pi/3$, 1.047). The volume of relatively few orchid seeds has been calculated. It can range from $< 2.00 \text{ mm}^3 \times 1000^3$ to $40 \times 1000^3 \text{ mm}^3$ (Table 1).

Prolate spheroids (Fig. 2b) are formed by rotating an ellipse about its major axis. As even cursory inspections with dissecting microscopes indicate, orchid embryos (Figs 1, 2d) are prolate spheroids. Moreover, their longer (main or major) and biological (smaller cells at the tip, larger ones at the suspensor end) axes are one and the same. The volume of prolate spheroids (Fig. 2b) is:

$$\frac{4}{3}\pi ab^2 = 4.19ab^2$$
 Eqn 2

 $(a = 0.5 \times \text{length} \text{ (or major axis)}; b = 0.5 \times \text{width} \text{ (or minor axis.)} \text{ Oblate spheroids (Fig. 2b) are formed when an ellipse rotates around its minor axis. Orchid seeds are clearly not oblate spheroids despite erroneous assertions to the contrary (Stoutamire, 1981). The volume of oblate spheroids is:$

$$\frac{4}{3}\pi a^2 b = 4.19a^2 b$$
 Eqn 3

 $(a = 0.5 \times \text{length} \text{ (or major axis)}; b = 0.5 \times \text{width} \text{ (or minor axis)}.$ An embryo with semi axes of a = 0.16 mm and b = 0.1 mm will have a volume of 0.670 mm³ if treated as a prolate spheroid and only 0.107 mm³ as an oblate one. The difference (6.26-fold in this example) is too large to ignore. Embryo volumes range from c. 0.0001 mm³ × 1000³ to c. 5.0 mm³ × 1000³ (Table 1). Note that a number of embryo volumes calculated as $4/3 \pi a^2 b$ (on the erroneous

assumption that they are oblate spheroids) are inaccurate and misleading, but cannot be recalculated because the lengths of the a and b semiaxes are not given in the original source (Stoutamire, 1983).

In addition to being small, orchid seeds are also very light, with the heaviest known being produced by the Asia-Pacific genus Galeola (14-17 µg) and the lightest coming from the South American Anguloa (0.3-0.4 µg). The ratio between the lightest and highest known weights is c. 40-60. Because of their large internal air space, such light seeds are very buoyant in both air and water. An estimate of this buoyancy as w/v is analogous to density. This means that heavier and larger seeds which have relatively bigger air spaces may be as or more buoyant (less 'dense') than more diminutive and lighter seeds with proportionally smaller air spaces. Also, the trend is for total seed volume (Fig. 5a) and percent air space (Fig. 7d) to increase as seeds become heavier and longer (Fig. 8a), but not necessarily wider (Fig. 8b,c).

Seed volumes do increase when embryo volumes are relatively small, but they quickly reach a plateau and further increases, although large, are limited to a few seeds (Fig. 5c). These observations (Figs. 5, 7a, 8a) suggest that embryos may become heavier and/or larger as seeds grow bigger and roomier, but the increases may not be directly proportional to seed volume. Given the large number of orchid species and the paucity of information (i.e. limited number of data points in the graphs, the largest being 123, in Fig. 5c, which is only a small fraction of the 25000-30000 orchid species) the generalizations presented here should be treated with caution. They may change as more information becomes available, but with the extent and direction of orchid research being what it is at present not much can be expected in the near future.

Free air space in seeds is variable (Tables 1,3). At 8.2%, the free air space in *Dactylorchis foliosa* is among the smallest, whereas the largest is 99% in *Cattleya* (a ratio of 12:1). Volume : weight ratios in orchid seeds have been described as being high due to the so-called scobiform shape of the seeds, without definition of the term numerically or provision of data to support the statement (Hirt, 1906; Rasmussen, 1995). A dictionary definition of 'scobiform' is 'resembling sawdust or filings', but orchid seeds do not resemble either. Other terms which should not be used to describe orchid seeds are filing-like (*Feilspansamen*) and sawdust-like (*Säges*-

seeds, $3.0 \ \mu g$; *Cephalanthera* seeds, $2.0 \ \mu g$; *Cypripedium* seeds, $2.0 \ \mu g$; *Cymbidium* seeds, $3.2 \ \mu g$. (c) Floatation time as affected by the ratio between weight and percent air space (i.e. unit weight per unit free volume). (d) Seed weight vs percent air space. The regression line is steep because some seeds have a relatively large free air space.

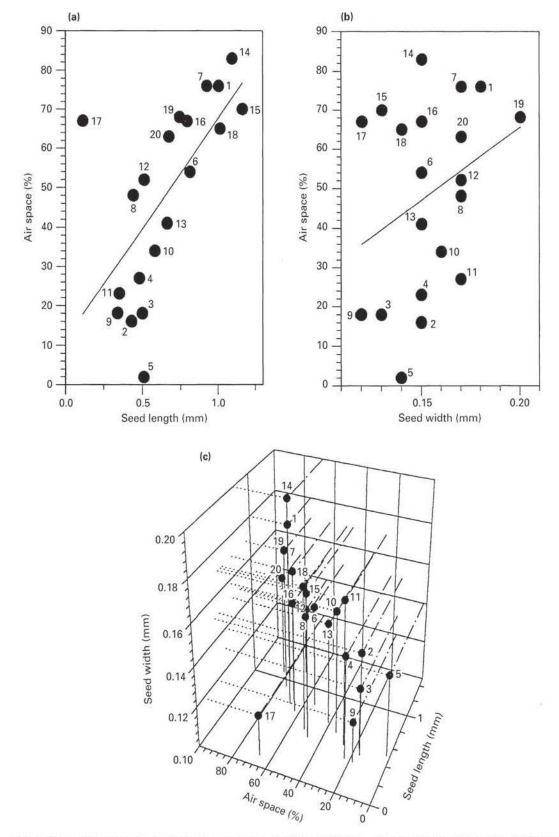


Fig. 8. Testa dimensions and percent free air space in *Paphiopedilum* seeds. (a) Length vs air space. (b) Width vs air space. (c) Percent air space vs seed width and length. Species: 1, *P. delenatii*; 2, *P. belatulum*; 3, *P. concolor*; 4, *P. godefroyae*; 5, *P. philippinense*; 6, *P. rothschildianum*; 7, *P. stonei*; 8, *P. haynaldianum*; 9, *P. lowii*; 10, *P. hisutissimum*; 11, *P. insigne*; 12, *P. fairrieanum*; 13, *P. druryi*; 14, *P. callosum*; 15, *P. ciliolare*; 16, *P. acmodontum*; 17, *P. purpuratum*; 18, *P. venustum*; 19, *P. glaucophyllum*; 20, *P. lemianum* var. *purpuratum*.

pansamen; Rauh *et al.*, 1975; Barthlott, 1976). A point to keep in mind is that dust, filings or sawdust have no internal air spaces. Furthermore, volume :

weight ratios can be high or low even in seeds or other objects that are not scobiform. Volume : weight (Fig. 5a), air space : volume (Fig. 5b) and air

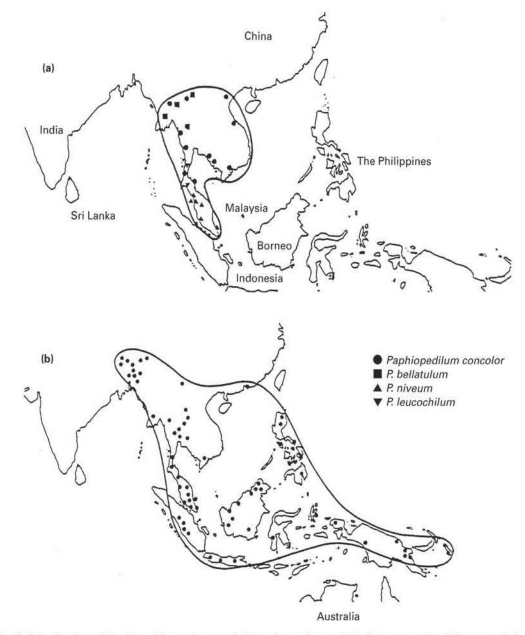


Fig. 9. Distribution of *Paphiopedilum* subgenera (a) *Brachypetalum* and (b) *Sigmatopetalum* (Karasawa & Saito, 1982).

space : weight ratios (Fig. 7d) are determined mostly by testa/embryo length (Fig. 8a,c) and to some extent width (Fig. 8b) relations, not by their shapes (Fig. 2a) and can vary considerably.

V. FLOATATION AND DISPERSAL

1. Air

(a) *Physical considerations.* Because of their small size, shape and large air space, orchid seeds can float in the air for long periods (Figs 6a-c, 7a-d; Tables 1-3). Hans Burgeff seems to have been the first to study the floatation of orchid seed in the air (Burgeff, 1936). Unfortunately very few if any studies of orchid seed floatation were carried out after Burgeff.

Therefore additional quantitative information on the subject is not available. Burgeff assumed that the ability of seeds to float in the air depended on their weight. This is true for Didymoplexis pallens, Ludisia and Gymnadenia (Figs 6a-c, 7c), but not for Schomburgkia (Fig. 6a). Heavier seeds like those of Phaius and to some extent Eulophia float for longer than the lighter ones of Cephalanthera and Laelia, for example (Figs 6a-c, 7c). Seed volume (Fig. 6b) and the percentage of free air space inside the seeds (Fig. 7a,b) seem to be more important as factors that affect and perhaps even determine floatation time in air (Tables 1, 3, 4). This is clearly the case with Paphiopedilum curtissii seeds, which are heavier than those of Cypripedium calceolus var. pubescens (Cypripedium pubescens) but have a higher percentage of air space and float longer (Fig. 7b). Cymbidium seeds

Table 1. Physical characteristics of some orchid seeds

	Dimensions						Free	
	Seed			Embryo			air	
Orchid	Length (mm)	Width (mm)	Volume (mm ³)	Length (mm)	Width (mm)	Volume (mm ³)	space in testa (%)	Weight (µg)
Acanthephippium	1.61 ± 0.36							
Acianthus	1.01 ± 0.49							
Anacamptys	0.4 ± 0.01							
Bletilla	0.43 ± 0.29	0.18 ± 0.06		0.27 ± 0.12	0.12 ± 0.05			
Bletilla striata	0.19 ± 0.07	-						
Bulbophyllum	0.37 ± 0.19	0.2 ± 0.10	5.90 ± 4.88	0.32 ± 0.08	0.2 ± 0.11			
Calanthe	0.77 ± 0.21	0.14 ± 0.25	3.81 ± 0.94	0.14 ± 0.02	0.09 ± 0.02	0.87 ± 0.57	77.67 ± 8.28	
Calopogon	0.07 1 0.01	0.72 ± 0.03	0.01 - 0.77	0.11 1 0.00	0107 1 0102	0101 1 0101	11101 10120	
Calypso bulbosa*	0.8 ± 0.4	0.14 ± 0.01	2.5 ± 0.66	0.14 ± 0.02	0.08 ± 0.01			
Catasetum	0.0 1 0.1	0.45 ± 0.09	0.12 ± 0.02	1.7	0.00 1 0.01			
Cattleva	0.70 ± 0.29	0.16 ± 0.18	1.25 ± 0.99	0.12 ± 0.11				
Cephalanthera damasonium	0.94 ± 0.2	0.23 ± 0.04	13.1 ± 5.1	0.12 ± 0.11				
Corallorhiza	0.90 ± 0.44	0.18 ± 0.04	8.80 ± 2.28	0.24 ± 0.23	0.11 ± 0.02	5.85 ± 4.02	79 ± 6.75	
Cryptostylis	0.63 ± 0.26	0.10 10.04	0.00 1 2.20	0.24 10.25	0.11 1 0.02	5.05 1 4.02	17 10.75	
Cymbidium	0.94 ± 0.33	0.22 ± 0.08	14.68 ± 4.73	0.25 ± 0.06	0.15 ± 0.05			3.2 ± 1.6
Cypripedium	1.07 ± 0.43	0.22 ± 0.08 0.24 ± 0.06	14.68 ± 4.73 14.68 ± 4.73	0.25 ± 0.06 0.25 ± 0.06	0.13 ± 0.03 0.34 ± 0.42	3.53	68	1.97 ± 0.0
			23	0.23 ± 0.06 0.26 ± 0.01	0.34 ± 0.42 0.18 ± 0.02	4.9	78	1.97 ± 0.0
Cypripedium acaule	1.53 ± 0.06	0.21 ± 0.04			0.18±0.02	4.9	10	
Cypropedium calceolus	0.94 ± 0.29	0.2 ± 0.03	8.73 ± 4.98	1.61 ± 0.69				
Cypripedium candidum	0.88 ± 0.07							
Cypripedium parviflorum var. pubescens†	1.19 ± 0.17	0.2 1 0.04	01.16.1.1.45	0.02 1.0.02	0 15 1 0 02	2.8	87	
Cypripedium reginae	1.16 ± 0.26	0.3 ± 0.04	21.16 ± 1.45	0.23 ± 0.03	0.15 ± 0.03			
Dactylorchis	0.78 ± 0.15	0.28 ± 0.08	14.96 ± 6.27	0.26 ± 0.04	0.78 ± 0.02	4.18 ± 1.63	71.02 ± 15.48	
Dactylorhiza	A 44 1 A A4	0.00 1.0.01	0.000	0.23 ± 0.04	0.15 ± 0.03	0.45	93	
Dendrobium	0.66 ± 0.86	0.22 ± 0.34	0.53 ± 0.04	0.29 ± 0.34	0.19 ± 0.27			
Dendrobium nobile	0.33 ± 0.09	0.08 ± 0.02						
Tribe Diseae	0.43	0.16						
Disa	0.31 ± 0.3	0.26 ± 0.18						
Disa, Group 1	0.32	0.14						
Disa, Group 2	1.09	0.52						
Tribe Epidendreae	0.18-3.85							
Epidendrum	1.31 ± 1.41							6
Epipactis	1.16 ± 0.38	0.25 ± 0.07	21.94 ± 8.14	0.19 ± 0.01	0.15 ± 0.05	1.46	89.62 ± 2.6	
Epipactis atrorubens	0.9 ± 0.06	0.29 ± 0.02	19.28 ± 3.28				85	
Epipactis gigantea	0.17 ± 0.17	0.28 ± 0.13	38.17 ± 2.71				88	
Epipactis helleborine‡	1.15 ± 0.15	0.25 ± 0.02	18.23 ± 4.34				92	
Epipactis latifolia	1.19 ± 0.18	0.21 ± 0.02	14 ± 1.1				90	
Epipactis palustris	1.38 ± 0.42	0.26 ± 0.08	25.81 ± 5.25	0.22 ± 0.05	0.4 ± 0.02		92	
Epipactis rubiginosa§	and Barris and Arcal area	and a second		and an an a state	ande stat ere versiker:		90	
Epipogium	0.34 ± 0.21							

Galeola	1.25 ± 0.97	0.79 ± 0.22	434				18.33 ± 5.13	
Geodorum	1.3 ± 0.49		41000	0.25 6.5.5				
Goodyera	0.98 ± 0.37	0.12 ± 0.03	4 ± 1.39	0.2 ± 0.05	0.09 ± 0.02	0.78 ± 14	71.42 ± 9.07	
Goodyera oblongifolia	0.96 ± 0.07	0.13 ± 0.02	4.1 ± 1.3		10.000 0.000		77	
Goodyera pubescens	2.5 ± 1.9	0.1 ± 0.01	2.5 ± 0.5	0.22 ± 0.08	0.25 ± 0.35	0.73 ± 0.07	63.04 ± 5.13	
Goodyera repens	0.73 ± 0.23	0.14 ± 0.03	4.15 ± 1.3	0.19 ± 0.04	0.09 ± 0.02	0.72	82	
Gymnadenia	0.62 ± 0.14						98	8
Habenaria	0.84 ± 0.62	0.26 ± 0.16	3.83	0.26 ± 0.16	0.16 ± 0.09	2.02	47	
Hataeria	1.47 ± 0.17	0.09 ± 0.01						
Lecanorchis	3.38 ± 0.67	0.13 ± 0.03	15.07 ± 2.88					
Limodorum abortivum	0.78 ± 0.63	0.29 ± 0.23	81	0.04	0.02			5.7
Liparis	0.49 ± 0.26	0.15 ± 0.02	3.43 ± 3.1	0.15 ± 0.07	0.11 ± 0.05	0.86 ± 0.87	67.99 ± 32.49	
Listera cordata	0.69 ± 0.17	0.19 ± 0.07						
Listera ovata	0.77 ± 0.16	0.22 ± 0.01	9.7 ± 2.46					
Malaxis	0.39 ± 0.1							
Microtatorchis	0.48 ± 0.10							
Monadenia	0.28 ± 0.08	0.13 ± 0.02						
Neottia	0.88 ± 0.15	0.28 ± 0.09	21.68 ± 5.99					
Neottia nidus-avis	0.90 ± 0.15	0.27 ± 0.09	22.67 ± 5.41					
Nervillia	0.29 ± 0.07							
Neuwedia	0.66 ± 0.25	0.26 ± 0.07	38					
Nigritella	0.36 ± 0.03	0.19 ± 0.02	3.67	0.18 ± 0.01	0.12 ± 0.02			
Oberonia	0.18 ± 0.03							
Odontoglossum	0.42 ± 0.05	0.1 ± 0.02	1.14 ± 0.40	0.24 ± 0.11	0.09 ± 0.01	0.6	48	
Oncidium	0.42 ± 0.22	0.10 ± 0.05	1.83 ± 2.35					
Ophrys	0.50 ± 0.11	0.15 ± 0.04	4.58 ± 2.58	0.16 ± 0.05	0.10 ± 0.03			
Orchis	0.49 ± 0.17	0.17 ± 0.06	3.93 ± 3.24	0.18 ± 0.05	0.12 ± 0.04	1.22 ± 0.77	43.01 ± 35.16	
Orchis coriophora	0.46 ± 0.06	0.18 ± 0.05	4.17 ± 2.7	0.2 ± 0.02	0.14 ± 0.2	1.11	77	
Orchis mascula	0.39 ± 0.13	0.18 ± 0.03	6.43 ± 3.98	0.21 ± 0.1	0.14 ± 0.01	1.64	71	
Orchis morio	0.56 ± 0.16	0.18 ± 0.05	5.73 ± 4.52	0.18 ± 0.04	0.12 ± 0.03	1.35 ± 0.13	76.05 ± 12.87	
Orchis purpurea	0.38 ± 0.04	0.15 ± 0.07	2.65 ± 1.69	0.15 ± 0.03	0.09 ± 0.03	0.78 ± 0.34	and the second second second	
Orchis saccata	0.52 ± 0.01	0.13 ± 0.05	2.9 ± 1.5	0.14 ± 0.03	0.11 ± 0.01	1.72	58	
Orchis simia	0.38 ± 0.09	0.15 ± 0.08	4.2	0.16 ± 0.03	0.11 ± 0.03	1.55	73	
Paphiopedilum	0.7 ± 0.35	0.15 ± 0.04	4.36 ± 2.06	0.23 ± 0.05	0.124 ± 0.03	1.78 ± 0.66	62.38 ± 21.08	
Paphiopedilum barbatum	0.6 ± 0.43	0.10 ± 0.01	1.6 ± 1.21	0.20 1 0100	01121 ± 0100	1110 7 0100		
Paphiopedilum insigne	0.0 ± 0.15	0.17 ± 0.05	1.0 ± 1.21					
Phaius	0.96 ± 0.18	0.117 ± 0.000						
Phalaenopsis	0.35 ± 0.05	0.08 ± 0.01	0.59 ± 0.34	0.20 ± 0.05	0.08 ± 0.01		53	2.91
Phreatia	0.15 ± 0.03	0.00 1 0.01	0.07 1 0.01	0.20 1 0.00	0.00 ± 0.01			
Piperia	0.15 ± 0.05 0.54 ± 0.1	0.18 ± 0.04	4.69 ± 1.84	0.18 ± 0.03	0.11 ± 0.03	0.53 ± 0.66	75 ± 10.33	
Platanthera	0.54 ± 0.10 0.64 ± 0.20	0.25 ± 0.04	13.13 ± 4.54	0.13 ± 0.05 0.25 ± 0.05	0.16 ± 0.03	2.38 ± 0.6	81.21 ± 3.27	
Pogonia	1.04 ± 0.20 1.04 ± 0.27	0.18 ± 0.01	8.73 ± 2.11	0.18 ± 0.09	0.10 ± 0.03 0.1 ± 0.04	4.88	16	
Ponerorchis	1.04 ± 0.27 0.39 ± 0.02	0.18 ± 0.01 0.19 ± 0.01	3.57 ± 0.26	0.13 ± 0.09 0.21 ± 0.04	0.16 ± 0.04	1.00	10	
Pterostylis	0.39 ± 0.02 0.66 ± 0.33	0.17 _0.01	5.57 ± 0.20	0.21 10.04	0.10 1 0.04			
Sarcochilus	0.00 ± 0.03 0.39 ± 0.09							
Sarcochilus	0.39±0.09							

	Dimensions	Dimensions							
	Seed			Embryo			Free air		
Orchid	Length (mm)	Width (mm)	Volume (mm ³)	Length (mm)	Width (mm)	Volume (mm ³)	space in testa (%)	Weight (µg)	
Satyrium	0.39 ± 0.06	0.14 ± 0.02	2.64	0.21 ± 0.08	0.09 ± 0.03				
Schizochilus	0.28 ± 0.05	0.22 ± 0.04							
Spathoglottis	10.09 ± 0.21	0.30 ± 0.04	22.13 ± 9.95	0.26 ± 0.06	0.14 ± 0.03	5.69	72		
Spiranthes	0.51 ± 0.16	0.13 ± 0.03	3.1 ± 3.78	0.18 ± 0.05	0.10 ± 0.05		69		
Stanhopea	0.39 ± 0.25	0.28 ± 0.17		0.17 ± 0.13	0.09 ± 0.06				
Taeniophyllum	0.34 ± 0.11								
Thelymitra	0.31 ± 0.06								
Vanda	0.31 ± 0.13								
Vanilla	0.25 ± 0.08	0.22 ± 0.06	2.87 ± 2.71						
Zeuxine	1.23 ± 0.52	0.1 ± 0.01							

There are few published systematic, statistical and/or morphometric studies of orchid seeds. Information of the kind presented in this table must be compiled from numerous sources that: vary in accuracy and reliability; report on seeds from natural populations or plants that were apparently cultivated under different and usually not fully described conditions; probably refer to seeds and fruits in various stages of development and maturation; span more than 100 years; and are in many languages. A number of the values were obtained by measuring illustrations (drawings, light photographs and SEM micrographs). Even after a wide literature search only partial information is available for most genera and species.

To allow for useful statistical parameters (mean \pm SD, as shown), calculations were limited to genera and species for which there were at least three data points. Averages based on one or two data points, and therefore without SD, are included in some instances to provide a more complete picture. Percent free air space in the testa was calculated as: ((seed volume – embryo volume)/seed volume) × 100. Seeds were assumed to be two cones joined at their bases and their volume was estimated as 2((width/2)² × (length/2) × 1.047). Embryos were assumed to be prolate spheroids and their volume was approximated as $\frac{4}{3}\pi ab^2$ (a = length/2; b = width/2). Measurements or calculations (i.e. volume) are inaccurate or wrongly calculated in several previous reports (for example when orchid embryos were treated as oblate spheroids by Stoutamire (1981)). When errors were obvious the data were not used to make calculations. Further, the limited amount of available information precluded statistical calculations that could reflect the frequency and distributions of dimensions (for example, determine if certain seed length(s) are predominant, common or rare within a genus); that such differences do occur is obvious from the *Disa* entries. Calculations that were possible with the available data imply equal and uniform distribution of dimensions within genera and species.

Orchid nomenclature and taxonomy are very fluid – nomenclatural and classification changes of taxa are common. Since this table contains many orchid names, a determination of currently 'valid' taxonomic concepts would require a nomenclatural study of the entire family, and thus all names used here are the ones that appear in the original sources, except as indicated. The sources are: Link, 1839–1842; Beer, 1863; Beccari, 1877; Anonymous, 1879; Curtiss, 1893; von Marilaun, 1896; Bernard, 1909, 1937; Schlechter, 1915; Knudson, 1921, 1922; Ames, 1922a, b. 1948; Fleischer, 1929, 1930; Linder, 1930; Quednow, 1930; Curtis, 1932; Francini, 1932; Gratiot, 1934; Burgeff, 1936; Carlson, 1936; Ziegenspeck, 1936; Hoene, 1945; Davis, 1946; Maheshwari & Narayanaswami, 1952; Poddubnaya-Arnoldi & Selezneva, 1953, 1957a,b; Ito, 1955; Scott & Arditti, 1959; Leuschner, 1960; Kupper & Linsenmaier, 1961; Zimmerman & Dougoud, 1961; Hey, 1962; Mitsuura *et al.*, 1962; Nakamura, 1962, 1964; Hamada & Nakamura, 1963; Stoutamire, 1964a,b, 1983; Rauh *et al.*, 1965; Hey & Hey, 1966; Zinger & Poddubnaya-Ardnoldi, 1966; Clifford & Smith, 1969; de Vogel, 1969; Fuller, 1969a,b, 1970a, 1972, 1981; Veyret, 1969; Voth, 1959, 1963, 1964, 1969; Fuller, 1969, 1970b; Vanseveren-Van Espen, 1971; Wildhaber, 1969, 1970, 1972, 1974; Harvais, 1973; Leuchs, 1973; Barthlott, 1974; Harley, 1975; Haas, 1977; Hallé, 1977; Chua & Rao, 1978; Hasegawa *et al.*, 1978; Jordão *et al.*, 1988; Arditti *et al.*, 1979, 1980; Kim *et al.*, 1979; Mohana *et al.*, 1979; Healey *et al.*, 1980; Sheviak & Bye, 1980; Dafni, 1981; Barthlott & Ziegler, 1981; Lucke, 1981, 1982a,b, 1983, 1984a,b; Arditti, 1992; Karasawa & Saito, 1982; Gandawijaja & Arditti, 1983; Tohda, 1983, 1985, 1986; Cherevchenko & Kushnir, 1986; Chase & Pippen, 1988, 1990; Singh, 1981; Sood, 1989; Benzing & Clements, 1991; Petersson, 1991; Stewart, 1992; Kurzweil, 1994; Dressler, 1993; Proctor & Harder, 1994; Raghavan & Goh, 1984; Shoushtari *et al.*, 1994; Wagner & Hansel, 1994; Zettler & McInnis, 1994; Mo

*This is a monotypic genus.

†Cypripedium pubescens is another name for this *Cypripedium* species.

[†]This species is also known as *Epipactis helleborine* var. *helleborine*.

§Another name for this species is Epipactis atrorubens var. atrorubens.

||It is not uncommon to see this name spelled as Phajus.

that weigh not much more than those of *Paphio-pedilum* but have considerably less air space float for much shorter periods (Fig. 7b). Also, seeds of *Cephalanthera damasonium* (*Cephalanthera grandi-flora*) and *Cypripedium calceolus* (which are of equal weight, but have a smaller proportion of free air space than *Cymbidium*) float for an even shorter time (Fig. 7b).

Within the genus Paphiopedilum (Ando & Tomiyama, 1990), the percentage of air space seems to increase more in proportion to seed length (Fig. 8a,c) than width (Fig. 8b). As a consequence the longer seeds of subgenus Sigmatopetalum (some taxonomists consider this group to be section Barbata in the subfamily Paphiopedilum) have larger air spaces (Fig. 8a-c) and float for longer periods (Table 3; Karasawa & Saito, 1982). For example, floatation time for seeds of a hybrid with Paphiopedilum callosum (seed length, 1.09 mm; width, 0.15; air space, 83 %) as the seed parent is 9.6 s (to fall 1.5 m). By contrast, seeds produced by a cross with Paphiopedilum insigne (length, 0.48 mm; width, 0.17 mm; air space, 27 %) as the seed parent floated for only 4.2 s (Burgeff, 1936; Karasawa & Saito, 1982).

Floatation time is inversely proportional to the seed weight : percent air space ratio (Fig. 7c). This ratio becomes smaller as percent air space (the divisor) becomes larger (i.e. mass per unit of air space or a property which can be equated to density decreases). Altogether it is clear that weight probably plays a less important role in floatation time than percent (i.e. free) air space inside the seeds and seed volume.

To measure floatation time, Burgeff dropped seeds through a glass cylinder (1.5 m long, 40 mm diameter) which was placed over a 50 mm (presumably diameter) white or black glass plate (Burgeff, 1936). He timed the fall with a stop watch. His measurements are still the only ones available and because of that they are referred to often. Burgeff was generally a careful worker and a good technician, but not necessarily a good theoretician (as indicated by his Orcheomyces classification for mycorrhizal fungi isolated from orchids (Burgeff, 1909, 1911, 1932, 1936) and his failure to develop asymbiotic germination of orchid seeds). Lewis Knudson (1884-1958) of Cornell University developed the asymbiotic method for orchid seed germination in his very first reported experiment with orchids (see Arditti, 1990), when Burgeff had been working on seed germination and mycorrhizae of orchids for c. 15 yr.

There is no doubt that Burgeff's measurements are accurate as they stand, but they are not representative of what may happen under natural conditions because the air in his glass tube and laboratory was probably still and the seeds simply fell down due to gravity. Under natural conditions air is seldom still. Seeds which are released into the

air may float for a while and start to descend, but they can be (and most probably are) repeatedly blown in all directions by air turbulence and winds of varying intensities. Seeds with larger air spaces and longer vertical floatation times in Burgeff's tube may be affected differently by wind and turbulence than those that are smaller, less 'dense' and/or drop faster. As a result, floatation times of seeds in their natural habitats are undoubtedly different and more varied than in the laboratory and their trajectory is almost certainly not vertically down in a nearly straight line as it must have been in Burgeff's cylinder. Consequently, his measurements and data should be viewed only as approximations and indications of time periods during which the seeds can be affected by air currents.

More realistic information and experimental results about the floatation and travel distance is available in two illuminating and interesting recent reports that describe well-designed and clever experiments (Carey, 1998; Murren & Ellison, 1998). In addition to the intrinsic value of each report, the two when taken together have an added value because one deals with a tropical epiphyte and the other with a European terrestrial species.

(b) Dispersal. Orchid seeds can be assumed to be very small and very light balloons. In fact, the seeds of the following, for example, even look like balloons: Epipogium nutans (Fig. 2d; current name Epipogium roseum); Epistephium parviflorum (Fig. 1b); Eulophia strepropetala (Fig. 1b); Gongora maculata var. pallida (Fig. 1b); Haematorchis altissima (Fig. 1b); Listera australis (Fig. 2d); Listera convallarioides (Fig. 2d); Neottia nidus-avis (Fig. 1a); Pleione formosana (Fig. 2d, now Pleione bulbocodioides); Schizochilus zeyheri (Fig. 2d); Stanhopea insignis (Fig. 1b); Stanhopea oculata (Fig. 1b, 2d); Stanhopea oculata (Fig. 1b, 2d); Stanhopea saccata (Fig. 2d); Stanhopea tigrina (Fig. 1a, 2d); Vanilla planifolia (Fig. 1b, 2d), and Zygopetalum intermedium (Fig. 2d).

One of the two recent reports (Murren & Ellison, 1998; A. E. Ellison, pers. comm.) describes well thought out and interesting experiments with seeds of the Central American orchid Brasavola nodosa (also known as Dama de la Noche, or Lady of the Night, because it becomes fragrant in the evening). Their shape is similar to those of Brassavola cordata, Fig. 1c; they measure $640 \pm 40.6 \times 56 \pm 4.8 \mu m$, and weigh 1.6 µg. This report provides theoretical, physical and mathematical underpinnings for considerations of floatation, flight and dispersal of orchid seeds (Murren & Ellison, 1998; A. E. Ellison, pers. comm.). Starting with the assumption that wind dispersal of orchid seeds may be similar to a ballistic model proposed for Eucalyptus and described by the equation:

$$X_{\rm m} = h \frac{U_{\rm c}}{U_{\rm t}}$$
 Eqn 4

Table 2. Or	rchid seeds	: numbers	and	weights
-------------	-------------	-----------	-----	---------

Seed	ds (capsule ⁻¹ , mg ⁻¹ , fruit ⁻¹ ,
or µ	g seed ⁻¹

	or pg over				
Species	Weight	Number	Plant ⁻¹	Refs	
Acropera (Gongora)		3712577	4000000	Darwin, 1904	
Anacamptis pyramidalis		1935	35000	Tournay, 1960*	
Anguloa clowesii	1100 seeds mg ⁻¹			Cherevchenko & Kushnir, 1986	
	0.91 $\mu g \text{ seed}^{-1}$				
Anguloa ruckeri	2574	3932948		Thomale, 1954	
Anguloa ruckeri	2564 seeds mg ⁻¹			Cherevchenko & Kushnir, 1986	
Anguloa ruckeri	1524 mg fruit ⁻¹			Arditti, 1992	
Anguloa ruckeri	$0.39 \ \mu g \ seed^{-1}$			11 II I	
Calanthe	25-100 mg fruit ⁻¹			Cherevchenko & Kushnir, 1986	
Calanthe vestita	2000 seeds mg ⁻¹ 0.5 μg seed ⁻¹			Cherevchenko & Kushnir, 1986	
Calypso bulbosa Calypso bulbosa		6800		Wildhaber, 1974 Proctor & Harder, 1994	
Seeds with embryos					
Half pollinium		4909			
One pollinium		10980			
Two pollinia		15720			
Cattleya		Millions		Malguth, 1901, citing F. Cohn & E. Pfitzer.	
Cattleya		500000-6000000		Poddubnaya-Arnoldi & Selezneva, 1957	
Cattleya aurantiaca		256 000		Knudson, 1956; Lenz & Wimber, 1956; Tournay, 1960*	
Cattleya bowringiana	330 seeds mg^{-1} 3.03 mg seed ⁻¹			Arditti, 1992 Cherevchenko & Kushnir, 1986	
Cattleya gigas	5.05 mg seeu	500000-700000		Arditti, 1992	
Cattleya guttata	870 seeds mg ⁻¹ 1.15 μg seed ⁻¹	688264.123680		Cherevchenko & Kushnir, 1986	
Cattleya labiata		2000000-3000000 (929000 viable)		Withner, 1959	
Cattleya labiata Cattleya labiata	400 seeds mg ⁻¹ 2323 fruit ⁻¹			Cherevchenko & Kushnir, 1986 Arditti, 1992	
Cattleya labiata	2.5 μ g seed ⁻¹			Andrea, 1772	
Cephalanthera		6020	24000	Tournay, 1960*	
damasonium†				÷.1	
Cephalanthera grandiflora†		6020	24080	Darwin, 1904; Lenz & Wimber, 1959	
Coeloglossum viride		1330	12000	Lenz & Wimber, 1959; Tournay 1960*	
Cycnoches chlorochilon	3770000 fruit ⁻¹	3770000	3770000	Ames, 1946; Lenz & Wimber, 1959; Tournay, 1960*; estimate by Rolfe, 1912	
Cycnoches chlorochilon		3751000		Bechtel <i>et al.</i> , 1986	
Cycnoches chlorochilon		3932948		11 II I	
Cycnoches chlorochilon	14400 seeds mg^{-1}	4000000		Arditti, 1992	
Cycnoches chlorochilon	3.6 µg seed ⁻¹	4000000		0	
Cymbidium	300-700 mg fruit-1			Cherevchenko & Kushnir, 1986	
Cymbidium		1 500 000		Arditti, 1992	
<i>Cymbidium</i> hybrid	730 seeds mg^{-1} 1.37 mg seed ⁻¹			Cherevchenko & Kushnir, 1986	
Cymbidium traceyanum		850000		Weiss, 1916; Arditti, 1982	
Cypripedium acaule		14000–54180 28160	28160	Tournay, 1960*	
Cypripedium acaule	526 mg ⁻¹	54180		Stoutamire, 1964b	
Cypripedium acaule	108 mg fruit ⁻¹			Arditti, 1992	
Cypripedium acaule	2 $\mu g \text{ seed}^{-1}$				
Cypripedium calceolus Seeds with embryos		1425		Proctor & Harder, 1994	
Natural pollination		1435			
Quarter anther Half anther		1867 3660			
One anther		3158			
Dactylorhiza		6200	186300	Tournay, 1960*	

Dactylorhiza (Dactylorchis) fuchsii (Druce) Soó Dendrobium U Dendrobium antennatum 2 5 Dendrobium antennatum 1 5 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 0 Epidendrum (radicans) 3 ibaguense 3 Epidendrum (radicans) 1 ibaguense 5 Epipactis, average for several species Eulophia horsfallii 5 Galeola altissima 1 3 Galeola septentrionalis 3	Weight			
(Dactylorchis) fuchsii (Druce) SoóDendrobiumUDendrobium antennatum25Dendrobium antennatum15Dendrobium antennatum11Dendrobium phalaenopsis11Dendrobium phalaenopsis11Dendrobium phalaenopsis11Dendrobium phalaenopsis32Dendrobium phalaenopsis11Dendrobium phalaenopsis11Dendrobium phalaenopsis32Epidendrum (radicans)11Baguense52Epipactis, average for several species11Galeola altissima13Galeola septentrionalis33Galeola septentrionalis32Gongora sp.2		Number	Plant ⁻¹	Refs
(Druce) SoóDendrobiumUDendrobium antennatum25Dendrobium antennatum15Dendrobium phalaenopsis11Dendrobium phalaenopsis11Dendrobium phalaenopsis11Dendrobium phalaenopsis31Dendrobium phalaenopsis31Dendrobium phalaenopsis31Dendrobium phalaenopsis11Dendrobium phalaenopsis31Dendrobium phalaenopsis11Dendrobium phalaenopsis31Dendrobium phalaenopsis11Dendrobium phal		3294	> 56 000	Tournay, 1960*
Dendrobium U Dendrobium antennatum 2 5 Dendrobium antennatum 1 5 Dendrobium antennatum 1 5 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 3 1 Dendrobium (radicans) 3 1 Baguense 5 2 Epipactis, average for several species 1 1 Galeola altissima 1 3 Galeola septentrionalis 3 2 Gongora sp. 3				
Dendrobium antennatum 2 5 Dendrobium antennatum 1 5 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 0 Epidendrum (radicans) 3 ibaguense 3 Epidendrum (radicans) 1 ibaguense 5 Epipactis, average for several species Eulophia horsfallii 5 Galeola altissima 1 Galeola septentrionalis 3 Gongora sp.				
5 Dendrobium antennatum 1 5 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 0 Epidendrum (radicans) 3 ibaguense 3 Epidendrum (radicans) 1 ibaguense 5 Epipactis, average for several species 5 Eulophia horsfallii 5 Galeola altissima 3 Galeola septentrionalis 3 Gongora sp.	Jp to 500 mg fruit ⁻¹			Arditti, 1992
5 Dendrobium phalaenopsis 1 1 Dendrobium phalaenopsis 1 0 Epidendrum (radicans) 3 ibaguense 3. Epipactis, average for several species Eulophia horsfallii 5. Galeola altissima 1 Galeola septentrionalis 3 Gongora sp.	200 seeds mg ⁻¹ μg seed ⁻¹			Pfitzer, 1882
1 Dendrobium phalaenopsis 1 0 Epidendrum (radicans) 3 ibaguense 3 Epidendrum (radicans) 1 ibaguense 5 Epipactis, average for several species Eulophia horsfallii 5 Galeola altissima 1 Galeola septentrionalis 3 Gongora sp.	.65 mg seed ⁻¹			Cherevchenko & Kushnir, 1986
0 Epidendrum (radicans) 3 ibaguense 3 Epidendrum (radicans) 14 ibaguense 5 Epipactis, average for several species Eulophia horsfallii 5 Galeola altissima 14 Galeola septentrionalis 33 Gongora sp.	mg seed ⁻¹			0
ibaguense 3. Epidendrum (radicans) 10 ibaguense 5. Epipactis, average for several species Eulophia horsfallii 5. Galeola altissima 11 Galeola septentrionalis 33 Gongora sp.	.89 mg seed ⁻¹			
ibaguense 5. Epipactis, average for several species Eulophia horsfallii 5. Galeola altissima 11 Galeola septentrionalis 33 Gongora sp.	20 seeds mg^{-1} 1.13 mg seed ⁻¹			0
several species Eulophia horsfallii 5. Galeola altissima 1 Galeola septentrionalis 3. Gongora sp.	67 seeds mg ⁻¹ .99 μ g seed ⁻¹			0
Eulophia horsfallii 5. 1. Galeola altissima 1 Galeola septentrionalis 3. 2. Gongora sp.		4500		Wildhaber, 1970
Galeola septentrionalis 3 Gongora sp.	58 seeds mg ⁻¹ .79 mg seed ⁻¹	204454		Tournay, 1960*
Galeola septentrionalis 3: 2: Gongora sp.	7 mg seed ⁻¹ 06 mg fruit ⁻¹	18000		Hamada & Nakamura, 1963; Arditti, 1992
Gongora sp.	84 mg fruit ⁻¹ 24 μg seed ⁻¹	16000		Hamada, 1939; Nakamura, 1964 Rasmussen, 1995; Arditti, 1992
Gongora sp.	10,000	317250		Darwin 1904; Withner, 1959; Tournay, 1960*
			740000001	Tournay, 1960*
Goodyera repens		1700	25 500	Tournay, 1960*
Goodyera repens				Rasmussen, 1995
Gymnadenia conopsea 2	l μg seed ⁻¹	2000	32000	Tournay, 1960*
Himantoglossum		1200	28800	Ziegenspeck, 1936
	60 seeds mg ⁻¹ .32 mg seed ⁻¹			Cherevchenko & Kushnir, 1986
Leporella fimbriata		5000		Peakall <i>et al.</i> , 1987; Peakall & James, 1989
Limodorum abortivum 5	.7 mg			Rasmussen, 1995
Listera cordata		376	2860	Stoutamire, 1964
Listera ovata		544	21000	Tournay, 1960*
Malaxis monophyllos		1572	19490	Stoutamire, 1964b
Maxillaria		1756440	10538640	Darwin, 1904
Maxillaria Maxillaria		1756440	10538640 < 10000000	Tournay, 1960* Poddubnaya-Arnoldi & Selezneva, 1957
Oeceoclades maculata				González-Díaz & Ackerman, 1988
Cultivated plants				1222224
Control flowers		2011 seed fruit ⁻¹ 12067 per six fruits		
Self-pollinated flowers		69% fertile seeds 893 seeds fruit ⁻¹ 30346 per thirty- four fruits		
Cross-pollinated flowers		92% fertile seeds 896 seeds fruit ⁻¹ 34034 per thirty- eight fruits		
5	50.7 mg fruit ⁻¹	signi nuns		
2		88 % fertile seeds		

390 TANSLEY J. Arditti and A. K. A. Ghani

Table 2 (cont.)

	Seeds (capsule ^{-1} , or μ g seed ^{-1})	mg ⁻¹ , fruit ⁻¹ ,		
Species Weight		Number	Plant ⁻¹	Refs
Self-pollinated, one pollinium		619 seeds fruit ⁻¹		González-Díaz & Ackerman, 1988
		6808 per eleven		
	200 - 10 - 10 - 10 - 10 - 10 - 10 - 10 -	fruits		
	23 mg fruit ⁻¹	000/ 0 11		
Calf walling and true		93 % fertile seeds 623 seeds fruit ⁻¹		
Self-pollinated, two pollinia		025 seeds fruit		
ponnna		6850 per eleven		
		fruits		
	46.2 mg fruit ⁻¹			
		92 % fertile seeds		
Flowers on plant under		515 seeds fruit ⁻¹		
artificial rain		5666 man alawan		
		5666 per eleven fruits ⁻¹		
	27.8 mg fruit ⁻¹	Iruits		
		99% fertile seeds		
Plants in the field				
Control 1		870 seeds fruit ⁻¹		
		24356 per twenty-		
	65.8 mg fruit ⁻¹	eight fruits ⁻¹		
	05.8 mg mun	97% fertile seeds		
Control 2		511 seeds fruit ^{-1}		
		28123 per fifty-five		
		fruits		
	60.7 mg fruit ⁻¹			
Insect exclusion		96 % fertile seeds 498 seed fruit ⁻¹		
Theet exclusion		4,483 per nine		
		fruits ⁻¹		
	66.3 mg fruit ⁻¹			
		98% fertile seeds		
Oncidium		6,200	10.000	m 10/0#
Ophrys apifera Orchids	÷	10,000 'Thousands'	40000	Tournay, 1960* Rasmussen, 1995
Orchids, epiphytic		< 6 000 000		Poddubnaya-Arnoldi &
oreinus, epipilytie		0000000		Selezneva, 1957, 1964, 1976
Orchids, terrestrial		< 10000		0 · · · · · ·
Orchis maculata§		6200	186300	Darwin, 1904; Lenz & Wimber,
01.1.5			5(000	1959
Orchis maculata§ Orchis morio		> 40 000	56000 32000	Lenz & Wimber, 1959 Tournay, 1960*
Platanthera chlorantha		25000	13750	in in the second
Rhizanthella gardenri		20-50	10100	George & Cooke, 1981
Stanhopea		Millions		Malguth, 1901 citing F. Cohn &
				E. Pfitzer
Stanhopea oculata	333 seeds mg^{-1}			Cherevchenko & Kushnir, 1986
Stanhopea tinning	3 mg seed ⁻¹ 600 seeds mg ⁻¹			
Stanhopea tigrina Stanhopea tigrina	$1.67 \text{ mg seed}^{-1}$			
Subtropical Cymbidium	the the obed			
Hybrid	$1.37 \ \mu g \ seed^{-1}$			
Temperate orchid sp.	9.84±10.22 μg			Calculated from data in this
T	seed ⁻¹			table. Range: 24–0.5 µg seed ⁻¹ ¶
Tropical orchid spp. Tropical orchids	$< 1 \ \mu g^{**}$			Rasmussen, 1995 Calculated from data in this
ropical orenids	2.56±1.76 μg seed ⁻¹			table. Range: 0.39–5.99
	0004			seed ⁻¹ ††

Table note: see facing page.

 $(X_m, approximate mean distance a seed will disperse;$ h, height of release; U_e , average wind velocity between point of release and ground; and U_{t} , terminal velocity (in this case $U_t = 0.157$ m s⁻¹). They (Murren & Ellison, 1998) measured seed dispersal in the field, wind tunnel and the laboratory. Based on their experimental evidence they concluded that modal dispersal distance (i.e. floatation and movement or travel in air) of dust-like orchid seeds was predicted adequately by this ballistic model (Eqn 4) at low wind velocities and low release heights, but that it underestimated the increasing importance of turbulence at higher wind velocities and greater release heights (Murren & Ellison, 1998). On estimating the relative importance of one measure of turbulence, vertical mixing velocity (W^*) on $X_{\rm m}$ in wind tunnel experiments (i.e. floatation distance) they found that under high turbulence adding mixing velocity to the ballistic model overestimated modal seed dispersal distances (Murren & Ellison, 1998). The reasons for this can be found in the following equations:

$$\lambda = \overline{1 + \left(\frac{W^*}{2U_t}\right)^2 + \frac{W^*}{2U_t}}$$
 Eqn 5

$$\lambda \approx 1 + \frac{W^*}{2U_t} \approx l$$
 Eqn 6

 $(W^*$, rate at which a suspended particle would reach the ground by eddy diffusion in the absence of gravity (Murren & Ellison, 1998); λ , dimension parameter which includes the effects of W^* and U_t and represent the spread of seed around their major location). By contrast, Burgeff's seeds fell to the ground strictly due to gravity and were not affected by eddies.

In these formulas, the value of λ will increase approximately in proportion to one third the power of airspace and this should reduce terminal velocity, U_t (A. M. Ellison, pers. comm.). In general, according to Eqns 5 and 6, the consequence of larger air spaces in orchid seeds would be an increase in the effects of turbulence (because the seeds will float longer) on modal dispersal distance and increased variance. This means that increased air space may increase floatation time yet reduce the average travel distance of a seed, but the stray seed could go a lot further (A. M. Ellison, pers. comm.). Given the large number of seeds produced by orchids, the 'stray seed' may actually be a large number in any given year and even more over longer periods.

A somewhat different approach was used in another carefully designed and illuminating study of the spread of *Himantoglossum hircinum* (Fig. 1a) in the south of England (Carey, 1998). This study tested two dispersal (i.e. floatation and air travel) models for seeds of this species. They measure $130 \times 30 \,\mu\text{m}$, weigh $5 \times 10^{-9} \,\text{kg}$ (Carey, 1998) and are produced at 1200 fruit⁻¹ or 28800 plant⁻¹ (Ziegenspeck, 1935, 1936). One of these, a diffusion model, represents dispersal as a bivariate normal distribution including root-mean-square displacement (Carey, 1998):

$$Q_{(x)} = \frac{M}{\pi R^2} \exp\left(-\frac{x^2}{R^2}\right)$$
 Eqn 7

 $(Q_{(x)})$, deposition density of seeds at distance x from the parent; M, number of seeds produced by parent plant; R, root-mean-square displacement for seed dispersal). This equation takes into account the number of seeds produced by a plant (M), but not their volume and/or weight. It indicates that a large M can lead to higher $Q_{(x)}$ values over short distances and probably lower ones over longer ranges and distributions (i.e. floatation periods) similar to the ones defined by Eqns 5 and 6. For *Himantoglossum hircinum* this equation is less satisfactory that another one called the tilted plume model (Carey, 1998):

$$\begin{split} Q_{(\mathrm{x})} = & \left(\frac{MW_s}{H\bar{U}\Gamma(1+\beta)}\right) \left(\frac{H^2\bar{U}}{(1+\alpha)\bar{A}}\right)^{1+\beta} x^{-\beta-1} \\ & \exp - \left(\frac{H^2\bar{U}}{2(1+\alpha)\bar{A}_{\mathrm{x}}}\right) \quad \mathrm{Eqn} \ 8 \end{split}$$

 $(Q_{(x)})$, deposition density of seeds at distance x from the parent; M, number of seeds (many in orchids); W_s , settling velocity of the seed (orchid seeds with large air spaces drop more slowly and therefore should have a lower settling velocity); H, height at

Except as noted below the names used here are those in the original literature. In most cases there are not enough reports (at least three) for the calculation of statistical parameters (mean, SD).

^{*}Tournay (1960) contains information from several sources - hence different values for the same species.

[†]This species has three names and is a typical example of the unending taxonomic gyrations that plague orchid systematics. At one time it was known as *Cephalanthera pallens*. This name was reduced to synonymy under *Cephalanthera grandiflora* and *Cephalanthera damasonium*.

[†]The figure 74000000 was obtained by multiplying 200 (the number of capsules found on one plant) by 317250 (the number of seeds per capsule of another specimen of this orchid).

[§]Another name for this species is Dactylorhiza maculata.

PCymbidium was excluded from these calculations because not all species are fully tropical. If it is included the values change to $8.43 \pm 9.77 \ \mu g \ seed^{-1}$.

^{**}This figure is incorrect as is obvious from calculations based on data in this table. It appears in a book on the seeds of terrestrial orchids (Rasmussen, 1995), may therefore be accepted as being accurate and must be corrected.

 $[\]pm Cymbidium$ was excluded from these calculations because not all species are fully tropical. If it is included the values change to $2.49 \pm 1.73 \,\mu\text{g seed}^{-1}$.

Table 3. Weight, air space and floatation of some orchid seeds in air

Species	Floatation in air (s)	Weight (µg)	Air space (%)	Refs
Acanthephippium sylhetense	13.0	0.66	97.35†	Burgeff, 1936
Angraecum cv. Veitchii	9.0		0.70	10
$(Angraecum \ eburneum \times A. \ sesquipedale)$				
Anguloa ruckeri	13.0		0.39	.0.
Anguloa uniflora × A. ruckeri	10.0			2
Anguloa mean	11.5	0.39		
Calanthe sylvestris* $\times C$. cv. Darblayana	12.0			
Calanthe cv. Veitchii		0.5		Table 1; J. Arditti & A. K. A. Ghani, unpublished
Calanthe veratrifolia $\times C$. vestita	13.0			Burgeff, 1936
Calanthe mean	12.5	0.5	78.75†	
Cattleya aurantiaca	2.65	99.00	0.00200.0004	Shoushtari et al., 1994; Table 2
Cattleya elongata		4.1		n , , , , , , , , , , , , , , , , , , ,
Cattleya forbesii		3.5		τí α.
Cattleya guttata $\times C$. autumnalis	5.0	010		Burgeff, 1936
Cattleya guttata var. leopoldinae‡	5.15			Shoushtari et al., 1994
Cattleya harrisoniana	0110	5.71		0
Cattleya harrisoniae $\S \times C$. gaskelliana	5.2			Burgeff, 1936
Cattleya intermedia		2.33		Shoushtati et al., 1994
Cattleva labiata		2.5		Hager, 1954
Cattleya labiata $\times C$. gaskelliana	5.6	2.0		Burgeff, 1936
Cattleya lawrenceana	5.0	3.45		Shoushtari et al., 1994
Cattleya lobata		0.10	76.86	Schlechter, 1915
Cattleya mossiae		3.12	70.00	Shoushtari et al., 1994
Cattleya percivalliana $\times C$. aurea¶	4.3	0.12		Burgeff, 1936
Cattleya reinekiana	1.5	5.05		Shoushtari et al., 1994
Cattleya schroederae		6.76		
Cattleya trianae		0.70	61.85	Schlechter, 1915; Shoushtari et al., 1994
Cattleya velutina $\times C$. gigas	4.6		01.05	Burgeff, 1936
Cattleya mean	4.94 ± 0.51	3.92 ± 1.48	79.24†±4.24	bugen, 1950
Cephalanthera austinae	4.94 ± 0.51	3.74 1 1.40	90.14	Tables 1,2
Cephalanthera damasonium		2.00	78.63	Rasmussen, 1995
Cephalanthera damasonium		2.00	60	Tables 1,2
Cephalanthera grandiflora	6.0		75	Burgeff, 1936
Cephalanthera mean	6.0	2.00	75.94±12.43†	burgen, 1950
Corallorhiza innata	7.2	2.00	75.94 ± 12.43 83	11
	4.9		00	II
Cymbidium devonianum	4.9	0 77		
Cymbidium eburneum		3.77	70.26	Shoushtari et al., 1994
Cymbidium	2.0		79.26	Dunger († 1026
Cymbidium giganteum** $\times C$. cv. Doris	3.0			Burgeff, 1936

Cymbidium hybrid		1.36		Cherevchenko & Kushnir, 1986
Cymbidium insigne \times C. lowianum	4.6			Burgeff, 1936
Cymbidium madidum		4.46		Shoushtari et al., 1994
Cymbidium traceyanum \times C. insigne	5.0			Burgeff, 1936
Cymbidium mean	4.38 ± 0.93	3.20 ± 1.63	79.26†	
Cypripedium++		2.0		Table 2
Cypripedium acaule		1.9	78	Tables 1,2
Cypripedium calceolus L. ^{‡‡}	5.9		72.79	Mean from Tables 1,2
Cypripedium californicum			92	Table 1
$Cypripedium \ californicum \times C. \ acaule$			90	13
Cypripedium candidum			89	
Cypripedium guttatum	7.3			Burgeff, 1936
Cypripedium montanum			89	Table 1
Cypripedium parviflorum			93	11
Cypripedium passerinum ^{‡‡}			92	0.
Cypripedium pubescens ¹¹	8.1		94.5	Burgeff, 1936
$Cypripedium$ pubescens $\times C$. acaule			93	Table 1
Cypripedium reginae			87	G ^{aren e}
Cypripedium mean	7.1 ± 1.11	2.0	$88.21 \pm 6.8 \pm$	
Cyrtopodium punctatum	5.1	3.4	1.5.7.5.7.7.7.7 .5. 7.9.9.5.1	Burgeff, 1936; Shoushtari et al., 1994
Didymoplexis pallens	32.0	0.45		Burgeff, 1936
Epipactis atrorubens			85	н
Épipactis gigantea			88	Table 1
Epipactis helleborine			92	ii anna anna anna anna anna anna anna a
Epipactis latifolia	9.1		90	Burgeff, 1936; Tables 1,2
Epipactis palustris	7.5		92	
Epipactis rubiginosa	6.8		90	14
Epipactis mean	7.8 ± 0.96		89.5 + 2.67 +	
Eulophia	1.1.1. - 1 .1.1.1.1	2.73	2.10 ± 2121	Table 2
Eulophia × Phaius flavus	23.0			Burgeff, 1936
Eulophia mean	23.0	2.73		Table 2
Goodyera mean		2.00	71+	Tables 1,2
Grammatophyllum speciosum	7.8	2.48		Burgeff, 1936
Gymnadenia conopsea§§	4.8	8	981	
Laelia	1.1.1	3.05		Table 2
Laelia anceps \times L. cinnabarina	4.3	0.00		Burgeff, 1936
Laelia purpurata \times Cattleya citrina	4.5			in the second se
Laelia tenebrosa \times Cattleya mendelii	2.3			**
Laelia mean	3.7 ± 1.22	3.05		
Limodorum abortivum	3.7	5.7	98.68†	";Tables 1,2
Liparis mean	7.0	500 C	68†	, a moreor a jar

Species	Floatation in air (s)	Weight (µg)	Air space (%)	Refs
Odontoglossum mean	6.3	12	48†	
Orchis mean	5.8		66.87+	
Paphiopedilum++		3.00		Table 2
Paphiopedilum cv. Albertianum $\times P$. cv. Sallieri	4.1			Burgeff, 1936
Paphiopedilum argus	6.1			
Paphiopedilum callosum $\times P$. curtisii	9.6			0
Paphiopedilum charlesworthi	3.5			
Paphiopedilum curtisii	10.5		96	";Tables 1,2
Paphiopedilum insigne			74	Table 1
Paphiopedilum insigne var. sanderae $\times P$. concolor	4.2			Burgeff, 1936
$Paphiopedilum \ lawrenceanum \times P. \ glaucophyllum$	13			
Paphiopedilum spicerianum	5.0			0
Paphiopedilum cv. Superciliare $\ \ \times P$. bellatulum	11.0			n:
Paphiopedilum cv. Susan Tucker × P. parishii			99	Arditti et al., 1979
Paphiopedilum tonsum $\times P$. niveum	12.0			Burgeff, 1936
Paphiopedilum villosum $\times P$, tonsum	3.8			0.
Paphiopedilum mean	7.53 ± 3.7	3.00	89.67 ± 13.65 +	
Phaius flavus	29.0		1970 - 1971 - 19	
Phaius maculatus		3.53	76	Table 2
Phaius $\times P$. wallachii	16.0			Burgeff, 1936
Phaius mean	22.5	3.53	76†	
Phalaenopsis			68.42	Table 1
Phalaenopsis amabilis $\times P$. rosea			53.32	
Phalaenopsis aphrodite $\times P$. sanderiana	9.0			Burgeff, 1936
Phalaenopsis mannii		2.91		Shoushtari et al., 1994
Phalaenopsis sanderiana	9.4			Burgeff, 1936
Phalaenopsis mean	9.2	2.91	60.87+	un de carte
Platanthera bifolia	6.8		82	Burgeff, 1936; Tables 1,2
Platanthera chlorantha	6.6		82	n n n
Platanthera dilatata var. leucostachys			78	Healey et al., 1980
Platanthera hyperborea			77	H.
Platanthera integrilabia			57.55	Zettler & McInnis, 1994
Platanthera saccata			86	Healey et al., 1980
Platanthera mean	6.7		$77.09 \pm 10.10 \ddagger$	
Renanthera imshootiana	11.6			Burgeff, 1936
Schomburgkia undulata	6.6	0.3		
Sobralia macrantha	3.7	6.3		11.
Sobralla macranina	2.1	0.5		

Sobralia, mean	4.05	6.3		
Spathoglottis	7.8		72	Burgeff, 1936; Table 2
Stanhopea		2.95		Table 2
Stanhopea oculata		3.00	89.31	von Marilaun, 1896; Fleischer, 1929, 1930; Burgeff, 1936
Stanoppea oculata \times S. quandricornis	5.9			Burgeff, 1936
Stanhopea mean	5.9	2.95	89.31†	

Since information about weight, air space and floatation time for all taxa in this table is not available, data for related species or estimations for genera are included for comparison purposes. Mean and SD were calculated only when there were more than three values for any group. Given the small number of samples and the size of the Orchidaceae these numbers should be treated with caution. Names and their spelling used here are as in the original literature (except as indicated) because it is not possible to retroactively determine taxonomic identities.

Floatation in air (in seconds) was measured by H. Burgeff by dropping seeds in a glass tube (1.5 m long and 40 mm in diameter) and determining the time required for them to reach a white or black glass plate placed 50 mm below the top of the cylinder. First to reach the plate were seed clumps; they were followed by individual seeds; empty testae (i.e. chaff) were last. The error for descent times of 3–6 s was 0.1 s. For longer descent times (up to 30 s) the error was 5 s (Burgeff, 1936). A possible overall error in Burgeff's measurements may have been generated by the still air inside his cylinders. In nature there are always air swirls (even if sometimes very gentle ones) which could affect the floatation time of seeds. *Burgeff may have meant *Calanthe sylvatica*.

†These values and a number of others were calculated from data in Tables 1 and/or 2 and/or our unpublished data. Because of that they are should be viewed as approximations. Calculations like these were made in cases where floatation times were available but not weights and/or percent air space.

[‡]This is probably Laelia autumnalis.

§Burgeff probably meant C. harrisoniana. This species is also known as Cattleya loddigesii var. harrisoniana.

Another possible name for this orchid is Laelia lobata.

¶Yet another uncritical use of an orchid name by Burgeff; this is Cattleya dowiana var. aurea.

**Other names for this species are Cymbidium lowianum, Eulophia speciosa and Iridorchis gigantea.

++At one time Cypripedium included the genera that are presently known as Cypripedium and Paphiopedilum. When specific epithets and/or illustrations are not available in publications of that era it is often impossible to determine the identity of a 'Cypripedium'.

‡‡Some taxonomists consider Cypripedium calceolus, Cypripedium parviflorum and Cypripedium pubescens as separate species. Others reduce the status of the last two to varieties of the first.

§§It is not unusual to see Gymnadenia conopsea or Gymnadenia conopea in the literature.

|| ||Burgeff does indicate that this is a hybrid.

396 TANSLEY J. Arditti and A. K. A. Ghani

Table 4.	Floatation	of	some	orchid	seeds	in	water	and	air	
----------	------------	----	------	--------	-------	----	-------	-----	-----	--

			ime in air come wei				
		Water					
Species (when available also seed weight (μg) and free air space $({}^{0}_{0})$)		(d)					
	Air (s)	Dry	Wetted	Difference	Duration of floatation	n f/s	Refs
Acanthephippium sylhetense (0.66 mg, 97.35%)	13.0						Burgeff, 1936
Aceras anthropophora (Aceras 66.67%) Immediate wetting							Ziegenspeck, 1935
Wetting after 1 h						1	
Wetting after 24 h						2 0 2	
Sinking after 24 h						0	
Wetting after 60 h							
Sinking after 20 h						1	-
Aerides quenquevulnerum × Vanda tricolor	8.0						Burgeff, 1936
Amerorchis rotundifolia Anacamptys pyramialis Immediate wetting Wetting after 1 h						f	Stoutamire, 1981 Ziegenspeck, 1935
Wetting after 24 h Sinking after 24 h						0	
Wetting after 60 h						0	
Sinking after 20 h						0	0 1001
Angraecum eburneum Angraecum eburneum $\times A$. sesquipedale = A. cv. Veitchii (0.7 µg)	9.0					S	Stoutamire, 1981 Burgeff, 1936
Angraecum sesquipedale		0	0	0			1.1
Anguloa ruckeri (0.39 µg)	13.0	U	U	0			11
Anguloa uniflora × A. ruckeri Anguloa virginalis	10.0					f & s	Stoutamire, 1981
Anguloa mean	11.5	(0.39 µ	g)				
Ansellia africana		A	87			f	1.1
Aplectrum hyemale						f	7.1
Aretusa bulbosa						f	O.
Bletilla hyacinthina	3.5	2		2			Burgeff, 1936
Brassia verrucosa Brassia verrucosa × Oncidium	4.5 4.5						11
leucochilum						60	0
Brassavola nodosa						f & s	Stoutamire, 1981
Bulbophyllum macrobulbum Calanthe silvestris (this is probably $C.$ sylvatica) $\times C.$	12.0	1		1		1 & 5	Burgeff, 1936
cv. Darblayana (0.5 μ g) Calanthe veratrifolia × C. vestita	13.0	(0.5	70 740/				11
Calanthe mean	12.5	(0.5 µg	, 78.74%)			C.	0 1001
Calopogon pallidus Calopogon tuberosus						f f	Stoutamire, 1981
						f	1 0
Calypso bulbosa Catasetum macrocarpum			1		1	1	Burgeff, 1936
Cattleya		1–4 to 7–8			1		Fleischer, 1929
Cattleya aurantiaca		1. 1.	- 1 0			s	Stoutamire, 1981
Cattleya citrina		9	1	8		6	Burgeff, 1936
Cattleya guttata \times C. autumnalis	5.0		10 BB 10				11
Cattleya guttata var. leopoldinae	5.15						Shoushtari et al., 1994

Floatation time in air or water, or Time for dry seeds to become wet and continue to float Water (d) Species (when available Duration also seed weight (µg) Air and free air space (%)) Wetted Difference Refs (s) Dry of floatation f/s Cattleya harrisoniae \times C. gaskelliana 5.2 Burgeff, 1936 Cattleya labiata × C. gaskelliana 5.6 11 4.3 Cattleya percivalliana $\times C$. aurea (this is probably C. dowiana aurea) 11 Cattleya velutina \times C. gigas 4.6 Cattleya mean 4.94 (3.94 µg, 79.24%) Fleischer, 1929 Cephalanthera grandiflora 28 5 Cephalanthera grandiflora 6.0 28 5 23 Burgeff, 1936 23 5 Cephalanthera pallens Ziegenspeck, 1936 6.00 (2.00 µg, 76%) Cephalanthera mean Chysis bractescens 1 Burgeff, 1936 1 0 Cleistes divaricata f Stoutamire, 1981 Cochlioda neotzliana × Odontoglossum 6.0 Burgeff, 1936 crispum 19 8 Coeloglossum Ziegenspeck, 1936 8 19 Coeloglossum viride Fleischer, 1929 27 8 19 Burgeff, 1936 Coeloglossum viride Colegolossum viride 19 8 Ziegenspeck, 1936 Coeloglossum viride f Stoutamire, 1981 f & s Coelogyne cristata 1 Coelogyne huettneriana Burgeff, 1936 1 Coelogyne macrocarpum 1 Corallorhiza 60 6-8 Ziegenspeck, 1936 19 Corallorhiza 8 Corallorhiza innata 60 7 Fleischer, 1929 7 60 Corallorhiza innata 7.2 60 Burgeff, 1936 Corallorhiza maculata f Stoutamire, 1981 Corallorhiza odontorhiza f 11 11 Corallorhiza striata f 11 Corallorhiza trifidida f Corvanthes macrantha 11.0 30 Burgeff, 1936 Cymbidium devonianum 4.9 11 3 3 Cymbidium giganteum \times C. cv. Doris 3.0 11 Cymbidium insigne $\times C$. lowianum 4.6 3 11 1 Cymbidium lancifolium 11 Cymbidium traceyanum \times C. insigne 5.0 Cymbidium mean 4.38 (3.2 µg, 79.26%) Cypripedium acaule f Stoutamire, 1981 Cypripedium arietinum f 7 Cypripedium calceolus (72.79%) 17 Fleischer, 1929 7 Burgeff, 1936 Cypripedium calceolus 5.9 25 17 7 17 Ziegenspeck, 1936 Cypripedium calceolus Cypripedium calceolus var. pubescens f Stoutamire, 1981 Cypripedium californicum f 11 11 Cypripedium candidum f 11 Cypripedium fasciculatum f Cypripedium guttatum 60 Burgeff, 1936 7.3 60 6 Stoutamire, 1981 Cypripedium guttatum f Cypripedium montanum f 11 Cypripedium passerinum f Cypripedium pubescens (94.5%) Burgeff, 1936 8.1 f Cypripedium reginae Stoutamire, 1981 Cypripedium mean 7.1±1.11 (2 μg, 88.21±6.8) Cyrtopodium punctatum (3.4 µg) 5.1 1 Burgeff, 1936 Cyrtopodium punctatum f Stoutamire, 1981

		ation t s to be				
		Wate	r			
		(d)				
Species (when available also seed weight (µg)	Air	17-1-1-1			Duration	
and free air space (%))	(s)	Dry	Wetted	Difference	of floatation f/s	Refs
Dactylorchis incarnatus						Ziegenspeck, 1935
(probably Dactylorhiza incarnata)						1
Immediate wetting						
Wetting after 1 h					12	
Wetting after 24 h					$\frac{1}{2}$ $\frac{1}{2}$ 0 $\frac{3}{4}$ 0	
Sinking after 24 h					0	
Wetting after 60 h					$\frac{3}{4}$	
Sinking after 20 h					0	
Dactylorchis latifolius (probably						Ziegenspeck, 1935
Dactylorhiza latifolia)						
Immediate wetting					0	
Wetting after 1 h					1	
Wetting after 24 h					1	
Sinking after 24 h					0	
Wetting after 60 h					1	
Sinking after 20 h					0	
Dactylorchis maculatus (probably						Ziegenspeck, 1935
Dactylorhiza maculata)						
Immediate wetting						
Wetting after 1 h					1	
Wetting after 24 h					$\frac{\frac{1}{2}}{\frac{1}{2}}$	
Sinking after 24 h						
Wetting after 60 h					1	
Sinking after 20 h					0	
Dactylorchis sambucinus (probably					0	Ziegenspeck, 1935
Dactylorhiza sambucina)						Elegenspeek, 1995
Immediate wetting					0	
Wetting after 1 h					1	
Wetting after 24 h					1	
Sinking after 24 h					0	
Wetting after 60 h					1	
Sinking after 20 h					0	
						Charles 1001
Dactylorrhiza maculata		1	0	а. С	f	Stoutamire, 1981
Dendrobium nobile		1	0	1		Burgeff, 1936
Dendrobium speciosum	22.0				s	Stoutamire, 1981
Didymoplexis pallens* (0.45 mg)	32.0		1		1	Burgeff, 1936*
Encyclia tampensis		22		22	S	Stoutamire, 1981
Epidendrum Boundii		22	4	22		Burgeff, 1936
Epidendrum cochleatum		0	1	2	1	11
Epidendrum cochleatum		2	0	2		
Epidendrum nocturnum					s	Stoutamire, 1981
Epidendrum radicans		23		23		Burgeff, 1936
Epipactis gigantea					f	Stoutamire, 1981
Epipactis helleborine		14	<u>_</u>		f	DI 1 1 1000
Epipactis latifolia (9.1 µg, 90%)		11	5			Fleischer, 1929
(9.1 mg, 90%)	0.4	17	-	11		D (1007
Epipactis latifolia	9.1	16	5	11		Burgeff, 1936
Epipactis palustris		25	-	0.5		Fleischer, 1929
Epipactis palustris (92%)	7.5	32	7	25		Burgeff, 1936
Epipactis rubiginosa	10	16	7			Fleischer, 1929
Epipactis rubiginosa (90%)	6.8	23	7	16		Burgeff, 1936
Epipactis mean			89.5±2.67	10)		
Epipogium nutans	90-	120				Burgeff, 1936
Erythrodes querceticola					f	Stoutamire, 1981
Eulophia (2.73 µg)			1		1	Burgeff, 1936
Eulophia alta	Service and a				f	Stoutamire, 1981
Eulophia × Phaius flavus	23.0					Burgeff, 1936
Galearis spectabilis†					f	Stoutamire, 1981

Floatation time in air or water, or Time for dry seeds to become wet and continue to float Water (d) Species (when available Duration also seed weight (µg) Air and free air space (%)) Dry Wetted Difference of floatation f/s Refs (s) 60 6-8 Goodyera (2 µg, 71 %) Ziegenspeck, 1936 Goodyera (2 µg, 71 %) 19 8 Goodyera oblongifolia f Stoutamire, 1981 Goodyera pubescens f Goodyera repens 60 6 Fleischer, 1929 60 Goodyera repens 60 6 Burgeff, 1936 Goodyera repens f Stoutamire, 1981 Goodyera tesselata f Grammatophyllum speciosum 7.8 Burgeff, 1936 (2.48 µg) 9 5 Gymnadenia conopeat Fleischer, 1929 Gymnadenia conopeat Ziegenspeck, 1935 Immediate wetting Wetting after 1 h 1 Wetting after 24 h 1 0 Sinking after 24 h Wetting after 60 h 1 0 Sinking after 20 h 5 9 Gymnadenia conopea[‡] (8 µg) 4.8 14 Burgeff, 1936 9 5 Gymnadenia conopea‡ Ziegenspeck, 1936 Gymnadenia conopsea‡ f Stoutamire, 1981 Habenaria quinqueseta f 11 11 Habenaria repens f Haemaria discolor × H. dawsoniana 28.0 Burgeff, 1936 11 Haemeria discolor × H. rubrovenia 18 ίř. 2 2 Haemaria discolor var. trilineata × H. dawsoniana Harrisella porrecta s Stoutamire, 1981 Helleborine atropurpurea (now known Ziegenspeck, 1935 as Serapias atrorubens) Immediate wetting Wetting after 1 h Wetting after 24 h 0 Sinking after 24 h Wetting after 60 h Sinking after 20 h 0 7 Helleborine atropurpure (see 16 Ziegenspeck, 1936 Helleborine atropurpurea) Helleborine latifolia 11 11 5 (now known as *Epipactis helleborine*) 11 Heleborine palustris (now known as 25 6-8 Epipactis palustris) f Hexalectris spicata Stoutamire, 1981 9 Himantoglossum 12 Ziegenspeck, 1936 Himantoglossum hircinum 19 9 Fleischer, 1929 Himantoglossum hircinum Ziegenspeck, 1935 Immediate wetting Wetting after 1 h 1 Wetting after 24 h 1 0 Sinking after 24 h Wetting after 60 h 1 0 Sinking after 20 h 21 9 12 Burgeff, 1936 Himantoglossum hircinum Huntleya burtii f & s Stoutamire, 1981 Isotria verticillata f Laelia (~ 3.05 µg) 1-4 Fleischer, 1929 to

7-8

			ime in air come we				
		Water	r				
Species (when available also seed weight (μ g) and free air space (%))		(d)					
	Air (s)	Dry	Wetted	Difference	Duration of floatation	f/s	Refs
Laelia anceps×L. cinnabarina Laelia purpurata ×Cattleya citrina	4.3 4.5						Burgeff, 1936
(i.e. the hybrid genus Laeliocattleya)							
Laelia tenebrosa×Cattleya mendelii	2.3						11
(i.e. Laeliocattleya)							
Laeliocattleya		4	0	4			11
Laelia and							
Laeliocattleya mean Leucorchis albida	3.7±	1.22 (3	8.05 μg)				
(Leucorchis is now Didymoplexis) Immediate wetting							Ziegenspeck, 1935
Wetting after 1 h						2	
Wetting after 24 h						2 4 2 4	
Sinking after 24 h						2	
Wetting after 60 h						4	
Sinking after 20 h						2	D (C 102/
Limodorum abortivum	3.7						Burgeff, 1936
(5.7 μg, 96.58 %)						f	Stautomine 1091
Liparis elata Liparis forbesii			1		1	1	Stoutamire, 1981 Burgeff, 1936
Liparis Jordesti Liparis liliifolia			1		1	f	Stoutamire, 1981
Liparis loeselii (68%)	7.0	48	8	41		1	Burgeff, 1936
Liparis loeselii			0	1.0		f	Stoutamire, 1981
Listera borealis						f	
Listera convallariodes						f	11
Listera cordata						f	
Listera ovata		2	4				Fleischer, 1929
Listera ovata							Ziegenspeck, 1935
Immediate wetting							
Wetting after 1 h							
Wetting after 24 h						121	
Sinking after 24 h						0	
Wetting after 60 h						0	
Sinking after 20 h			4	2		0	D (C 102/
Listera ovata		6 2	4	2			Burgeff, 1936
Listera ovata Listera ovata		2	4			f	Ziegenspeck, 1936
Ludisia (0.85 mg)	23					1	Stoutamire, 1981
Luaisia (0.85 mg) Lycaste skinneri	23	1	0	1			Burgeff, 1936
Lycaste skinneri × L. lasioglossa	6.9		v				11
Malaxis ehrenbergii	14602					f	Stoutamire, 1981
Malaxis monophyllos						f	11
Neottia		4	2				Ziegenspeck, 1936
Neottia nidus avis		4	2 2 2				Fleischer, 1929
Neottia nidus	6.1	6	2	4			Burgeff, 1936
Nigritella nigra Immediate wetting							Ziegenspeck, 1935
Wetting after 1 h						2	
Wetting after 24 h						3	
Sinking after 24 h						1	
Setting after 60 h						3	
Sinking after 20 h	6.2					1	
Odontoglossum Odontoglossum historiense	6.3						D
Odontoglossum bictoniense	5.4						Burgeff, 1936

				Time for dry ue to float		
Species (when available also seed weight (μ g) and free air space (%))		Wate	r			
		(d)				
	Air (s)	Dry	Wetted	Difference	Duration of floatation f/s	Refs
Odontoglossum crispum × O. triumphans	7.2					Burgeff, 1936
Odontoglossum grande Odontoglossum grande	5.2	1	0	1	s	Stoutamire, 1981
Odontoglossum loochristiense§×O. inversum§	7.7					Burgeff, 1936
Odontoglossum mean	6.4	1.1				
Odontoglossum pulchellum × Miltonia warscewiczii						
Odontoglossum spp. and	6.3 <u>+</u>	1.0				
hybrids∥ Oncidium concolor × O. marshallianus	m 3 1					11
Oncidium incurvum \times O.	4.2					11
pulvinatum Oncidium floridanum (terrestrial race from Florida)					8	Stoutamire, 1981
Oncidium forbesii × O. marshallianum	72					Burgeff, 1936
Oncidium gartneri (probably	4.8					burgen, 1950
gardneri)§ × O. marshallianum						
Dncidium marshallianum	3.6					2.1
Oncidium rogersii§×O. concolor	5.9					11
Oncidium splendidum \times O.	4.3					4.4
cavendishianum						
Oncidium mean	4.7		12			
Ophrys apifera		3	3			Fleischer, 1929
Ophrys apifera						Ziegenspeck, 1935
Immediate wetting					4	
Wetting after 1 h					4	
Wetting after 24 h Sinking after 24 h					4	
Wetting after 60 h					4 0 4	
Sinking after 20 h					3	
Ophrys apifera		6	3	3		Burgeff, 1936
Ophrys apifera Ophrys aranifera		3	30			Ziegenspeck, 1936 Ziegenspeck, 1935
Immediate wetting Wetting after 1 h						
Sinking after 24 h					0	
Wetting after 60 h						
Sinking after 20 h					0	
Ophrys botteronii						Ziegenspeck, 1935
Immediate wetting						
Wetting after 1 h Wetting after 24 h					4	
Sinking after 24 h					4 0 4 3	
Wetting after 60 h					4	
Sinking after 20 h					3	
Ophrys fuciflora						Ziegenspeck, 1935
Immediate wetting Wetting after 1 h						1970 II) (1
Wetting after 24 h						
Sinking after 24 h					0	
Wetting after 60 h					222	
Sinking after 20 h					0	
Ophrys lutea						Ziegenspeck, 1935
Immediate wetting Wetting after 1 h					3	

402 TANSLEY J. Arditti and A. K. A. Ghani

				Time for dry ue to float		
	(11)	Water	r			
		(d)				
Species (when available	Air				Duration	
also seed weight (μg) and free air space (%))	(s)	Dry	Wetted	Difference	of floatation f/	s Refs
Wetting after 24 h					3	
Sinking after 24 h					0 3	
Wetting after 60 h					3	
Sinking after 20 h					0	
Ophrys muscifera		9	4			Fleischer, 1929
Ophrys muscifera						Ziegenspeck, 1935
Immediate wetting					2	
Wetting after 1 h					2	
Wetting after 24 h					2	
Sinking after 24 h					0	
Wetting after 60 h Sinking after 20 h					2 2 0 2 0	
Ophrys muscifera		13	4	9	0	Burgeff, 1936
Ophrys muscifera Ophrys muscifera		9	4			Ziegenspeck, 1936
Orchids, epiphytic		1-4	0			Fleischer, 1929
oremas, epipinyae		to 7-8				1 (clocher, 172)
Orchids (tropical) mean	(2.5)		µg seed ⁻¹)		Calculated from
orentate (cropical) mean	(2.00	11.00	<i>µ</i> 5 5000	,		data in Table 2.
						Range: 0.39-5.99
						seed ⁻¹
Orchis coriophorus (now coriophora)						Ziegenspeck, 1935
Immediate wetting						terrebutter provide a second
Wetting after 1 hour					3	
Wetting after 24 h					3	
Sinking after 24 h					3 3 3 3 3 3	
Setting after 60 h					3	
Sinking after 20 h					3	
Orchis incarnata	6.2					Burgeff, 1936
Orchis latifolia	5.6					1.1
Orchis longicornu	4.9					11
Orchis maculata		19	9			Fleischer, 1929
Orchis maculata	5.2	27	8	19		Burgeff, 1936
Orchis maculatus (maculata)		19	6-8			Ziegenspeck, 1936
Orchis mascula		11	7	123		Fleischer, 1929
Orchis mascula		18	7	11		Burgeff, 1936
Orchis masculus (mascula)						Ziegenspeck, 1935
Immediate wetting						
Wetting after 1 h					1	
Wetting after 24 h					1	
Sinking after 24 h					0 1 0	
Wetting after 60 h Sinking after 20 h					1	
Orchis masculus		11	7		0	Ziegenspeck, 1936
Orchis militaris		11				Ziegenspeck, 1935
Immediate wetting						Ziegenspeek, 1955
Wetting after 1 h					1	
Wetting after 24 h					1	
Sinking after 24 h						
Wetting after 60 h					0	
Sinking after 20 h					0	
Orchis morio						Ziegenspeck, 1935
Immediate wetting						o,,
Wetting after 1 h					2	
Wetting after 24 h					2	
Sinking after 24 h					2 2 1 2 2 2	
Wetting after 60 h					2	

Table 4 (cont.)

			'ime for dry ue to float			
		Water	r			
		(d)				
Species (when available also seed weight (µg) and free air space (%))	Air (s)	Dry	Wetted	Difference	Duration of floatation f/s	Refs
Orchis pallens		20	8			Fleischer, 1929
Orchis pallens		28	8	20		Burgeff, 1936
Orchis pallens		20	8	20		Ziegenspeck, 1936
Orchis paluster (now pallustris)						Ziegenspeck, 1935
Immediate wetting						
Wetting after 1 h						
Wetting after 24 h						
Sinking after 24 h					0	
Wetting after 60 h					(1 2 5)	
Sinking after 20 h					0	
Orchis provincialis						Ziegenspeck, 1935
Immediate wetting					1	Diegenopeen, tree
Wetting after 1 h						
Wetting after 24 h					4 5 3 5 3	
Sinking after 24 h					3	
Wetting after 60 h					5	
Sinking after 20 h					3	
Orchis purpureus (now purpurea)						Ziegenspeck, 1935
Immediate wetting						0
Wetting after 1 h					3	
Wetting after 24 h					3	
Sinking after 24 h					3 3 0 3 1	
Wetting after 60 h					3	
Sinking after 20 h					1	
Orchis sambucina	7.0					Burgeff, 1936
Orchis simia						Ziegenspeck, 1935
Immediate wetting						
Wetting after 1 h					2	
Wetting after 24 h					2	
Sinking after 24 h					2 2 0 2	
Wetting after 60 h					2	
Sinking after 20 h					1	
Orchis ustulatus (now ustulata)						Ziegenspeck, 1935
Immediate wetting					1	
Wetting after 1 h					4	
Wetting after 24 h					4 4 2 4	
Sinking after 24 h					2	
Wetting after 60 h					4	
Sinking after 20 h					3	
Orchis mean	5.8 <u>+</u>	0.8 (6	5.87%)			
Paphiopedilum cv.	4.1					Burgeff, 1936
Albertianum $\times P$. cv. Sallieri						
Paphiopedilum argus	6.1					31
Paphiopedilum	9.6					11
$callosum \times P.$ curtisii						
Paphiopedilum charlesworthi	3.5					11
Paphiopedilum curtisii	10.5		3		3	11
Paphiopedilum insigne var.	4.2					11
Sanderae $\times P$. concolor	IDEN.					24
Paphiopedilum	13					71
$lawrenceanum \times P.$ $glaucophyllum$						
Paphiopedilum spicerianum	5.0					11
Paphiopedilum cv.	11.0					11
Superciliare $\times P$. bellatulum	12/2011 21					11
Paphiopedilum	12.0					

Table 4 (cont.)

				'ime for dry ue to float			
	_	Water	r				
		(d)					
Species (when available	A :				D :		
also seed weight (μg) and free air space (%))	Air (s)	Dry	Wetted	Difference	Duration of floatation	f/s	Refs
Paphiopedilum	3.8						Burgeff, 1936
villosum × P. tonsum Paphiopedilum wallchii	16.0						11
Paphiopedilum mean		4313	0 μg, 89.62	70/_)			
Phaius flavus	29.0	. 1.5 (5.	2	/0/	2		1.1
Phaius cv. Marthae × Phalaenopsis	9.0		5075				11
aphrodite × Phalaenopsis sanderiana							
Phalaenopsis sanderiana	9.4						1.1
Phalaenopsis mean	92 (2	2.91 μg,	60.87%)				
Pholidota imbricata			1		1		11
Piperia unalascensis		-				f	Stoutamire, 1981
Platanthera bifolia		7	4				Fleischer, 1929
Platanthera bifolia						1	Ziegenspeck, 1935
Immediate wetting Wetting after 1 hour						22	
Wetting after 1 hour Wetting after 24 h						1 2 2 2	
Sinking after 24 h						0	
Wetting after 60 h						2	
Sinking after 20 h						2 1	
Platanthera bifolia	6.8						Burgeff, 1936
Platanthera bifolia		7	4				Ziegenspeck, 1936
Platanthera blephariglottis						f	Stoutamire, 1981
Platanthera chlorantha		11	5				Fleischer, 1929
Platanthera chlorantha	6.6	16	5	11			Burgeff, 1936
Platanthera chlorantha		11	5			6	Ziegenspeck, 1936
Platanthera dilatata						f	Stoutamire, 1981
Platanthera grandfilora						f	11
Platanthera hookeri						f	
Platanthera hyperborea						f f	11
Platanthera integra Platanthera lacera						r r	11
Platanthera leucophaea						f	0
Platanthera obtusata						f	11
Platanthera orbiculata						f	
Platanthera psycodes						f	11
Platanthera sparsiflora						f	1.1
Platanthera mean	6.7 (77.1 ± 1	0.1%)				
Pleurothallis pulchella	2	0	2				Burgeff, 1936
Pogonia ophioglossoides						f	Stoutamire, 1981
Polyrrhiza lindenii						s	11
Polystachya luteola	11.0					S	
Renanthera imschootiana Saccolahium bulehellum	11.6					0	Burgeff, 1936
Saccolabium pulchellum Sarcochilus australis						s	Stoutamire, 1981
Schomburgkia tibicinis						s	11
Schomburgkia undulata (0.3 µg)	6.6						Burgeff, 1936
Sobralia macrantha (6.3 µg)	3.7		1		1		11
Sobralia shoenbrunnensis × Sobralia macrantha	4.4						11
Spathoglottis (72 %)	7.8						10
Spathoglottis (12 76) Spathoglottis chrysantha	7.0		1		1		11
Spathoglottis chrysantha imes	6.1						11
Spathoglottis plicata	a anan Marco						
Spathoglottis fimbriata	8.4		1		1		11
Spathoglottis plicata	9.5		1		1		3.1

Table 4 (cont.)

				'ime for dry ue to float			
		Water	r				
Coursing (suban available		(d)					
Species (when available also seed weight (µg) and free air space (%))	Air (s)	Dry	Wetted	Difference	Duration of floatation	f/s	Refs
Spiranthes aestivalis Immediate wetting Wetting after 1 hour Wetting after 24 h Sinking after 24 h Wetting after 60 h Sinking after 20 h Spiranthes cernua Spiranthes cinnabarina Spiranthes lacera Spiranthes odorata Spiranthes romanzoffiana Stahopea oculata			1		1	3 3 1 3 2 f f f f f f	Ziegenspeck, 1935 Stoutamire, 1981
Stanoppea oculata \times S. quandricornis Stanhopea tigrina (2.95 µg, 89.31 %) Sturmia Sturmia Taeniophyllum sp. Thunia marshalliana Tipularia discolor Traunsteinera globosa Immediate wetting Wetting after 1 hour Wetting after 24 h Sinking after 24 h Sinking after 24 h Wetting after 20 h	6.1	3 41 41	1 6–8 2	2	2	s f 1 3 0 3 2	Fleischer, 1929 Ziegenspeck, 1936 Stoutamire, 1981 Burgeff, 1936 Ziegenspeck, 1935
Vanda suavis Vanda tricolor × V. burgefii§ Vanda tricolor × Vandopsis burgefii lissocholoides Vanda whiteana (now Vanda hindsii) Xylobium squalens Zeuxine strateumatica Zeuxine reflexa Zygopetalum mackayii	5.7 5.0 6.3 39.0	7	0 3	7	3	s f f & s	Stoutamire, 1981 Burgeff, 1936 Stoutamire, 1981 Burgeff, 1936 Stoutamire, 1981

Except in a few instances, all names and their spelling as well as systematic affiliations are as given in the original literature. Much of the published information on the subject cannot be used here because it is presented in the form of vague qualitative statements. Only a few reports are quantitative, and even some of these are flawed because of faulty experimental procedures, fragmentary data and numbers that are not validated by statistics (e.g. Stoutamire, 1981).

The 'dry' column lists the number of days during which half of the seeds that were thoroughly shaken daily remained on the surface. Floatation time in days of seeds pre-wetted with alcohol is shown in the 'wetted' column; it was presented originally as a measure of the duration of the effects of air in the testa on seed buoyancy. The 'difference' column being the difference between 'dry' and 'wetted', was described as a measure of the duration of the effect that air which adheres to cell walls has on buoyancy (Burgeff, 1909, 1936). This measure assumes (probably inaccurately) that bathing with alcohol does not affect wetting of the testa and/or its coating. Data in the 'duration' column are from another experiment in which the seeds were shaken daily (Burgeff, 1936). The numbers in this case represent the duration of seed floatation on the surface in days. Numbers in **bold** represent the time it takes for dry seeds to become wetted and their floatation periods. These experiments were carried out in laboratory water or solutions that may have been different from natural effluates. **Means are in bold italics**.

Some of the data (Burgeff, 1909, 1936; Fleischer, 1929; Ziegenspeck, 1935, 1936) seem to be repetitions, but they are presented separately due to several differences, which suggest that they may have been copied incorrectly, obtained independently or derived from similar or repeat experiments. It is difficult to determine which if any of these possibilities is correct because later authors do not always cite earlier ones for reasons which are not clear.

Floatation in air (in seconds) was measured by dropping seeds in a glass tube (1.5 m long and 40 mm in diameter) and recording the time required for them to reach a glass plate placed 50 mm under the cylinder. First to fall on the plate

which the seeds are released (in orchids this can vary from a few centimetres in terrestrials to many metres above ground for epiphytic species); \overline{U} , mean wind speed; α , parameter associated with horizontal diffusion; β , parameter associated with atmospheric diffusion; \overline{A} , mean vertical diffusivity (which should be low in slow-dropping orchid seeds).

The parameters can be calculated from measurements of: number of seeds released per unit time; height of release (H); wind speed at H; the settling velocity W_s ; and the dimensionless λ , which measures the relative spread about a mean for settling particles (Carey, 1998) that would be affected by air space in seeds (A. M. Ellison, pers. comm.). This equation produced a more satisfactory description of the distribution of *Himantoglossum hircinum* (Carey,

1998) despite the fact that air space in the seed (ca. 42%) and floatation time were not included in the equation. However, the calculation includes λ whose magnitude and effects can be expected to be the same as above (M. A. Ellison, pers. comm.). Altogether, orchid seeds can and do cover distances that may be as short as a few metres as suggested by Eqns 5 and 6, but also travel as far as thousands of kilometers (Table 5). The longer traveling will result from the increased variance and longer tail brought about by λ (Carey, 1998; Murren & Ellison, 1998). Given the many seeds produced by orchids, it is reasonable to assume that even if a small proportion will constitute the large variance and long tail, a sufficient number will travel far enough to colonize new areas: the available information is proof of that (Tables 4-6).

were seed clumps; these were followed by individual seeds; last to settle were empty testae. The error for descent times of 3–6 s was 0.1 s. For longer periods (up to 30 s) the error was 5 s (Burgeff, 1936). These measurements are not representative of natural conditions because the air inside the cylinder in Burgeff's laboratory was probably still. In nature the air moves constantly with varying velocities and directions even during a still day. These aerial eddies would tend to swirl the seeds and affect floatation time.

The tendency of orchid seeds to float or sink was investigated in several studies (Burgeff, 1909, 1936; Fleischer, 1929; Ziegenspeck, 1935, 1936; Stoutamire, 1981). In one of the the earlier studies (Ziegenspeck, 1935) 10 ml of the 'purest distilled water' were poured into glass containers and then seeds were carefully dusted onto the surface. The water was then stirred carefully for 2 min. Wetting, floating and sinking were evaluated after 1, 24 and 60 h and listed on a scale of '-' (no wetting or sinking), $\frac{1}{4}$ -4 (degrees of wetting) and 5 (complete wetting). The evaluations seem to have been purely subjective, but the scale was given an appearance of great accuracy by the use of quarter fractions. However, these fractions and whole numbers are purely subjective. Further, water under natural conditions is not 'purest distilled' and may contains substances that could affect wetting and/or floatation.

In the more recent 'f/s' report (Stoutamire, 1981) the only information presented is whether the seeds float ('f') or sink ('s') after being shaken for one minute in distilled water. This is not representative because there is no distilled water in nature. A more appropriate liquid would have been an effluate, even one collected from cultivated plants sprinkled with water. Additional details about these determinations are not given. In a batch of seeds of a species or a hybrid some will float and others will not, especially if the suspension is not agitated or shaken. Shaking for one minute, even in water only, may increase the number of seeds that sink. More seeds will sink after longer and/or repeated shaking. The addition of surfactants and/or media components (or leachates in nature) may increase the wetting and/or reduce the surface tension of water, and therefore reduce floatation time with or without agitation. Seeds that do not sink immediately may do so with time (hours or days) even if they are not shaken. Also, chaff which can appear to the naked eye like embryo-containing seeds, may never sink. However, seed or chaff could sink if they become filled (to some extent at least) or completely saturated with water. These facts and the single 'treatment' (shaking for one minute) raise questions about the data produced by this study (Stoutamire, 1981), the validity of its findings and the value of the information is presents. The information is presented here simply because it provides at least some indications in a field for which good data are scant.

*Burgeff visited the Bogor Botanical Gardens in Indonesia c. 1930 and worked with plants of this orchid which grew inside a clump of bamboo near one of the gates. These plants were still growing and flowering there in 1969. They did not reappear in 1972 and failed to show up again until at least 1981.

[†]The genus *Galearis* has been equated with parts of *Orchis* and *Habenaria*. There is no *Habenaria spectabilis*. On the other hand there is an *Orchis spectabilis*, but its synonym is *Galeorchis spectabilis*. Also, *Galeorchis* is now a synonym of *Orchis*. A reasonable conclusion is that the original paper (Stoutamire, 1981) is in error and should have given the genus as *Galeorchis* or *Orchis*.

[‡]It is not uncommon to see these spellings. The correct spelling seems to be G. conopsea.

§These are examples of Burgeff's lack of care in the use of orchid names: *Odontoglossum* cv. Loochritiense is not a species. It is a hybrid produced by the Belgian grower C. Vuylsteke who crossed *Odontoglossum crispum* (\mathfrak{P}) with an unknown pollen source. It was first described in the British *Orchid Review* in 1898.

A species named *Odontoglossum inversum* could not be traced. An *Odontoglossum* cv. Inversum has not been registered. The closest similarity in names is *Odontoglossum* cv. Inverness which was registered in 1958, 22 years after Burgeff's book was published.

Oncidium rogersii (no authority is given in the original) is actually Oncidium varicosum var. rogersii.

'Oncidium gartneri' (no authority is given in the original) probably represents confusion due to the similarity between 'gardener' and 'Gardner' and the German equivalent gärtner and Gartner. It is Oncidium gardneri.

'Paphiopedilum superciliare' is actually Paphiopedilum cv. Superciliare, a hybrid between Paphiopedilum barbatum and Paphiopedilum superbiens.

'Vanda burgeffii' is actually Vanda cv. Burgeffii, a hybrid registered by the Munich Botanical Gardens in 1928. In the early days of orchid hybridization, hybrids were often given Latinized names by recognized orchid taxonomists like John Lindley and H. G. Reichenbach filius.

If Burgeff followed the current practice of listing the seed parent first it is not surprising that the seeds of the hybrid are similar to those of *Odontoglossum*.

Table 5. 1	Distances	travelled	by	some	orchid	seeds
------------	-----------	-----------	----	------	--------	-------

Location or orchid	Distance covered	Refs
Agalega Island to Madagascar	600 km	Ridley, 1930
Agrostophyllum majus	81 m in Singapore	10
Aldabra to Madagascar	330 km	11
Anacamptis pyramidalis	'Some tens of kilometres' from the nearest	Willems, 1994
T.I. I	population of this species	D. U. 1020
Andaman Islands to Tenasserim	480 km	Ridley, 1930
Australia to New Zealand	2000 km	Close et al., 1978
Azores	1350 km	van Leeuwen, 1936
Azores to Madeira	825 km	Ridley, 1930
Azores to Portugal	1350 km	11
Bermuda to North Carolina, USA	870 km	11
Canary Islands to Cape Juby, Africa	75 km	11
Cape Verde Islands to Africa	750 km	11
Common dispersal distances		Lehaie, 1910 cited by Rasmussen, 1995
Christmas Island	200 km	van Leeuwen, 1936
Cymbidium acutum*	54 m in Singapore	Ridley, 1930
Cymbidium finlaysonianum	27 m in Singapore	11
Dendrobium pandaneti	1.5 km in Singapore	11
Epipactis latifolia†	12-15 km in the UK	Darwin, 1904
Epipactis latifolia†	12-14 km in England	Ridley, 1930
Galapagos	900 km	van Leeuwen, 1936
Hawaii	1000 km	
Iceland to Greenland	375 km	Ridley, 1930
Kermadak	900 km	van Leeuwen, 1936
Krakatau	4080 km	van Leeuwen, 1936; Went, 1957
Madeira to Morocco	675 km	Ridley, 1930
New Zealand from Australia		Close et al., 1978
No source or destination given	400 km	Garay, 1964
Ophrys apifera	60-75 km in Holland	Willems, 1994
Orchids, no names given	'Several hundreds of kilometres'	1
Orchis militaris	> 100 km	Willems, 1982 cited by Rasmussen, 1995
Orchis simia	<i>c</i> . 250 km	Crackles, 1975 cited by Willems, 1994
Orchis simia	< 25-200 km in Holland	Willems, 1994
Seychelles to Mauritius	1350 km	Ridley, 1930
Socotra to Cape Guardafui, Africa	750 km	
Trinidad	900 km	van Leeuwen, 1936
Trinidad (south) to the South American mainland	900 km	Ridley, 1930
Unidentified orchid seed Unidentified orchid seed	1350 km Found in the air 1800 m above sea level	Dafni, 1981

Spelling of all geographic locations as well as taxonomic affiliations and names (except as below) are as they appear in the original sources. For reviews see Ridley (1930), Gandawijaja & Arditti (1983), Arditti (1992), and Rasmussen (1995). Reported distances of travel between smaller and larger islands and/or land masses are presented here because orchid seeds may have traveled and/or still do travel between them (Ridley, 1930). * This orchid is now classified as *Cymbidium dayanum.*[†] The current name is *Epipactis helleborine*.

	Floatation in air	Weight	Air	Volume (µn	n ³)	Ratios					
	(s)	(ng)	space	Total	Free				12-95 (April 1		
Species	(F)	(W)	(%)	(TV)	(FV)	W:TV	W:FV	TV/W	FV/W	W/%	Refs
Acanthephippium sylhetense	13.0	660	98.45*	3465696	3412143	1.9×10^{-4}	1.93×10^{-4}	5251	5170	6.70	Burgeff, 1936
Angraecum eburneum × A. sesquipedale (A. Veitchii)	9.0	700									0
Anguloa ruckeri	13.0	390									0
Anguloa uniflora × A. ruckeri	10.0										11
Anguloa means	21.5	390									
Calanthe silvestris $\times C$. Darblayana	12.0										- 11
Calanthe Veitchii	12.0	500									J. Arditti & A. K. A. Ghani, unpublished
$Calanthe \ veratrifolia \times C. \ vestita$ $Calanthe \ means$	13.0	500	00 EE*	5222172	4721 707	0 28 10-5	1.06 - 10-4	106642	0442	ECE	Burgeff, 1936
Catantne means $Cattleva \ percivalliana \times C$, aurea	12.5 4.3	500	88.55*	5332172	4721707	$9.38 imes 10^{-5}$	$1.06 imes 10^{-4}$	106643	9443	5.65	0
Cattleya aurantiaca	4.5	2290	20.48	1095232	224363	2.09×10^{-3}	1.02×10^{-2}	478	98	112	Shoushtari et al.,
		2290	20.40	1095252	224303	2.09 × 10	1.02 × 10	470	90	112	1994
Cattleya guttata var. leopoldinae		5150									
Cattleya guttata \times C. autumnalis	5.0										Burgeff, 1936
Cattleya harrisoniae \times C.	5.2										
gaskelliana											
Cattleya labiata		2500									Hager, 1952
$Cattleya \ labiata \times C. \ gaskelliana$	5.6		12233/02250		121224231231232372						Burgeff, 1936
Cattleya lobata			76.86	5199840	3996601						Schlechter, 1915
Cattleya velutina \times C. gigas	4.6	1000								20.00	Burgeff, 1936
Cattleya trianae		4740	61.85	2469610	1527535	1.92×10^{-3}	3.1×10^{-3}	521	322	76.64	Schlechter, 1915;
											Shoushtari et al.,
0.00	4.94	3176	51.67*	2921561	1916166	1.09×10^{-3}	1.66×10^{-3}	920	603	61.46	1994
Cattleya means Cephalanthera damasonium	4.94	2000	78.63	10343552	8133326	1.09×10^{-4} 1.93×10^{-4}	2.46×10^{-4}	5172	4067	25.44	Rasmussen, 1995
Cephalanthera grandiflora	6.0	2000	75.03	10343 552	8133320	1.95 × 10	2.40 × 10	5172	4007	23.44	Burgeff, 1936
Cephalanthera granathora	6.0	2000	76.82	10343552	8133326	$1.93 imes 10^{-4}$	2.46×10^{-4}	5172	4067	26.03	Burgen, 1950
Corallorhiza innata	0.0	2000	83	10545552	8133320	1.93 ~ 10	2.40 × 10	5172	4007	20.05	Gandawijaja &
coratorniza innata			05								Arditti, 1983
Corallorhiza innata	7.2		74.36	9966360	7410749						Burgeff, 1936
Corallorhiza means	7.2		78.68	9966360	7410749						burgen, 1700
Cymbidium (estimated)		4000	87.49	21 535 586	18841545	1.85×10^{-4}	2.13×10^{-4}	5384	4710	45.72	J. Arditti &
2.1. Automotion second constrained Transformation of a second se Second second second second second se											A. K. A. Ghani, unpublished
Cymbidium Combidium	4.0		91.03								Dunneff 1026
Cymbidium devonianum Cymbidium giganteum × cv. Doris	4.9 3.0										Burgeff, 1936
Cymolaium giganleum x cv. Doris	5.0										

Table 6. Weight, volume, air space and floatation time in air of orchid seeds

Cymbidium insigne \times C. lowianum Cymbidium traceyanum \times C. insigne	4.6 5.0										Burgeff, 1936
Cymbidium means	4.38	4000	89.26*	21535586	18841545	$1.85 imes10^{-4}$	$2.13 imes 10^{-4}$	5384	4710	44.81	
Cypripedium acaule			78								J. Arditti & A. K. A. Ghani, unpublished
Cypripedium acaule Cypripedium calceolus		2000	85.51 93	33912000	28999602	5.90×10^{-5}	6.9×10^{-5}	16956	14500	23.39	0 0
Cypripedium calceolus Cypripedium californicum	5.9		90.98 92	16249274	14784645						Burgeff, 1936 J. Arditti & A. K. A. Ghani, unpublished
Cypripedium californicum Cypripedium californicum × C. acaule			95.03 90	40931470	38897111						n n
Cypripedium candidum			89								1.4
Cypripedium candidum			93.20	28617960	26672052						11
Cypripedium guttatum	7.3										Burgeff, 1936
Cypripedium montanum			87								J. Arditti & A. K. A. Ghani, unpublished
Cypripedium montanum			91	22608000	20573641						
Cypripedium passerinum			94.58	34265250	32407792						
Cypripedium pubescens	8.1		96								Burgeff, 1936
Cypripedium $pubescens \times C.$ acaule			93								J. Arditti & A. K. A. Ghani, unpublished
Cypripedium reginae			87								11
Cypripedium means	7.1	2000	90.35	29430659	27055897	$6.8 imes 10^{-5}$	7.39×10^{-5}	147159	1352	22.14	

	El	XX7 · 1 ·	A :	Volume (µm	³)	Deties					
	Floatation in air (s)	Weight (ng)	Air space	Total	Free	Ratios					
Species	(F)	(W)	(%)	(TV)	(FV)	W:TV	W:FV	TV/W	FV/W	W/%	Refs
Cyrtopodium punctatum	5.1	3400									Burgeff, 1936; Shoushtari <i>et al.</i> , 1994
Dendrobium antennatum		5650									Anonymous, 1879
Dendrobium attenuatum		6500									Ames, 1922b, 1948
Dendrobium crumenatum		8000									Shoushtari et al., 1994
Dendrobium insigne			60.64	785274750	476156391						Benzing & Clements, 1991
Dendrobium nobile		1000									Zinger &
											Poddubnaya
											Arnoldi, 1966
Dendrobium nobile			87.63	5023045	4401516						Ito, 1955
Dendrobium means		1000	87.63	5023045	4401516	$2.0 imes 10^{-4}$	$2.3 imes10^{-4}$	5023	4402	11.41	12 July 10 July 2020 1
Didymoplexis pallens	32.0	450									Burgeff, 1936
Epipactis gigantea			63.93	9286550	5936950						Arditti, 1992
Epipactis gigantea			87								J. Arditti &
											A. K. A. Ghani, unpublished
Epipactis helleborine			99.29	35026115	34778234						Curtiss, 1893
Epipactis nelleoorine Epipactis latifolia	9.1		99.29	55020115	34770234						Burgeff, 1936;
Epipacus iaujona	9.1		90								I. Arditti &
											A. K. A. Ghani,
											unpublished
Epipactis palustris	7.5		93								
Epipactis rubiginosa	6.8		90								-11
Epipactis means	7.8		87.20	22156332	20357592						
Eulophia horsfallii		2730									J. Arditti &
											A. K. A. Ghani,
											unpublished
Eulophia yashuiana				13687643							

Eulophia × Phaius flavus Eulophia means	23.0 23.0	2730		13687643		$2.0 imes 10^{-4}$		5014			Burgeff, 1936
Galeola altissima		17000		257900760		6.6×10^{-5}		15171			Hamada & Nakamura, 1963
Galeola lindleyana		14000									Burgeff, 1936
Galeola nudiflora				104333392	3						Barthlott & Ziegler, 1981
Galeola septentrionalis		24000									Nakamura, 1962, 1964
Galeola means		18333		650617342		$2.8\times\mathbf{10^{-5}}$		35489			
Goodyera procera			88.84	8374908	7439875						Link, 1839–1842
Goodyera pubescens			68.24	2520227	1719765						Curtiss, 1893
Geodorum densiflorum			63.78	18780820	6801782						Singh &
											Thimmappaiah, 1982
Geodorum means	1.000		63.78	18780820	11978038						
Geodorum pictum	1.000										Hallé, 1977
Goodyera repens		2000	67.65	32149336	21747334						Curtiss, 1893;
											Downie, 1940;
											Arditti, 1992;
											Rasmussen, 1995
Grammatophyllum speciosum	7.8	2480									Burgeff, 1936
Gymnadenia conopsea	4.8	8000	93.92*							85.19	
Laelia		3630									J. Arditti &
											A. K. A. Ghani,
											unpublished
Laelia anceps \times L. cinnabarina	4.3										Burgeff, 1936
Laelia purpurata × Cattleya citrina	4.5										11
Laelia tenebrosa × Cattleya mendelii	2.3										
Laelia means	3.7	3630									
Limodorum abortivum	3.7	5715	91.10*	40694400	37478784	1.4×10^{-4}	1.5×10^{-4}	7121	6559	62.73	Burgeff, 1936;
	5.7	5415	51.10	10071100	57470704	1.17.10	1.5 × 10	1121	0007	02.70	Veyret, 1969; J. Arditti & A. K. A. Ghani, unpublished

Table 6 (cont.)

	DI	XX7 · 1 ·	A.'.	Volume (µ	m ³)	D					
	Floatation in air (s)	Weight (ng)	Air space	Total	Free	Ratios					
Species	(F)	(W)	(%)	(TV)		$W\!:\!\mathbf{T}V$	$\mathbf{W}:\mathbf{F}\mathbf{V}$	TV/W	FV/W	W/%	Refs
Paphiopedilum Albertianum $\times P$. Sallieri	4.1										Burgeff, 1936
Paphiopedilum argus Paphiopedilum barbatum	6.1			569125							Beer, 1863; Barthlott, 1976
Paphiopedilum callosum × P. curtisii Paphiopedilum charlesworthi	9.6 3.5										Burgeff, 1936
Paphiopedilum curtisii	10.5		96.45	9385460	9052070						Schlechrter, 1915; Burgeff, 1936; Hoene, 1949; J. Arditti & A. K. A. Ghani, unpublished
Paphiopedilum insigne Paphiopedilum insigne	4.2	3000	27.78	8556067	2376090	3.5×10^{-4}	1.3×10^{-4}	2852	792	107.99	Zinger & Poddubnaya- Arnoldi, 1966; Lucke, 1984a Burgeff, 1936
Sanderae $\times P$. concolor Paphiopedilum lawrenceanum \times	13										
P. glaucophyllum Paphiopedilum parishii			99								
Paphiopedilum spicerianum	5.0										
Paphiopedilum superciliare × P. bellatulum	11.0										
Paphiopedilum tonsum × P. niveum Paphiopedilum villosum × P. tonsum	12.0 3.8										11 11
Paphiopedilum means Phajus	7.53	3000	97.5 92.58	6170217 7065000	5714080 6540578	$3.5 imes10^{-4}$	$1.3 imes10^{-4}$	2852	792	107.99	Davis, 1948
Phajus flavus	29.0		0.7717.72								Burgeff, 1936
Phajus maculatus		3530									Shoushtari et al., 1994
Phajus Marthae $\times P$. wallchii	16.0										Burgeff, 1936
Phajus means	22.5	3530	92.58*	7065000	6540578	$5.0 imes10^{-4}$	$5.4 imes 10^{-4}$	2001	1853	38.13	

Stanhopea means	5.9	2955	89.31*	12028398	10743062	2.5×10^{-4}	2.8×10^{-4}	4070	3635	33.09	
Stanoppea oculata× S. quandricornis	5.9										1929, 1930; Burgeff, 1936
Stanhopea oculata		3000	89.31	12028398	10743062	2.5×10^{-4}	2.8×10^{-4}	4010	3581	33.59	von Marilaun, 1896; Fleischer,
Sobralia means Stanhopea	5.05	6300 2910									Shoushtari et al., 1994
S. macrantha	1,1										
Sobralia shoenbrunnensis ×	4.4	0300									11
Schomburgkia undulata Sobralia macrantha	6.6 3.7	300 6300									
Renanthera imschootiana	11.6										Burgeff, 1936
Platanthera means	6.7		73.85	20232758	1164335						
Platanthera integrilabra			57.55	20232758	1164335						unpublished Zettler & McInnis, 1994
Flataninera chioranina	6.6		82								Burgeff, 1936; J. Arditti & A. K. A. Ghani,
Platanthera chlorantha			82								J. Arditti & A. K. A. Ghani, unpublished
Platanthera bifolia	6.8		82								Burgeff, 1936;
Phalaenopsis means	9.2	2910	42.17	1361975	216727	$2.1 imes 10^{-3}$	$1.3 imes 10^{-2}$	468	74.48	69.01	Burgen, 1950
Phalaenopsis mannii Phalaenopsis sanderiana	9.4	2910									Shoushtari et al., 1994 Burgeff, 1936
Phalaenopsis aphrodite × P. sanderiana	9.0										Burgeff, 1936
Phalaenopsis amabilis × P. rosea			15.91	1 361 975	216727						Bernard, 1909
· · · · · · · · · · · · · · · · · · ·											Linder, 1930; Bernard, 1937
Phalaenopsis			68,42	4108799	2811298						Linder 1020

Since information about weight, air space and floatation time for all orchids in this table is not available, figures for related species or estimations for genera are included for comparison purposes. The accuracy and reliability of these wide comparisons are clearly subject to errors. Total volume (TV) is the entire air space inside the testa. Free volume (FV) is the total volume minus the embryo volume.

Names used here are the ones given in the original literature.

Floatation in air (in seconds) was measured by dropping seeds in a glass tube (1.5 long and 40 mm in diameter) and measuring (with a stop watch) the time required for them to reach a white or black glass plate placed 50 mm under the cylinder. First to fall on the plate were seed clumps; individual seeds were next; the last to drop were empty testae. The error for drop times of 3-6 s was 0.1 s; for longer drop times, up to 30 s, the error was 5 s (Burgeff, 1936).

*These values and a number of others here were calculated from data for related species. Because of that some of them should be viewed as estimates. Calculations like these were made in cases where floatation times were available, but not sizes, weights and/or percent air space. The missing values generated by the calculations were used to construct several graphs (Figs 6,7) which depict relationships between floatation in air, weight, volume and percent air space ratios.

Box 2. Darwin's The Various Contrivances by Which Orchids are Fertilized by Insects

Darwin's *The Various Contrivances by Which Orchids are Fertilized by Insects* (Darwin, 1904) contains a valuable discussion of orchid seed number – despite the fact that he drew inaccurate conclusions. Note that Darwin did not italicize scientific names:

'The final end of the whole flower, with all its parts, is the production of seed; and these are produced by Orchids in vast profusion. Not that such profusion is anything to boast of; for the production of an almost infinite number of seeds or eggs, is undoubtedly a sign of lowness of organisation [inaccurate: orchids are now considered to be advanced plants]. That a plant, not being an annual, should escape extinction, chiefly by the production of a vast number of seeds or seedlings, shows a poverty of contrivance [inaccurate: orchids use almost all known contrivances to attract pollinators and disperse their seeds], or a want of some fitting protection against other dangers. I was curious to estimate the number of seeds on a long ruled line as equably as I could in a narrow hillock; and then counted the seeds in an accurately measured length of one-tenth of an inch. In this way the contents of the capsule were estimated at 6020 seeds, and very few of these were bad; the four capsules borne by the same plant would have therefore contained 24,080 seeds. Estimating in the same manner the smaller seeds of Orchis maculata [Fig. 1a], I found the number nearly the same, viz., 6200; and, as I have often seen above thirty capsules on the same plant, the total amount would be 186,300. As this Orchid is perennial, and cannot in most places be increasing in number, one seed alone of this large number yields a mature plant once in every few years.

To give an idea what the above figures really mean, I will briefly show the possible rate of increase of O. maculata: an acre of land would hold 174,240 plants, each having a space of six inches square, and this would be just sufficient for their growth; so that, making the fair allowance of 400 bad seeds in each capsule, an acre would be thickly clothed by the progeny of a single plant. At the same rate of increase, the grandchildren would cover a space slightly exceeding the island of Anglesea; and the great grand children of a single plant would nearly (in the ratio of 47 to 50) clothe with one uniform green carpet the entire surface of the land throughout the globe [see below]. But the number of seeds produced by one of our common British orchids is as nothing compared to that of some of the exotic kinds. Mr. Scott found that the capsule of an Acropera [Fig. 1b] contained 371250 seeds; and judging from the number of flowers, a single plant would sometimes yield about seventy-four millions of seeds. Fritz Müller [an early Darwin follower who wrote a book entitled Für Darwin; they corresponded at length, never met but held each other in high esteem – see Avadhani et al., 1994] found 1756440 seeds in a single capsule of a Maxillaria [Fig. 1a-c]; and the same plant sometimes bore half-a-dozen such capsules. I may add that by counting the packets of pollen (one of which was broken up under the microscope) I estimated that the number of pollen-grains, each of which emits its tube, in a single anther of Orchis mascula [Table 1] was 122400. Amici estimated the number in O. morio [Tables 2, 3] at 120300 [in some cases this orchid can flower 23 months after sowing: Frosch, 1983]. As these two species apparently do not produce more seed than the allied O. maculata [Tables 2, 3], a capsule of which contained 6200 seeds, we see that there are about twenty pollen-grains for each ovule. According to this standard, the number of pollen-grains in the anther of a single flower of the Maxillaria [Fig. 1a-c] which yielded 1756440 seeds must be prodigious.'

Darwin's calculations for clothing the globe with *O. maculata* are impressive, particularly because they are for a plant which produces only 6200 seeds capsule⁻¹. The process would be much faster for an orchid like *Cychnoches ventricosum* var. *chlorochilon* (formerly *Cycnoches chlorochilon*), which produces 4000000 seeds per fruit (Table 2). Theoretically this species can produce $65\ 536 \times 10^{24}$ individuals in four generations. The numbers for *Acropera* (*Gongora*) and *Maxillaria* would be 9×10^{51} and 144×10^{36} , respectively. *O. maculata* could produce as many as 81×10^{40} .

In the case of *Paphiopedilum*, the seeds of subgenus Brachypetalum are almost oval and 0.5-0.6 mm long (Karasawa & Saito, 1982). Examples of this subgenus are Paphiopedilum concolor (seeds oval, length 0.5 mm, air space 18%; Fig. 8), Paphiopedilum bellatulum (oval, 0.43 mm, 16 %; Fig. 8) and their natural hybrid (according to Karasawa & Saito, 1982), Paphiopedilum godefroyae (oval, 0.35 mm and 23 %; Fig. 8; percent air space calculated from measurements by Karasawa & Saito, 1982). The average air space of these seeds is 19%, and they probably have relatively short floatation periods. Species with such seeds can be expected to have a limited dispersal due to the constraints of the equations above and, indeed, this subgenus is found only in a contiguous land mass consisting of Burma, Thailand, southwest China, Yunan, Laos, Cambodia and Vietnam (Fig. 9; Karasawa & Saito, 1982; Cribb, 1987; Braem, 1988). Seeds of the subgenus Sigmatopetalum are longer. Examples are Paphiopedilum callosum (1.09 mm, 83%; Fig. 8), Paphiopedilum ciliolare (1.16 mm, 70%; Fig. 8) and Paphiopedilum acmodontum (0.79 mm, 69%; Fig. 8). These seeds have larger air spaces (an average of 74% or 3.9 times as much as Brachypetalum) and longer floatation times. Therefore their dispersal can be expected to be wider due to the increased variance and longer tail, and this is indeed the case. This subgenus is found in an area stretching from China through Java, Borneo, The Philippines and New Guinea to the Solomon Islands, even if the distribution of some species may be more limited (Fig. 9; Karasawa & Saito, 1982; Cribb, 1987; Braem, 1988).

Complicating factors in these considerations are the constant disagreement on taxonomy and nomenclature, and the short half-life of every 'new and

Box 3. Darwin on fruit and seed set in orchids

In Darwin's research on orchids, he produced important consideration of fruit and seed set. Note that Darwin did not italicize scientific names:

'The frequency with which throughout the world members of various Orchideous tribes fail to have their flowers fertilised, though these are excellently constructed for cross-fertilisation, is a remarkable fact.

Fritz Müller informs me that this holds good in the luxuriant forests of South Brazil with most of the Epidendreae, and with the genus Vanilla. For instance, he visited a site where Vanilla creeps over almost every tree, and although the plants had been covered with flowers, yet only two seedcapsules were produced. So again with an Epidendrum, 233 flowers had fallen off unimpregnated and only one capsule had been formed; of the still remaining 136 flowers, only four had their pollinia removed. In New South Wales Mr. Fitzgerald does not believe that more than one flower out of a thousand of Dendrobium speciosum sets a capsule; and some other species there are very sterile. In New Zealand over 200 flowers of Coryanthes triloba yielded only five capsules [see below] and at the Cape of Good Hope only the same number were produced by 78 flowers of Disa grandiflora. Nearly the same result has been observed with some of the species of Ophrys in Europe. The sterility in these cases is very difficult to explain. It manifestly depends on the flowers being constructed with such elaborate care for cross-fertilisation, that they cannot yield seeds without the aid of insects. From the evidence which I have given elsewhere we may conclude that it would be far more profitable to most plants to yield a few cross-fertilised seeds, at the expense of many flowers dropping off unimpregnated, rather than produce many selffertilised seeds. Profuse expenditure is nothing unusual under nature, as we see with the pollen of wind-fertilised plants, and in the multitude of seeds and seedlings produced by most plants in comparison with the few that reach maturity. In other cases the paucity of the flowers that are impregnated may be due to the proper insects having become rare under the incessant changes to which the world is subject; or to other plants ... '

Note that the genus *Coryanthes* and its pollinators are indigenous to Central and South America. There is no reason why it should set seed in New Zealand (unless hand-pollinated). In discussing *Coryanthes triloba* Darwin probably meant *Corysanthes trilobus*, now reclassified as *Corybas trilobus*. *Corybas* (*Corysanthes*) is native to New Zealand. *Corysanthes* and *Coryanthes* do not even resemble each other.

improved' classification system (the *Paphiopedilum* classification used here is that of two Japanese cytologists, Karasawa & Saito, 1982). Another classification could produce different or no correlations. As with examples in all areas of orchid biology it is necessary to keep in mind that the Orchidaceae is a large, complex and varied family and all models must be treated with caution. Very often it is necessary to use limited or even single

samples because there are no others. Even competent studies (Carey, 1998; Murren & Ellison, 1998) are limited by technical and biological constraints to only a single species each (fusiform, $640 \pm 40.6 \times 56 \pm 4.8 \mu m$, 1.6 µg for *Brassavola nodosa* and fusiform but stubby, $130 \times 30 \mu m$ and 5×10^{-9} kg for *Himantoglossum hircinum*). Would larger or smaller and/or heavier or lighter and/or differently shaped seeds give the same result?

(c) Birds. It is a mistake to assume that all travel by orchid seeds is through floatation in the air. For example, it was generally assumed that orchid seeds reached Krakatau on wind currents (van Leewen, 1936; Went, 1957; Garay, 1964; Gandawijaja & Arditti, 1983) following the volcanic eruption and destruction of the island in 1883 (Simkin & Fiske, 1983). However, Went (Went, 1957, 1990; F. W. Went, pers. comm.) suggested that orchid seeds could have reached the island on the legs or feathers of birds (exoornithochory). If transported on the legs of birds in mud or even on feathers the seeds probably arrived together with spores of fungi which could establish a mycorrhizal association and make germination possible (Went, 1957; F. W. Went, pers. comm.; Mitra, 1971; see Gandawijaja & Arditti, 1983).

Transport by birds, which is relatively quick, is also favoured because of the rapid loss of viability by seed stored under conditions that are generally unfavourable (Lucke, 1985) and/or permit rapid desiccation (Arditti, 1967, 1979, 1992; Arditti & Ernst, 1984). Conditions during long air-floatation periods (i.e. flights) may also lead to desiccation (Garay, 1964). Transport on muddy feet of birds or their feathers could reduce desiccation and make possible wider dispersal (Garay, 1964; Gandawijaja & Arditti, 1983).

According to one view, transport inside birds (endoornithochory) is not a means of orchid seed dispersal because of their inability to 'withstand the digestive fluids of birds' (Garay, 1964). On the other hand there are indications that seeds of Vanilla planifolia and Galeola septentrionalis (probably similar to those of Galeola galeata and Galeola altissima Fig. 1b) may be endoornithochorous (Nakamura & Hamada, 1978). The seeds of the latter may also be or endozoochorous (Hamada & Nakamura, 1978) . There is also a report that orchid seeds are scattered widely with excrement (Beer, 1863; Nakamura & Hamada, 1978). Treatments with a variety of solvents and surface sterilants suggest that orchid seeds may be able to withstand digestive tracts at least for a while (see Arditti, 1967, 1979, 1992; Arditti & Ernst, 1984). It is possible that ingestion by birds may even enhance germination of Vanilla planifolia seeds because of chemical changes brought about by 'juices' in the intestinal canal (Bouriquet, 1943; Stoll, 1954; Nakamura & Hamada, 1978). Chemical attack by digestive fluids could facilitate

germination by rendering the testa more permeable to water. Thus, endoornithocory and endozoochory cannot be fully excluded without experimental evidence. Some pertinent albeit indirect evidence should be relatively easy to obtain because the nature of digestive tracts and fluids in birds and other animals are known and could be simulated in a laboratory where orchid seeds can be subjected to them.

2. Water

(a) *Physical considerations*. Orchid seeds can float on water for prolonged periods (Table 4; Burgeff, 1936; Ziegenspeck, 1935, 1936; Rauh *et al.*, 1975; Stoutamire, 1981; Rasmussen, 1995) for the following reasons:

- They are filled with air (Table 1; Ziegenspeck, 1935, 1936; Carlson, 1940; Arditti, 1967, 1979, 1992; Arditti *et al.*, 1979, 1980; Healey *et al.*, 1980; Barthlott & Ziegler, 1981; Tohda, 1983; Arditti & Ernst, 1984).
- Their testae are hard to wet (Burgeff, 1936; Ziegenspeck, 1936; Arditti *et al.*, 1982; Rasmussen, 1995).
- The structure of the testa enhances floatation as in the case of *Disa uniflora* and several related species (Kurzweil, 1994).
- Buoyancy is enhanced by air bubbles trapped in the sculpturing of the testa (Ziegenspeck, 1936; Rasmussen, 1995).

Agitation (shaking or stirring) was employed in most experiments designed to study the floatation of orchid seeds in water. This seems to be a reasonably good approximation of natural condition because seeds may float in and be carried by moving, eddying and bubbling tree effluates and/or streams or rivulets of various sizes and velocities, which provide agitation. However, treating seeds with solvents, surfactants and/or other chemicals prior to floatation experiments could not possibly produce representative results because these treatments change the nature of the testa and/or any coating that may cover it as well as the surface tension of water. For example, surfactants accelerate the sinking of seeds in sodium hypochlorite solutions (Arditti et al., 1982). Hypochlorite (calcium and sodium) solutions in themselves also alter the outer chemical nature of the testa (Rasmussen, 1995). These changes undoubtedly affect wetting of the seeds and, as a consequence, also their floatation. Even distilled water (Stoutamire, 1981) is not a good approximation because under natural conditions water that may carry orchid seeds is actually a solution containing minerals and organic leachates, which could affect the testa.

Orchid seeds are difficult to wet because the outer walls of their testa cells are hard, lignified and covered with a cuticle (Ziegenspeck, 1936). The inner walls have no cuticle and are thinner. As the seeds ripen testa cells lose moisture and their walls curve inwards (Ziegenspeck, 1936). Both the cells and testas are filled with air. A mechanism that brings about the wetting of these hard to wet seeds was proposed more than half a century ago (Ziegenspeck, 1936). When seeds that fall into water or drop onto a moist substrate are chilled by cool water or reduced atmospheric temperatures the air inside the testa contracts. This creates suction that draws water into the seed through the micropylar opening (Fig. 3, 4; Ziegenspeck, 1936). Once inside the seeds, water enters testa cells through their thin inner walls (Ziegenspeck, 1936). Such a mechanism probably works well wherever even a small temperature differential follows the release of of orchid seeds.

In addition to entering testa cells from the inside, water within the seed also hydrates the embryos. The hydration causes the embryos to enlarge and emerge from the testa. Limited metabolic activity is also initiated in the hydrated seeds. Embryos may even turn green but do not develop into seedlings until they are colonized by an appropriate mycorrhizal fungus. Under laboratory conditions seeds placed on a medium that contains only minerals use their resources very slowly. Even the limited reserves inside the seeds are sufficient for survival periods of a year or more (Arditti, 1967, 1979, 1993, 1992). In nature, such long survival (almost suspended animation) periods increase the chances of fungal penetration. Thus the resrves in orchid seeds function not in sustaining germination, but in supporting the seeds until seedling development becomes possible. Clearly the evolution of orchid seeds is an exquisite contrivance that ensures dispersal and survival.

(b) Dispersal. Orchid seeds may not have been carried to Krakatau by ocean currents because they were reported to 'lose their viability after extended immersion in salt water ' (Garay, 1964). At present there is no infomation about the tolerance (or lack of it) of orchid seeds to sea water and/or saline conditions. It is also not known whether seeds will germinate after immersion in sea water and subsequent deposition onto terrestrial substrates (saline or not) such as rocks of phorophytes. This could be tested by immersing seeds in sea water for various periods and then attempting to germinate them. Orchid plants sometimes grow on rocks or trees near or over oceans where they are subjected to salt water mist or spray (Arditti, 1992). Seeds obviously germinate under these conditions, but it is not clear whether such germination is limited only to a few species that have adapted to salinity.

There is no doubt that orchid seeds fall into oceans (on Malaita, Solomon Islands several orchids, some in bloom, others in fruit grew on tree branches hanging from trees on rocky beaches; J. Arditti, unpublished). If orchid seeds could be carried by ocean currents they would be washed onto sandy or rocky beaches, not the interior of land masses, except if blown inland by wind after being deposited at the water's edge; carried inward in ocean mist; jettisoned onto seaside rocks and/or trees in the spray that results from crashing waves; and/or moved to the interior on the feet of animals that come to beaches. These possibilities have not been studied.

Fresh water does not seem to affect orchid seed deleteriously. Seeds of *Disa cardinalis*, *Disa uniflora* and *Disa tripetaloides* 'fall into streams and float down the water surface (this can be directly observed; due to their large size the seeds can be seen with the naked eye!)...and most of the seedlings occur on the banks of streams' (Kurzweil, 1994). European orchids are carried by rain water to areas where the soils are appropriate for germination (Fleischer, 1929; Burgeff, 1936; Ziegenspeck, 1936). In the tropics orchid seeds germinate on tree barks (Frey & Dodson, 1972).

VI. CONCLUSIONS

Another quotation from Darwin points to one function of the small, light, air-filled and buoyant orchid seeds (scientific names not italicized):

'The minute seeds within their light coats are well fitted for wide dissemination; and I have several times observed seedlings springing up in my orchard and in a newlyplanted wood, which must have come from a considerable distance. This was especially the case with Epipactis latifolia; and an instance has been recorded by a good observer of seedlings of this plant appearing at the distance of between eight and ten miles from any place where it grew...'

He also noted that:

'species... are sparingly distributed; yet, if their seeds or seedlings were not largely destroyed, any one of them would immediately cover the whole land ... The number of the individuals which come to maturity does not seem to be at all closely determined by the number of seeds which each species produces; and this holds good when closely related forms are compared.'

Among post-Darwinian deliberations regarding the nature of orchid seeds and their role in the evolution, dispersal, life cycle and physiology of orchids none are as perceptive and lucid as those by David Benzing (Benzing, 1981). His discussions are based on five generally accepted premises regarding orchids '(1) Orchidaceae is probably the largest of all angiosperm families [but similar claims are being made for the Asteraceae]; (2) its seeds are among the smallest of any taxon; (3) the resulting seedlings are mycotrophic; (4) many of its species are rare or widely dispersed; and (5) they often rely on very specialized pollination relationships ... ' and on one question: '... what about the tiny seeds and unusual mode of seedling nutrition? Individual seeds ... [may weigh]...0.01-0.1 mg for many orchids...'.

Benzing enumerates 'three distinct requirements' which necessitate 'close adherence to an optimal profile: (1) the heterotrophic demand of a seedling before it achieves autotrophy; (2) the need for dispersal to propitious sites; and (3) the necessity for sufficient fecundity to maintain a parent population.' He also points out that the characteristics of orchids seeds vary among taxa due to tradeoffs that are brought about by the need to survive under numerous 'combinations of constraints in native habitats.' One of these habitats, 'the patchy and infertile nature of the epiphytic biotope' is 'the rationale for the high fecundity and vagility required of plants living in the crowns of trees'.

Studies by Benzing and his associates (Benzing, 1981, 1987) showed that for epiphytes small seeds are advantageous because they increase the 'mineral economy and [counter] the high rates of juvenile mortality'. Terrestrial orchids also have to contend with inhospitable environments like 'infertile acid bogs and porous soils,' and 'physical constraints for some time'. On the basis of his considerations Benzing suggested 'that the seeds of orchids have become unusually small in response to strong reselecting forces imposed by unpredictably disturbed, patchy and stressed habitats. The miniaturization required to achieve a 'dust' type seed - the basis of the impressive fecundity mounted by these organisms - seems to have been made possible by major shifts in juvenile nutrition that, in effect, obviated the need for maternally supplied food at germination and for varying periods of time afterwards.'

The dependence of orchid seeds on mycorrhizal fungi for germination supports Benzing's views because it explains how young orchid seedlings survive in the field with very limited food reserves and an impaired ability to use what little they do have (Arditti, 1967, 1979, 1992; Arditti & Ernst, 1984; Arditti et al., 1990). During the early stages of germination orchid seedlings obtain nutrients from their fungi. In fact the orchids parasitize the fungi and this relieves the orchid embryo of its reliance on endogenous sources of nutrients. To put it differently, orchid seeds are small because they can get away with not having an internal endosperm. They obtain nutrients from an exogenous 'endosperm' (i.e. the fungal symbiont). Thus, Benzing concludes, 'a maternal parent, freed from the usual task of provisioning its young with substantial nutrient stores, is amenable to selective forces that encourage the production of unusually large numbers of small seeds'.

These 'unusually large numbers of small seeds' are neither wasteful burdens on the Orchidaceae nor 'lowness of organisation' or 'a poverty of contrivance'. They are two reasons for the success of the family.

ACKNOWLEDGEMENTS

We thank Roberk K. Josephson for allowing us to use his computing facilities to prepare figures; Wolfgang and Heidi Zierau and Robert Ernst for translations from German; Syoichi Ichihashi and Ken Cho for translations from Japanese; and Mort Arditti for gifts of computer software, literature and peripherals.

REFERENCES

- Ackerman JD, Zimmerman JK. 1994. Bottlenecks in the life histories of orchids: resources, pollination, population structure, and seedling establihsment. In: Pridgeon AM, ed. Proceedings of the 14th World Orchid Conference, Glasgow 1993. London, UK: HMSO, 125-129.
- Ames O. 1922a. Notes on New England orchids. II. The mycorrhiza of *Goodyera pubescens*. Rhodora 24: 37–46.
- Ames O. 1922b. Observations on the capacity of orchids to survive in the struggle for existence. Orchid Review 30: 229–234.
- Ames O. 1946. The evolution of the orchid flower. American Orchid Society Bulletin 14: 355–360.
- Ames O. 1948. Orchids in retrospect. Cambridge, MA, USA: Botanical Museum, Harvard University, USA.
- Ando T, Tomiyama M. 1990. Factors affecting on the germinaion of *Paphiopedilum* seed. In: Kimura T, Ichihashi S, Nagata H, eds *Proceeding of the Nagoya International Orchid Show*. Nagoya, Japan: The Organizing Committee of the Nagoya International Orchid Show, 1–1 Kawaseyama, Igayacho, Kariya-shi, Aichi-ken 448, 108–112.
- Anonymous. 1879. Weight of seeds. The Gardener's Chronicle Ser. 2, Vol 11: 178.
- Arditti J. 1961. Cycnoches ventricosum Batem. var. chlorochilon (Klotzsch) P. H. Allen comb. nov. Ceiba 9: 11–22.
- Arditti J. 1967. Factors affecting the germination of orchid seeds. Botanical Review 33: 1–97.
- Arditti J. 1979. Aspects of the physiology of orchids. Advances in Botanical Research 7: 241–665.
- Arditti J. 1984. An history of orchid hybridization, seed germination, and tissue culture. *Botanical Journal of the Linnean Society of London* 89: 359–381.
- Arditti J. 1990. Lewis Knudson (1885–1958): his science, his times, and his legacy. *Lindleyana* 5: 1–79.
- Arditti J. 1992. Fundamentals of orchid biology. New York, USA: John Wiley & Sons Inc.
- Arditti J. 1993. Storage and longevity of orchid seeds. Malayan Orchid Review (Singapore) 27: 59-63, 82-87.
- Arditti J, Clements MA, Fast G, Hadley G, Nishimura G, Ernst R. 1982. Orchid seed germination and seedling culture. In: Arditti J, ed. Orchid Biology Reviews and Perspectives, vol I. Ithaca, NY, USA: Cornell University Press, 243–370.
- Arditti J, Ernst R. 1984. Physiology of germinating orchid seeds. In: Arditti J, ed. Orchid biology reviews and perspectives, vol. III. Ithaca, NY, USA: Cornell University Press, 179–222.
- Arditti J, Ernst R, Yam TW, Glabe C. 1990. The contribution of orchid mycorrhizal fungi to seed germination: a speculative review. *Lindleyana* 5: 249–255.
- Arditti J, Michaud JD, Healey PL. 1979. Morphometry of orchid seeds. I. Native California and related species of *Cypripedium. American Journal of Botany* 69: 1129–1139.
- Arditti J, Michaud JD, Healey PL. 1980. Morphometry of orchid seeds. II. Native California and related species of Calypso, Cephalanthera, Corallorhiza and Epipactis. American Journal of Botany 67: 347-360.
- Avadhani PN, Nair H, Arditti J, Hew CS. 1994. Physiology of orchid flowers. In: Arditti J, ed. Orchid biology reviews and perspectives, vol. VI. New York, USA: John Wiley and Sons, Inc., 189–362.
- Barthlott W. 1974. Morphologie der Samen. Die Orchidee 25: 162–164.
- Barthlott W. 1976. Morphologie der Samen von Orchideen in Hinblick auf taxonomische und functionelle Aspekte. In: Senghas K, ed. Proceedings of the 8th World Orchid Conference Frankfurt. Frankfurt, Germany: German Orchid Society, 444– 455.

- Barthlott W, Ziegler B. 1980. Über ausziehbare Zellwandverdickungen als Haft-Apparat der Samenschalen von Chiloschista lunifera (Orchidaceae). Berichte der Deutsche Botanische Gesellschaft 93: 391–403.
- Barthlott W, Ziegler B. 1981. Mikromorphologie der Samenschalen als systematisch Merkmal bei Orchideen. Berichte der Deutsche Botanische Gesellschaft 94: 267–273.
- Beccari O. 1877. Malesia vol. 1. Genova, Italy: Tipografia del R. Instituto Sordo-Muti, 217.
- Bechtel H, Cribb P, Launert E. 1986. The manual of cultivated orchid species. Cambridge, MA, USA: MIT Press.
- Beer JG. 1863. Beiträge zur morphologie und biologie der familie der orchideen. Vienna, Austria: Druck und Verlag von Carl Gerold's Sohn.
- Benzing DH. 1981. Why is Orchidaceae so large, its seeds so small, and its seedlings mycotrophic? *Selbyana* 5: 241–242.
- Benzing DH. 1987. Major patterns and processes in orchid evolution: a critical synthesis. In: Arditti J, ed. Orchid biology, reviews and perspectives, vol. IV. Ithaca, NY, USA: Cornell University Press, 33-77.
- Benzing DH, Clements MA. 1991. Dispersal of the orchid Dendrobium insigne by the ant Iridomyrmex cordatus in Papua New Guinea, Biotropica 23: 604–607.
- Bernard N. 1909. L'evolution dans la symbiose, les orchidées et leur champignons commensaux. Annales des Sciences Naturelles, Botanique Ser. 9, Vol. 9: 1–196.
- Bernard N. 1937. Symbiose et parasitisme, l'œuvre de Noël Bernard. Paris, France: Masson et Cie.
- Bouriquet G. 1943. Sur la germination des graines de vanillier (Vanilla planifolia Andr.). Bulletin de l'Academie Malagache 16: 1–10.
- Bouriquet G. 1947. Sur la germination des graines de vanillier (Vanilla planifolia Andr.). L'Agronomie Tropicale 2: 150–1.
- Braem GJ. 1988. Paphiopedilum. Hildesheim Germany: Brücke-Verlag Kurt Schmersow.
- Burgeff H. 1909. Die Wurzelpilze der Orchideen, ihre Kultur und ihr Leben in der Pflanze. Jena, Germany: Verlag von Gustav Fischer.
- Burgeff H. 1911. Die Anzucht tropischer Orchideen aus Samen. Jena, Germany: Verlag von Gustav Fischer.
- Burgeff H. 1932. Saprophytismus und Symbiose. Studien an tropischer Orchideen. Jena, Germany: Verlag von Gustav Fischer.
- Burgeff H. 1936. Samenkeimung der Orchideen und Entwicklung ihrer Keimpflanzen. Jena, Germany: Verlag von Gustav Fischer.
- Carey PD. 1998. Modelling the spread of Himantoglossum hircinum (L.) Spreng. At a site in the north of England. Botanical Journal of the Linnean Society of London 126: 159-171.
- Carlson M. 1936. The germination of the seed and development of the seedlings of Calopon pulchellus (SW.) R. BR. Transactions of the Illinois State Academy of Sciences 28: 85–86.
- Carlson MC. 1940. Formation of the seed of Cypripedium parviflorum. Botanical Gazette 102: 295–301.
- Chase MW, Pippen JS. 1988. Seed morphology in the Oncidiinae and related subtribes (Orchidaceae). Systematic Botany 13: 313–323.
- Chase MW, Pippen JS. 1990. Seed morphology and phylogeny in Catasetinae (Orchidaceae). *Lindleyana* 5: 126–133.
- Cherevchenko TM, Kushnir GP. 1986. Orhidei v kulture (Orchids in cultivation). Kiev, Ukraine: Naukova Dumka.
- Chua LG, Rao AN. 1978. Polyembryony and suspensor characteristics in Spathoglottis. Flora 167: 399–402.
- Clifford HT, Smith WK. 1969. Seed morphology and classification of Orchidaceae. *Phytomorphology* 19: 133–139.
- Close RC, Moore NT, Tomlinson AI, Low AD. 1978. Aerial dispersal of biological material from Australia to New Zealand. *International Journal of Biometeorology* 22: 1–19.
- Cribb P. 1987. The genus Paphiopedilum. Kew, UK: Royal Botanic Gardens in association with Twickenham, UK: Collingridge Books, Hamlyn Publishing Group, Ltd.
- Curtis JT. 1932. A new Cypripedium hybrid. Rhodora 34: 239-243.
- Curtiss CC. 1893. Seeds of Orchidaceae. Journal of the Royal Microscopic Society 13: 653.
- Dafni A. 1981. Hasahlabim beisrael (Orchids of Israel, in Hebrew). Israel: Massada Ltd.

- Darwin C. 1904. The various contrivances by which orchids are fertilised by insects. London, UK: John Murray.
- Davis A. 1946. Orchid seed and seed germination. American Orchid Society Bulletin 15: 218–223.
- Davis A. 1948. Studies on orchid seeds. American Orchid Society Bulletin 17: 589–592.
- de Lehaie JH. 1910. Observations pour servir à l'étude de la dissemination des orchidées indigènes en Belgique. Bulletin de la Société Royale de Botanique de Belgique 47: 45-52.
- de Vogel EF. 1969. Monograph on the tribe Apostasieae (Orchidaceae). Blumea 17: 313-350.
- De Wit HCD. 1977. Orchids in Rumphius' Herbarium Amboinense. In: Arditti J, ed. Orchid biology, reviews and perspectives, vol. I. Ithaca, NY, USA: Cornell University Press, 47-94.
- Diem K, (ed). 1962. Documenta Geigi. Ardsley, NY, USA: Geigy Pharmaceuticals, Geigy Chemical Corporation.
- **Downie DG. 1940.** On the germination and growth of *Goodyera* repens. Transactions and Proceedings of the Botanical Society of Edinburgh **33**: 36–51.
- Dressler RL. 1993. Phylogeny and classification of the orchid family. Portland, OR, USA: Dioscorides Press.
- Fernandez Perez A. 1985. Flora de la Real expedicion botanica del Nuevo Reino de Granada. Vol. IX, 3. Madrid, Spain: Ediciones Cultura Hispanica.
- Fleischer E. 1929. Zur Biologie feilspanförmiger Samen. Botanisches Archiv 26: 86–132.
- Fleischer E. 1930. Zur Biologie feilspanförmiger Samen. Botanisches Zentralblatt 158: 91–92.
- Francini E. 1932. Lo svilupo del systema conduttore nelle plantule di alcune Orchidaceae. Nuovo Giornale Botanico Italiano N. S. 39: 226–242.
- Frei JK, Dodson CH. 1972. The chemical effect of certain bark substrates on the germination and early growth of epiphytic orchids. *Bulletin of the Torrey Botanical Club* 99: 301–307.
- Frosch W. 1983. Asymbiotische Vermehrung von Orchis morio midt der ersten Blüte nach 23 Monaten. Die Orchidee 34 (Sonderheft): 101–104.
- Füller F. 1969a. Ophrys. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Füller F. 1969b. Platanthera, Gymnadenia, Leucorchis. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Füller F. 1970a. Frauenschuh und riemenzunge. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Füller F. 1970b. Aceras und Anacamptis. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Füller F. 1972. Alpine und nordisch-alpine orchideen. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Füller F. 1981. Frauenschuh und riemenzunge. Wittenberg Lutherstadt, Germany: Die Neue Brehm Bücherei. A. Ziemsen Verlag.
- Gandawijaja D, Arditti J. 1983. The orchids of Krakatau: evidence for a mode of transport. *Annals of Botany* 52: 127-130.
- Garay LA. 1964. Evolutionary significance of geographical distribution of orchids. In: Burkill HM, Yeoh BC, Scott R, eds. Proceedings of the World Orchid Conference, Singapore 1963. Singapore: Straits Times Press, 170–187.
- George A, Cooke J. 1981. Rhizanthella: the underground orchid of Western Australia. In: Lawler L, Kerr RD, eds. Proceedings of the Orchid Symposium, Thirteenth International Botanical Congress, Sydney, Australia, 77–78.
- González-Díaz N, Ackerman J. 1988. Pollination, fruit set, and seed production in the orchid, Oeceoclades maculata. Lindleyana 3: 150–155.
- Gratiot J. 1934. Les orchidées, leur culture. Paris, France: Libraire Agricole de la Maison Rustique, Libraire de la Maison Rustique.
- Haas NF. 1977. Asymbiontische Vermehrung europäscher Erdorchideen. II. Nigritella nigra (L.) Rchb. f. and Nigritella miniata (Cr.) Janchen. Die Orchidee 28: 69–73.
- Hager H. 1954. Growing cattleyas from seed to flower in 2 years. American Orchid Society Bulletin 23: 78–81.
- Hallé N. 1977. Flore de la Nouvelle-Calédonie et dependances. 8. Orchidacées. Paris, France: Muséum National D'Histoire Naturelle.

- Hamada M. 1939. Studien über die Mykorrhiza von Galeola septentrionalis Reichb. f. Ein neuer Fall der Mykorrhiza-Bildung durch intraradicale Rhyzomorpha. Japanese Journal of Botany 10: 151–211.
- Hamada M, Nakamura SI. 1963. Wurzelsymbiose von Galeola altissima Reichb. f. Einer Chlorophyll-freien Orchidee, mit dem Holzzerstorenden Pilz Hymenochaete crocieras Berk. et Br. Scientific Reports of Tohoku University 4th Series (Biology) 29: 227-233.
- Harrison CR. 1977. Ultrastructural and histochemical changes during the germination of *Cattleya auratiaca* (Orchidaceae). *Botanical Gazette* 138: 41–45.
- Harrison CR, Arditti J. 1978. Physiological changes during the germination of *Cattleya aurantiaca* (Orchidaceae). *Botanical Gazette* 139: 180–189.
- Harvais G. 1973. Growth requirements and development of Cypripedium reginae in axenic cultures. Canadian Journal of Botany 51: 327–332.
- Hasegawa A, Goi M, Sato M. 1978. Fundamental studies on the asymbiotic seed germination of *Calanthe. Technical Bulletin*, *Faculty of Agriculture, Kagawa University* 29: 251–259.
- Healey PL, Michaud JD, Arditti J. 1980. Morphometry of orchid seeds. III. Native California and related species of Goodyera, Piperia, Platanthera, and Spiranthes. American Journal of Botany 67: 508–518.
- Henriksson LE. 1951. Asymbiotic germination of orchids and some effects of vitamins on *Thunia marshalliana*. Svensk Botanisk Tidskrift 45: 447–459.
- Hey GL. 1962. The mixed orchid house. Orchid Review 70: 315-316.
- Hey GL, Hey MG. 1966. Growing seeds of rare orchids. In: De Garmo LR, ed. Proceedings of the Fifth World Orchid Conference. Long Beach, CA, USA: Fifth World Orchid Conference Inc., and West Palm Beach, FL, USA: American Orchid Society, 35–38.
- Hirt W. 1906. Semina scobiforma, ihre Verbreitung im Pflanzenreich, Morphologie, Anatomie und biologische Bedeutung. Mitteilungen aus dem Botanischen Museum der Universität Zürich 30: 1–108.
- Hoene FC. 1945. Flora Brasilica Fasc. 8, Vol. XII, II: plate 69.
- Hoene FC. 1949. Inconographia de orchidaceas do Brasil. Sao Paulo, Brasil: Secretaria de Agricultura.
- Ito I. 1955. Germination of seeds from immature pod and subsequent growth of seedlings in *Dendrobium nobile* Lindl. *Scientific Reports of the Saiko University*, Agriculture 7: 35–42.
- Jacquet P. 1994. History of orchids in Europe from antiquity to the 17th century. In: Arditti J, ed. Orchid biology, reviews and perspectives, vol. VII. Dordrecht, The Netherlands: Kluwer Academic Publishers, 35–38.
- Jordão LR, Lopes VB, Takaki M. 1988. Selection of viable seeds in *Hormidium coriaceum* Ldl. (Orchidaceae) by density separation. Seed Science and Technology 16: 515-519.
- Karasawa K, Saito K. 1982. A revsion of the genus Paphiopedilum (Orchidaceae). Bulletin of the Hiroshima Botanical Garden 5: 1-69.
- Kim YJ, Lee JS, Yi YD, Moon RS. 1979. Exploitation of native orchid plants and their propagation for the floricultural crops. I. Wild orchids survey and propagation. *Journal of the Korean Society for Horticultural Science* 20: 94–105.
- Knudson L. 1921. La germinación no simbiótica de las semillas de orquídeas. Boletin de la Real Sociedad Española de Historia Natural 21: 250-260, plate 16.
- Knudson L. 1922. Nonsymbiotic germination of orchid seeds. Botanical Gazette 73: 1–25.
- Knudson L. 1929. Physiological investigations on orchid seed germination. Proceedings of the International Congress of Plant Science 2: 1183–1189.
- Knudson L. 1956. Self pollination in Cattleya aurantiaca (Batem.) P. N. Don. American Orchid Society Bulletin 25: 528–532.
- Kull T. 1998. Fruit-set and recruitment in populations of Cypripedium calceolus L. in Estonia. Botanical Journal of the Linnean Society of London. 126: 27-38.
- Kupper W, Linsenmaier W. 1961. Orchids. London, UK: Thomas Nelson and Sons Ltd.
- Kurzweil H. 1993. Seed morphology in Southern African Orchidoideae (Orchidaceae). *Plant Systematics and Evolution* 185: 229–247.

- Kurzweil H. 1994. The unusual seeds of the Disa uniflora group with notes on their dispersal. In: Pridgeon AM, ed. Proceedings of the 14th World Orchid Conference, Glasgow 1993. London, UK: HMSO, 397–399.
- Lenz LW, Wimber DE. 1959. Hybridization and inheritance in orchids. In: Withner CL, ed. *The orchids : a scientific survey*. New York, USA: Ronald Press, 261–314.
- Leuchs O. 1973. Bletilla striata die Japanorchidee. Der Palmengarten 37: 18.
- Leuschner JD. 1960. The culture of an orchid seed. Orchid Society of Southern California Review 2 (1): 8-11.
- Light MHS, MacConaill M. 1998. Factors affecting germinable seed yield in Cypripedium calceolus var. pubescens (Willd.) Correll and Epipactis helleborine (L.) Crantz (Orchidaceae). Botanical Journal of the Linnean Society of London 126: 3-26.
- Linder EA. 1930. The growing of orchids from seed. Missouri Botanical Garden Bulletin 18: 29–36.
- Link HF. 1839-1842. Icones selectae anatomico-botanicae. Berlin, Germany: Haude and Spener.
- Lucke E. 1981. Samenstruktur und Samenkeimung europäischer Orchideen nach Veyret sowie weitere Untersuchungen (Teil 1). Die Orchidee 32: 182–188.
- Lucke E. 1982a. Samenstruktur und Samenkeimung europäischer Orchideen nach Veyret sowie weitere Untersuchungen (Teil 2). Die Orchidee 33: 8–16.
- Lucke E. 1982b. Samenstruktur und Samenkeimung europäischer Orchideen nach Veyret sowie weitere Untersuchungen (Teil 3). Die Orchidee 33: 108–115.
- Lucke E. 1983. Eine Vakuumfritte f
 ür Orchideensamen. Die Orchidee 34: 72–74.
- Lucke E. 1984a. Samenstruktur und Samenkeimung europäischer Orchideen nach Veyret sowie weitere Untersuchungen (Teil 4). Die Orchidee 35: 13–20.
- Lucke E. 1984b. Samenstruktur und Samenkeimung europäischer Orchideen nach Veyret sowie weitere Untersuchungen (Teil 5). Die Orchidee 35: 153–158.
- Lucke, E. 1985. Zur Lagerung und Vitalitätsprüfung von Orchideensamen. Die Orchidee 36: 111–112.
- Maheshwari P, Narayanaswami S. 1952. Embryological studies on Spiranthes australis Lindl. Transactions of the Linnean Society of London 53: 474–486.
- Malguth R. 1901. Biologische Eigentumlichkeiten der Fruchte epiphytischer Orchideen. Inaugural dissertation, Königliche Universität Breslau. Breslau, Germany: Royal Botanic Garden.
- Mitra GC. 1971. Studies on seeds, shoot-tips and stem-discs of an orchid grown in aseptic culture. *Indian Journal of Experimental Biology* 9: 79–85.
- Mohana Rao PR, Sood S. 1979. Life history of Satyrium nepalense (Orchidaceae). Norwegian Journal of Botany 26: 285-294.
- Molvray M, Kores PJ. 1995. Character analysis of the seed coat in Spiranthoideae and Orchidoideae, with special reference to the Diurideae (Orchidaceae). American Journal of Botany 82: 1443–1454.
- Montalvo AM, Ackerman JD. 1987. Limitations to fruit production in *Ionopsis utricularoides* (Orchidaceae). *Biotropica* 19: 24-31.
- Murren JC, Ellison AM. 1996. Effects of habitat, plant size, and floral display on male and female reproductive success of the neotropical orchid Brassavola nodosa. Biotropica 28: 30–41.
- Murren JC, Ellison AM. 1998. Seed dispersal characteristics of Brassavola nodosa (Orchidaceae). American Journal of Botany 85: 675-680.
- Mutsuura O, Ito I, Nakahira R. 1962. Studies on the germination and the development of seedlings of *Taeniophyllum* aphyllum. (Makino) Makino. Scientific Reports of Kyoto Prefectual University (Natural Sciences and Living Sciences) Ser A, Vol. 3, No. 4: 13-18.
- Nakamura SI. 1962. Zur samenkeimung einer Chlorophyllfreien Erdorchidee Galeola septentrionalis Reichb. f. Zeitschrift für Botanik 20: 487–497.
- Nakamura SI. 1964. Einige experimente zur Samenkeimung einer Chlorophyllfreien Erdorchidee Galeola septentrionalis Reichb. f. Memoirs of the College of Agriculture, Kyoto University 86 (Botany Series) 4: 1-48.
- Nakamura SI, Hamada M. 1978. On the seed dispersal of an achlorophyllous orchid, Galeloa septentrionalis. The Journal of Japanese Botany 53: 260–263, plate III.

- Neiland MRM, Wilcock CC. 1995. Maximization of reproductive success by European Orchidaceae under conditions of infrequent pollination. *Protoplasma* 187: 39–48.
- Neiland MRM, Wilcock CC. 1998. Fruit set, nectar reward, and rarity in the Orchidaceae. American Journal of Botany 85: 1657–1671.
- Nilsson LA. 1979. Anthecological studies on the lady's slipper, Cypripedium calceolus (Orchidaceae). Botaniska Notiser 132: 329-347.
- Peakall R, Beattie AJ, James SH. 1987. Pseudocopulation of an orchid by male ants: a test of two hypotheses accounting for the rarity of ant pollination. *Oecologia* 73: 533–534.
- Peakall R, James SH. 1989. Outcrossing in an ant-pollinated clonal orchid. *Heredity* 62: 161–167.
- Perez-Arbelaez E, Alvarez Lopez E, Uribe Uribe L, Balguerias de Quesada E, Sanches Bela A, Riva Goday S. 1954. Flora de la Real expedicion botanica del Nuevo Reino de Granada, vol. I. Madrid, Spain: Ediciones Cultura Hispanica.
- Petersson B. 1991. The genus Nervilia (Orchidaceae) in Africa and the Arabian peninsula. Orchid Monographs 5: i-iv, 1-90, plates 1-4.
- Pfitzer E. 1882. Grundzuge einer Vergleichenden Morphologie der Orchideen. Heidelberg, Germany: Carl Winter's Universitätsbüchlandlung.
- Poddubnaya-Arnoldi VA. 1964. Obshtaya embriologia pokritosemenyih rastenyi (General embryology of angiosperms). Moscow, Russia: Nauka.
- Poddubnaya-Arnoldi VA. 1976. Tsitoembriologia pokritosmenyih rastenyi (Cytoembryology of angiosperms, principles and perspectives). Moscow, Russia: Nauka.
- Poddubnaya-Arnoldi VA, Selezneva VA. 1953. Viyrazivanye orhidei iz semyan (Cultivation of orchids from seeds). *Trudy Glavyi Botanicheskii Sad* 3: 106–124.
- Poddubnaya-Arnoldi VA, Selezneva VA. 1957a. Metodika semenogo razmonozenya orhidei (Methods for propagation of orchids from seeds). Biuleten Glavyi Botanicheskii Sad 27: 33-40.
- Poddubnaya-Arnoldi VA, Selezneva AV. 1957b. Orhidei i ih kultura (Orchids and their cultivation). Moscow, Russia: Akademia Nauki.
- Proctor HC, Harder LD. 1994. Pollen load, capsule weight, and seed production in three orchid species. *Canadian Journal of Botany* 72: 249–255.
- Quednow KG. 1930. Beiträge zur Frage der Aufnahme gelöster Kohlenstoffverbindlungen durch Orchideen und andere Pflanzen. Botanisches Archiv 30: 51–102.
- Raghavan V, Goh CJ. 1994. DNA synthesis and mRNA accumulation during germination of embryos of the orchid Spathoglottis plicata. Protoplasma 183: 137–147.
- Rasmussen HN. 1995. Terrestrial orchids, from seed to mycotrophic plant. Cambridge, UK: Cambridge University Press.
- Rauh W, Barthlott W, Ehler N. 1975. Morpholgie und Funktion der Testa staubförmiger Flugsamen. Botanische Jahbucher fur Systematik, Pflanzengeschichte und Pflanzengeographie 96: 353– 374.
- Ridley HN. 1930. The dispersal of plants throughout the world. Ashford, UK: Reeve & Co.
- Rolfe RA. 1912. Evolution of the Orchidaceae. Orchid Review 20: 260–264.
- Schlechter R. 1915. Die Orchideen. Berlin, Germany: Paul Parey.
- Schweinfurth C, Fernandez Perez A, Garay LA. 1969. Flora de la Real expedicion botanica del Nuevo Reino de Granada, vol. VIII. Madrid, Spain: Ediciones Cultura Hispanica.
- Schweinfurth C, Fernandez Perez A, Schultes RE. 1963. Flora de la Real expedicion botanica del Nuevo Reino de Granada, vol. VII. Madrid, Spain: Ediciones Cultura Hispanica.
- Scott RJ, Arditti J. 1959. Cymbidiums from pod to pot. American Orchid Society Bulletin 28: 823–829.
- Sheviak CJ, Bye RA Jr. 1980. Spiranthes dendroneura, a new species from Chihuahua. Brittonia 32: 368–371.
- Shoushtari BD, Heydari R, Johnson GL, Arditti J. 1994. Germination and viability staining following prolonged storage. *Lindleyana* 9: 77-84.
- Simkin T, Fiske RS. 1983. Krakatau 1883: the volcanic eruption and its effects. Washington, DC, USA: Smithsonian Institution Press.

- Singh G. 1981. Differential staining of orchid seeds for viability testing. American Orchid Society Bulletin 30: 416–481.
- Singh F, Thimmappaiah M. 1982. Polyembryony in orchid seeds. Seed Science and Technology 10: 29-33.
- Snow AA, Whigham DF. 1989. Costs of flower and fruit production in *Tipularia discolor* (Orchidaceae). *Ecology* 70: 11286–1293.
- Sood SK. 1989. Embryology and systematic position of *Liparis* (Orchidaceae). *Plant Systematics and Evolution* 166: 1-9.
- Stewart J. 1992. Orchids at Kew. London, UK: HMSO Publications Centre.
- Stoll S. 1954. Étude des éléments a caractères différentiels permettant la recherche et l'identification des différentes espèces de vanilla dans les produits alimentaires. In: Bouriquet G, ed. Le vanillier et la vanille dans le monde. Paris, France: Editions Paul Lechevlier, 616–622, plates XI–XVI.
- Stoutamire WP. 1963. Terrestrial orchid seedlings. Australian Plants 2: 119–122.
- Stoutamire WP. 1964a. Terrestrial orchid seedlings. II. Australian Plants 2: 264–266.
- Stoutamire WP. 1964b. Seeds and seedlings of native orchids. Michigan Botanist 3: 107–119.
- Stoutamire WP. 1981. Early growth in North American terrestrial orchid seedlings. In: Plaxton EH, ed. Proceedings from Symposium II and Lectures: North American Terrestrial Orchids. Southfield, MI, USA: Michigan Orchid Society and West Palm Beach, FL, USA: American Orchid Society, 14–31.
- Thomale H. 1954. Die Orchideen. Stuttgart, Germany: Eugen Ulmer.
- Tohda H. 1983. Seed morphology in Orchidaceae. I. Dactylorchis, Orchis, Ponerorchis, Chondradenia and Galeorchis. The Science Reports of the Tohoku University, 4th Ser. Biology 38: 253-268.
- Tohda H. 1985. Seed morphology in Orchidaceae II. Tribe Cranichideae. The Science Reports of the Tohoku University, 4th Ser. Biology 39: 21-43.
- Tohda H. 1986. Seed morphology in Orchidaceae III. Tribe Neottieae. The Science Reports of the Tohoku University, 4th Ser. Biology 39: 103-119.
- Tournay R. 1960. Orchidaceae Africanae. III. Le nombre de graines d'une capsule d'*Eulophia horsfallii. Bulletin du Jardin Botanique d'Etat* (Belgium) 30: 407–410.
- Tremblay RL. 1997. Lepanthes caritensis, an endangered orchid: no sex, no future. Slebyana 18: 160–166.
- van Leeuwen WMD. 1936. Krakatau 1883–1933. Leiden, The Netherlands: E. J. Brill.
- Vanseveren-Van Espen N. 1971. Aspects morphologiques de la germination de graines de Cattleya Lindl. (Orchidaceae) in vitro. Bulletin de la Société Royale du Botanique de Belgique 104: 29–45.
- Veyret Y. 1969. La structure des semences des Orchidaceae et leur aptitude à la germination in vitro en cultures pures. Travaux du Laboratoire de La Jaysinia, Muséum Nationale D'Histroire Naturelle, Paris Fascicle 3: 89–98: Plates III, IV.
- Vij SP, Pathak P, Mahant KC. 1995. Green pod culture of a therapeutically important species *Dactylorhiza hatagirea* (D. Don) Soo. *Journal of the Orchid Society of India* 9: 7–12.
- Vij SP, Sood A, Plaha KK. 1981. In vitro seed germination of some epiphytic orchids. In: Verma SC, ed. Contemporary trends in plant sciences. New Delhi, India: Kalyani Publishers, 473–481.
- von Marilaun A K. 1896. The natural history of plants (translated by Oliver FW, Busk M, Ewart MF). London, UK: Blackie and Son.
- Voth W. 1959. Die Aussaat und Sämlingsentwicklung von Peristeria elata Hook. Obst und Garten Ser B, 9: 6-9.

- Voth W. 1963. Asymbiotische Pflanzenaufzucht von Huntleya burtii und Pescatorea cerina. Obst und Garten Ser. B, 13: 224–229.
- Voth W. 1964. Asymbiotische Ausaat und Jungpflanzenanzucht einiger botanischer Orchideen. Obst und Garten Ser. B, 14: 250–258.
- Voth W. 1969. Über Aussaat und Sämlinge einiger Bulbophyllumund Cirrhopetalum-Arten. Die Orchidee 20: 272–277.
- Wagner J, Hansel A. 1994. In vitro seed germination of Cypripedium calceolus L. at various embryogenic stages. Angewante Botanik 68: 5-9.
- Weatherhead MA, Zee SY, Barretto G. 1986. Some observations on the early stages of development of *Eulophia* yushuiana. Memoirs of the Hong Kong Natural History Society 17: 85-90.
- Weiss FE. 1916. Presidential address. Annual Report and Transactions, Manchester Microscopical Society: 32–43.
- Went FW. 1957. The plants of Krakatau. In: Piel G, ed. Plant life - a Scientific American book. New York, USA: Simon and Schuster, 137–145.
- Went FW. 1990. Orchids in my life. In: Arditti J, ed. Orchid biology, reviews and perspectives. vol. V. Portland, OR, USA: Timber Press, 21–36.
- Wildhaber OJ. 1969. Zur Kenntnis von Epipogium aphyllum (Schmidt) Sw. Die Orchidee 20: 135–137.
- Wildhaber OJ. 1970. Beitrag zur Karpologie von Epipactis-Arten. Probleme der Orchideengattung Epipactis. Sonderheft der Zeitschrift Die Orchidee, May 1970: 109–113.
- Wildhaber OJ. 1972. Zur Karpologie von Orchis. Die Orchidee 23: 61–66.
- Wildhaber OJ. 1974. Zur Karpologie von Calypso bulbosa (L.) Oakes. Die Orchidee 25: 230.
- Willems JH. 1992. Establishment and development of a population of Orchis simia Lamk.in the Netherlands. New Phytologist 91: 757–765.
- Willems JH. 1994. Bottle-necks in establishment and survival of small populations of orchids in Western Europe. In:Brederoo P, Kapteyn den Boumeester DW, eds. Euroorchis 92, Proceedings of the International Symposium on European Orchids. Utrecht, The Netherlands: Stichting Uitgeverij Koninklijke Nederlandse Natuurhistorische Vereniging, 72–82.
- Withner CL. 1959. Orchid physiology. In: Withner, CL, ed. The orchids: a scientific survey. New York, USA: Ronald Press, 315-360.
- Zettler L, McInnis, TM Jr. 1994. Light enhancement of symbiotic seed germination and development of an endangered terrestrial orchid (*Platanthera integrilabra*). *Plant Science* 102: 133-138.
- Ziegenspeck H. 1935. Was bedingt die Schwimmfähigkeit der Samen der einheimischen Orchideen und der Sporen von Lycopodium. Botanisches Archiv 37: 373-382.
- Ziegenspeck H. 1936. Orchidaceae. In: von Kirshner O, Loew E, Schröter C, eds. Lebensgeschichte der Blütenpflanzen Mitteleuropas, vol 1, part 4. Stuttgart, Germany: Eugen Ulmer Verlag.
- Zimmerman JK, Aide TM. 1989. Patterns of fruit production in a neotropical orchid: pollinator vs. resource limitation. *Ameri*can Journal of Botany 76: 67–73.
- Zimmermann A, Dougoud R. 1961. Tropische orchideen. Creatura Naturkundliche K + F-Taschenbücher. Bern, Switzerland: Kümmerly & Frey Geographischer Verlag
- Zinger NV, Poddubnaya-Arnoldi VA. 1966. Application of histochemical techniques to the study of embryonic processes in certain orchids. *Phytomorphology* 16: 111–124.