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Development and evolution of extreme synorganization in angiosperm flowers and diversity: a comparison of Apocynaceae and Orchidaceae

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- **Background and Aims** Apocynaceae and Orchidaceae are two angiosperm families with extreme flower synorganization. They are unrelated, the former in eudicots, the latter in monocots, but they converge in the formation of pollinia and pollinaria, which do not occur in any other angiosperm family, and for which extreme synorganization of floral organs is a precondition. In each family extensive studies on flower development and evolution have been performed; however, newer comparative studies focusing on flower synorganization and involving both families together are lacking.
- **Scope** For this study an extensive search through the morphological literature has been conducted. Based on this and my own studies on flowers in various Apocynaceae and Orchidaceae and complex flowers in other angiosperms with scanning electron microscopy and with microtome section series, a review on convergent floral traits in flower development and architecture in the two families is presented.
- **Key Findings** There is a tendency of protracted development of synorganized parts in Apocynaceae and Orchidaceae (development of synorganization of two or more organs begins earlier the more accentuated it is at anthesis). Synorganization (or complexity) also paves the way for novel structures. One of the most conspicuous such novel structures in Apocynaceae is the corona, which is not the product of synorganization of existing organs; however, it is probably enhanced by synorganization of other, existing, floral parts. In contrast to synorganized parts, the corona appears developmentally late.
- **Conclusions** Synorganization of floral organs may lead to a large number of convergences in clades that are only very distantly related. The convergences that have been highlighted in this comparative study should be developmentally investigated directly in parallel in future studies.

Key words: Apocynaceae, Orchidaceae, angiosperm flower development, flower evolution, flower symmetry, synorganization, congenital fusion, postgenital fusion, pollinium, pollinarium, species diversity.

INTRODUCTION

The integration of floral organs, resulting in functionally fitting positions of all floral organs and covariation of fitting parts, is an evolutionarily important trend in angiosperms (Armbruster *et al.*, 2009, 2014). In contrast to ecological aspects, developmental aspects of integration have found much less attention (Wagner, 2014). A major developmental and evolutionary mechanism to increase such integration is synorganization of modules by highly symmetrical arrangement and tangential and radial congenital and postgenital fusion. Such synorganization is extreme in Apocynaceae (especially Asclepiadoideae) and Orchidaceae among angiosperms.

When we analyse a flower we commonly focus on the four organ categories sepals, petals, stamens and carpels. They are the organs that make up a flower. They are used in basic descriptions of flowers and are also the focus of molecular developmental studies on flowers. The classical ‘ABC model’ of flower development rests on them (Coen and Meyerowitz, 1991). In most flowers we encounter, these organs are easily seen and easy to distinguish from each other. However, looking at the flower of an orchid or an asclepiad, it is difficult to

recognize the basic organs immediately, especially in androecium and gynoecium.

Harder and Johnson (2008) discussed the function and evolution of aggregated pollen in angiosperms, also addressing Apocynaceae and Orchidaceae. It was shown that pollen loss during pollination in plants with pollinia is considerably lower than in plants with granular pollen (Harder and Routley, 2006). However, the evolution of pollinia and pollinaria needed complex morphological preconditions by intimate synorganization of floral organs, and this was only achieved in Apocynaceae and Orchidaceae convergently as a prominent novelty. The present paper focuses on the morphology, development and evolution of the highly synorganized flowers of these two families. Without intimate knowledge of the developmental processes and diversity it is difficult to understand the structural evolution of these flowers. There is an extensive and exciting literature on the functional aspects of flowers of Apocynaceae and Orchidaceae. However, the literature on comparative floral development and morphology in the two families is much smaller (e.g. Vogel, 1959; Schick, 1980, 1982a, b, 1988, 1989; Kunze, 1981, 1990, 1991, 1994, 1995, 1996, 1997, 2005; Fallen, 1986;

Kurzweil, 1987a, b, 1988, 1993, 1995, 1998; Kurzweil and Weber, 1992; Liede and Kunze, 1993; Endress, 1994, 2011; Liede, 1994; M. E. Endress, 2003; Kocyan and Endress, 2001; Kurzweil and Kocyan, 2002; Kunze and Wanntorp, 2008). As Apocynaceae and Orchidaceae are the only angiosperms that have pollinia and pollinaria, it is not surprising that initial comparisons between the two families were made long ago (Brown, 1833).

An important aspect of the convergent evolution of pollinia is economical use of available pollen and male fitness, which appears to be an important principle in floral biology (Barrett and Harder, 2006; Harder and Johnson, 2008). Various structural devices have evolved to this end (Erbar and Leins, 1995; Leins and Erbar, 2006, 2010), and the advent of pollinia is a particularly conspicuous trend in this respect.

Both families are species-rich, and the advent of pollinia and pollinaria may have been an important factor for this diversity. However, floral synorganization may also have led to other features driving speciation, such as devices forcing pollinators into precise positions on the flowers for pollinaria removal and deposition, or various ways of pollinator deception, especially in orchids. Apocynaceae comprises almost 5000 species. The subclade of Asclepiadoideae plus Secamonoideae, which is nested in Apocynaceae, has more species (3180) (Meve, 2002) than all other subclades of the family together (Periplocoideae, Apocynoideae and the basal grade of Rauvolfioideae), and it has 171 genera (M. E. Endress *et al.*, 2014), whereas the rest of the family has 194 genera (M. E. Endress *et al.*, 2014). Orchidaceae are one of the two most species-rich angiosperm families, with approx. 25 000 species and 735 genera (Chase *et al.*, 2015). Apostasioideae have two genera and 14 species (Chase *et al.*, 2015), Vanilloideae 14 genera and 245 species (Chase *et al.*, 2015), Cypripedioideae five genera and 169 species (Chase *et al.*, 2015), Orchidoideae 198 genera and 4575 species (Chase *et al.*, 2015), Epidendroideae 516 genera (Chase *et al.*, 2015) and 21 160 species (Freudenstein and Chase, 2015), and Cymbidieae and Vandaeae of Epidendroideae together have 300 genera and 4528 species (Chase *et al.*, 2015). Family stem ages were calculated as 52 Mya for Apocynaceae and 109 Mya for Orchidaceae (Magallón *et al.*, 2015). However, when each of the salient morphological innovations first appeared within each family is largely unknown.

FLOWER SYNORGANIZATION IN GENERAL

The general evolutionary trend of synorganization in flowers

Angiosperm flowers are characterized by basically three kinds of organs, which serve the following basic functions: (1) protection and optical attraction (the perianth organs: tepals or sepals and petals), (2) male function (the androecial organs: stamens) and (3) female function (the gynoecial organs: carpels). These floral organs are modular structures, repetitive units of the same kind. Each kind of organ occurs in different numbers and arrangement in a flower, although they always have the same sequence from the periphery to the centre of the flower: perianth→androecium→gynoecium. There is a common evolutionary trend in angiosperms that organs of the same kind or also organs of different kinds become more intimately associated. Such association or integration of ancestrally independent

organs into a complex structure is called synorganization. Thus, a common general evolutionary direction is from independent organs to a complex of organs. In the extreme, this complex of organs may become so synorganized that it behaves like a single complex unit or a single organ.

Preconditions for synorganization of floral organs

Precise localization of organs or organ parts is necessary for synorganization in development of the complex structure and at the same time for precise functioning of the flower [e.g. attachment of pollinaria to pollinators and deposition of pollinia on stigmas, such as in Catasetinae (Romero, 1990), see below]. An important precondition for such precise mutual position of the floral organs is whorled phyllotaxis, in contrast to spiral phyllotaxis (Endress, 1987, 1990, 2006). This allows (1) tangential, circumferential synorganization among the organs of a whorl (synsepal, sympetal, synstemon, syncarpy) and (2) sectorial synorganization (e.g. perianth organs and stamens in many monocots, and inner perianth organs and stamens in several basal eudicots) (Fig. 1). One of the most common tangential or circumferential synorganizations is syncarpy. It occurs in the majority of angiosperms (Endress, 1982). Also quite common is sympetal, which is present in many eudicots, especially asterids, and syntepal in several monocots. A combination of both tangential and radial synorganization also occurs, such as syntepal plus fusion between tepals and stamens on the same radii in many monocots, and between petals and stamens in many asterids and some rosids (e.g. Galipeinae of Rutaceae, El Ottra *et al.*, 2013), or between stamens and carpels in Orchidaceae and Apocynaceae (see below; Fig. 2).

Developmental processes and results of synorganization; different kinds of fusion

Synorganization most commonly occurs by means of (1) postgenital fusion or (2) congenital fusion, and (3) rarely without fusion. In postgenital fusion adjacent free organs or parts of organs make secondary contact and the contiguous epidermis fuses. In congenital fusion the primary meristems of adjacent organs become confluent so that these organs develop as a unity (confluence of meristems or meristem fusion); thus, the epidermis is not involved in the fusion process. Postgenital and congenital fusion often co-occur in gynoecia and at specific locations. Postgenital fusion has also been called epidermal or ontogenetic fusion, and congenital fusion has also been called phylogenetic fusion (Cusick, 1966). For a discussion on meristem fusion, see Hagemann (1973), and for molecular aspects of fusion, see the review by Specht and Howarth (2015).

A general advantage of congenital fusion of organs in a flower is that differential elongation of the basal and upper part of all organs makes possible a broad potential of shapes with various proportions of free and fused parts. Without congenital fusion this is not easily achieved.

An interesting feature of postgenital fusion is the potential for easy reopening at anthesis after tight closure in bud (Endress, 2006). Petals are often (partially or completely) postgenitally united in bud (and partially also at anthesis) in many asterids, especially campanulids (Araliaceae, Asteraceae,

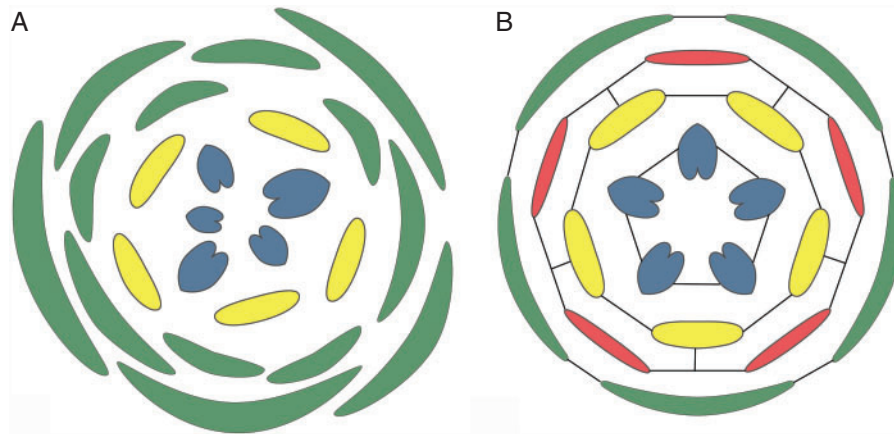


FIG. 1. Different potential for synorganization of floral organs in different phyllotaxis patterns. (A) Spiral phyllotaxis with limited potential. (B) Whorled floral phyllotaxis with high potential for radial and tangential synorganization. Black lines indicate preferred locations for synorganization.

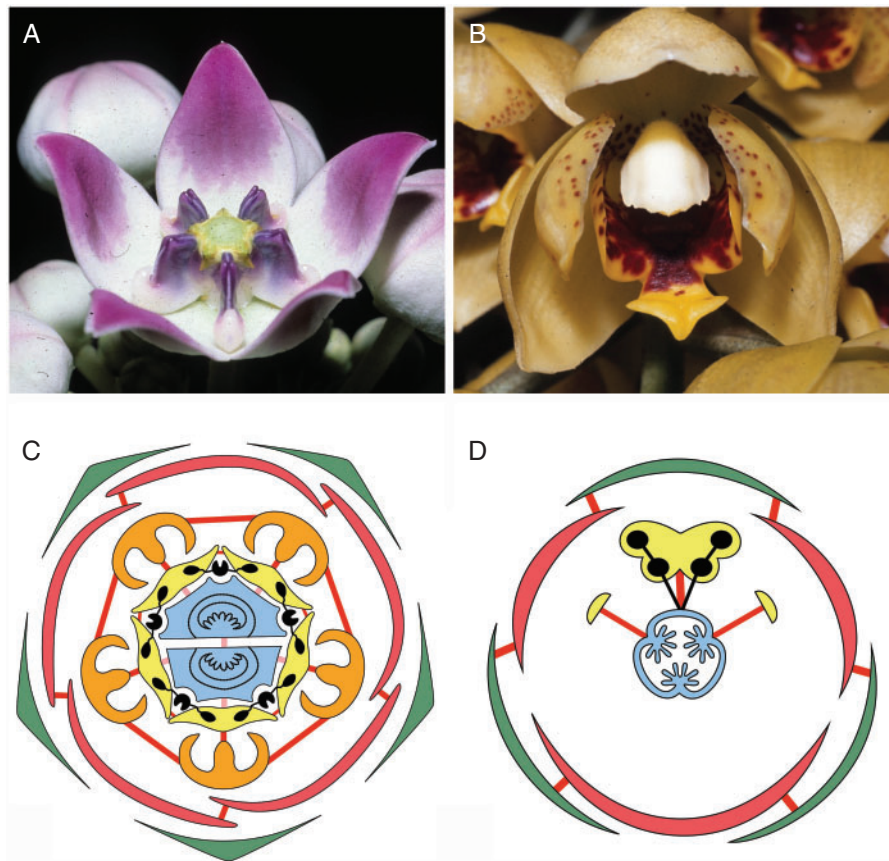


FIG. 2. (A, B) Flowers with high degree of symmetry and firm consistency of floral organs with plastic-like appearance. (A) *Calotropis procera* (Apocynaceae-Asclepiadoideae). (B) *Acineta densa* (Orchidaceae-Epidendroideae). (C, D) Floral diagrams with synorganizations indicated. Green: outer whorl of perianth; red: inner whorl of perianth; orange: corona; yellow: androecium; blue: gynoecium; black: pollinarium. Thick red lines: congenital fusion; thick pink lines: postgenital fusion. (C) *Asclepias* (Apocynaceae). (D) Monandrous Orchidaceae, flower shown in resupinate position.

Campanulaceae), in Gentianales (basal Apocynaceae *sensu stricto* (s.s.), [Fallen, 1986](#); Rubiaceae, [Robbrecht, 1988](#); basal lamiids: Icacinaceae, [Endress and Rapini, 2014](#)). Postgenital fusion is most common in gynoecia in intracarpellary and intercarpellary positions (e.g. [Baum, 1948](#); [Endress, 2015](#)). It is also

common in anthers, such as in many buzz-pollinated flowers ([Endress, 1994, 2006](#)) and in other complex flowers, such as in Balsaminaceae (e.g. [Vogel and Cocucci, 1988](#)). Rarely it occurs between stamens and carpels (balsaminoids, [von Balthazar and Schönenberger, 2013](#)).

A rare kind of intense synorganization without a particular fusion is known from *Geranium robertianum* (Endress, 2010). Here fusion only occurs in the gynoecium, which is syncarpous. All other synorganized parts (the sepals among themselves, and the sepals with petals, stamens and carpels) are free from each other but are held together by architectural modifications other than fusion.

General and conspicuous results of floral synorganization are an enhanced expression of the three-dimensional structure of the flowers and of internal morphological spaces. This may cause floral parts to become hidden. In addition, organs may become difficult to distinguish from their neighbours because their individuality becomes obliterated.

Also conspicuous is that synorganization leads to robustness or stability of the novel bauplan both in individual development and in evolution. Once the synorganized structures are established they are stable and are not easily lost again. This principle was nicely demonstrated by Simon (1962) with his watchmaker parable. At any new level of synorganization it is possible to experiment with variations at many single new points without harming the entirety of the bauplan and to attain new diversity.

EXTREME SYNORGANIZATION IN APOCYNACEAE AND ORCHIDACEAE

Apocynaceae (eudicots) and Orchidaceae (monocots) are the two families with the most extreme flower synorganizations among angiosperms. They are not phylogenetically related but have convergently evolved flowers with pollinia and pollinarium. A conspicuous difference is that the flowers of Apocynaceae are polysymmetric, those of Orchidaceae monosymmetric (Fig. 2).

Flowers of the most elaborate Apocynaceae (clade of Asclepiadoideae plus Secamonoideae)

It is easiest to begin with the most extremely complex flowers, thus flowers with the highest synorganization, and later show how the complexity develops during ontogeny and also show some of the evolutionary steps that led to the increasing synorganization. Floral organ numbers in the Asclepiadoideae plus Secamonoideae (this clade here called ‘ascleps’) are absolutely fixed: five sepals, five petals, five stamens and two carpels (as far as I know) in all 3100 species, without exception. Within some species, single aberrant flowers with four or six petals have been reported but are exceptional (Fuchs, 2013). In the basal grade of Apocynaceae it is similar, but there are a few taxa that have normally more than two (up to five) carpels (e.g. M. E. Endress *et al.*, 1997), or rare mutants with surplus petals in species with normally 5-merous flowers (Wang *et al.*, 2011). In ascleps, in addition, the floral organs of all floral whorls (except the sepals) are tangentially congenitally fused. Corona and stamens are also radially congenitally fused. Postgenital fusion occurs between the anthers and the style head and in the upper zone between the two carpels. In Apocynaceae in general, the corolla tube often has a postgenitally fused zone above the congenitally fused zone (e.g. Fallen, 1986; Kunze, 2005). This stability of the bauplan is necessary for synorganization. However,

there is plasticity at other structural levels to attain the present diversity.

The complexity of the flowers in asclepiads is highlighted by three structures: (1) gynostegium, (2) pollinarium (five per flower) and (3) corona. These structures are not basic floral organs, but rather are already synorganized organs or new parts enabled by synorganization of other structures (see also Endress, 1994).

The uppermost part of the gynoecium, the style head, and the anthers are postgenitally fused. This organ complex is called gynostegium. On the surface of the gynostegium the functional units for pollen transport are formed. The style head itself develops by postgenital fusion of the two carpel tips. The corona is a novel structure between corolla and androecium; it is often highly subdivided into a number of parts, which together form a complex apparatus with several functions. In its simplest form there is a corona element behind each stamen and is fused with it, but other parts of the corona are often also present between the stamens. The pollinaria are formed by the synorganized androecium and gynoecium.

The pollinaria are the strangest parts of the flowers. Each pollinarium is a composite apparatus for pollen transfer. Pollen from each theca is united to form a compact pollinium, and is thus not dispersed as single grains. In addition, pollinia are not transported singly, but always two together, one each from two neighbouring anthers. They become connected by a translator. Each translator consists of a clip and two arms. Each pollinium is connected to an arm. The translator does not consist of tissue, but is composed of sculpted secretion. It is unusual for secreted materials to attain such a complex and precisely formed structure.

This raises several questions: Where and how is this translator formed? What are the mechanisms to take the translators out of a flower? What are the mechanisms to position the pollinia at the right site in the recipient flower? The answers to these questions are in the precise synorganization of the floral parts, especially the corona and the gynostegium.

The five translators are formed each on a corner of the expanded, pentagonal style head (Fig. 2A, C). Each corner has a longitudinally directed concavity, which functions as the mould for the secretion of a clip (Fig. 3A). The two flanks of the corner secrete the arms. Each arm contacts the adjacent pollinium, which is presented at this site from the opened anther and is attached to the arm. In this way the clip is presented exactly in the middle of the upper end of a guide rail (Figs 4 and 5A).

In Asclepiadoideae, each guide rail is formed by the adjacent flanks of two neighbouring anthers. It guides pollinator body parts to the clip or to the stigma. Thus, there are five guide rails, five clips and five functional stigmatic units in each flower. The anther flanks that form the guide rails are the transformed (i.e. sterilized and histologically reinforced) dorsal pollen sacs of each theca. Thus, each anther produces only two pollinia (from the ventral pollen sacs). The anther flanks are postgenitally connected with the style head below the translator glands (Fig. 5B). The stigma is not on top of the gynoecium, but is located on the underside of the style head. It is subdivided into five functional units. Each unit is placed exactly in the radius of a guide rail. The stigma units are not visible from the outside because they are hidden in a concavity behind the guide rails on the underside of the style head (Fig. 5C). Below the stigma,

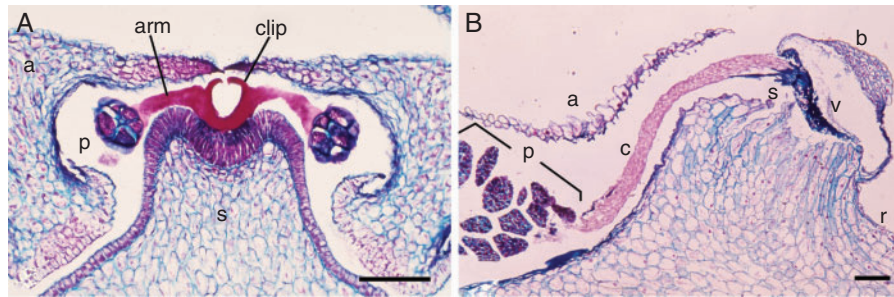


FIG. 3. Pollinaria and their position in the flower. (A) *Vincetoxicum nigrum* (Apocynaceae-Asclepiadoideae), microtome transverse section of gynostegium, showing one corner of style head (s) with secreted translator consisting of clip and two arms, which have become attached to the pollinium (p) of the two adjacent anthers (a) and forming the pollinarium. See also overview of this section in Fig. 5A. (B) *Ophrys fusca* (Orchidaceae-Orchidoideae), microtome longitudinal section of gynostegium, showing part of the pollinarium, consisting of sectile pollinium (p) subdivided into massulae, caudicula (c) and viscidium (v) with scutellum (s), covered by bursicula (b); anther wall (a); rostellum (r). Scale bars: A = 100 μ m, B = 200 μ m.

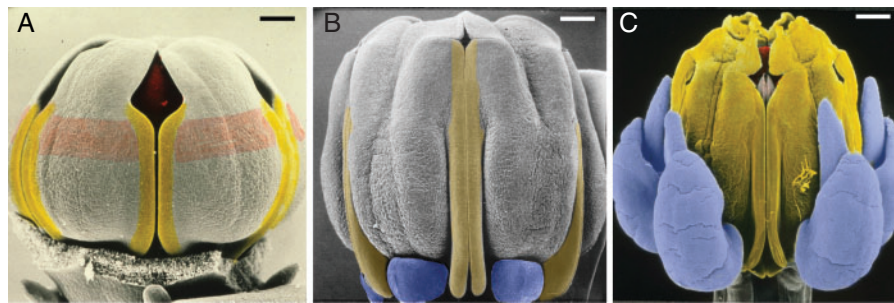


FIG. 4. Young flower buds of *Asclepias curassavica* (Apocynaceae-Asclepiadoideae) showing developmental steps of synorganization, scanning electron micrographs, all from the side, with one guide rail in the centre. (A) The anther flanks that later form the guide rails are marked with yellow. The zone of postgenital fusion of the anther flanks with the style head marked with pink. The corona elements are not yet formed. (B) Later stage. The anther flanks that later form the guide rail are marked with yellow. The corona elements, marked with blue, beginning to be formed. (C) Still later stage. The entire anthers are marked with yellow; the lower entrance of the guide rail has become wider. Corona elements are now much longer. Clip of the pollinarium above the guide rail is marked with pink. Part of the style head above the clip is marked with red. Scale bars: A = 100 μ m, B = 200 μ m, C = 500 μ m.

at the base of the guide rails, there are five nectaries in niches between the (at this level) congenitally fused stamens (and stamens fused with the corona elements in the same radii) (Fig. 5D). The nectaries may also secrete additional substances, perhaps in conjunction with pollen tube growth (Christ and Schnepf, 1985; Vieira and Shepherd, 2002). Thus, in the five radii of the guide rails, exactly aligned from bottom to top, are the nectaries, the stigmas and the clips of the translators. The guide rails are broadest at the base and taper toward the top where the clip is located (Fig. 5A–D). Thus, body parts of visiting insects will easily get caught in the basal part of a guide rail and then be drawn upwards exactly into the clip when the insect moves about on the flower or leaves the flower. Below the guide rails the corona elements are, in addition, congenitally fused with each other (Fig. 5E, F).

Movements of insects on the flower are greatly encouraged by the compartmentalized location of the nectar. In *Asclepias* nectar is stored (and presented) in ten sectors of the flower by troughs formed by coronal elements, which are provided with nectar by capillary forces from the five nectaries (Galil and Zeroni, 1965). These flowers are thus intricately differentiated revolver flowers, i.e. flowers in which nectar is available for pollinators not from a single position, but in this case from five or ten different positions.

When a pollinarium is drawn out of a flower it is at first positioned perpendicular to the insect, but it soon (about 1 min) bends forward by a differential drying process of the translator. The translator is not homogeneous but is composed of lipophilic and hydrophilic components, which cause bending by the drying process (Schnepf *et al.*, 1979). In this way it becomes optimally positioned for insertion into a guide rail of a recipient flower. If a pollinium comes into contact with the stigma, it sticks there and breaks off from the translator. The antestaminal elements of the corona also act as holding devices for insects and, because they protrude at the periphery of the floral centre, often the ends of the legs automatically come to lie into the guide rails, and most often the pollinaria become attached to the legs.

In most cases, an inserted pollinium provides only one of the two carpels with pollen tubes (Sage *et al.*, 1990; survey in Vieira and Shepherd, 2002; Vieira *et al.*, 2012). Thus, there is no functional compitum. Only rarely are both carpels served by the pollen tubes of a single pollinium (Kunze, 1991). There is a strange asymmetry because one carpel may be served from three stigmatic chambers, but the second carpel only from two. Often only one carpel develops into a fruit.

How can the style head of a dimerous gynoecium (Fig. 6A) produce five pollinaria? The mismatch between the

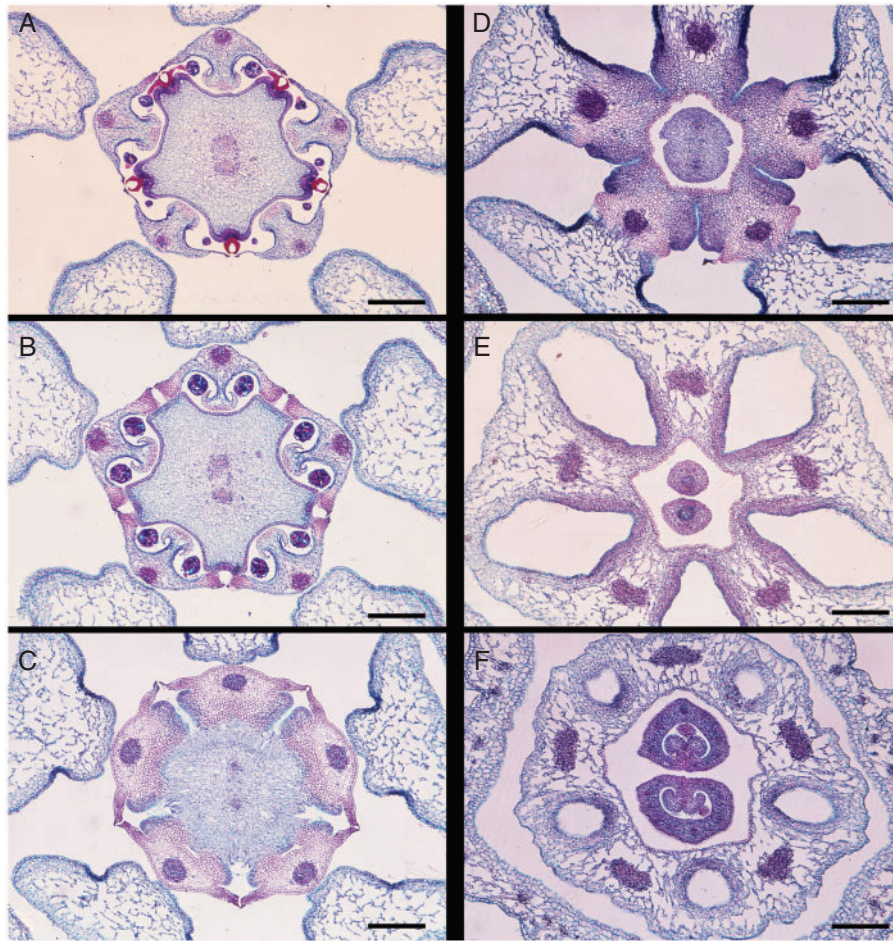


FIG. 5. Centre of a flower with gynostegium of *Vincetoxicum nigrum* (Apocynaceae-Asclepiadoideae), microtome transverse section series at six levels, from top, downwards. (A) Level of the five translator glands at style head. The two carpels postgenitally fused and forming five edges, each with a translator gland, the two main carpellary vascular bundles still distinct. (B) Level of postgenital fusion of anther flanks with style head and histological reinforcement of anther flanks for guide rail function. (C) Level of the five stigmas at lower end of style head. (D) Level of the five nectaries in the five grooves below the guide rails. The five corona portions congenitally fused with the stamens. (E) Level of the five corona/stamen portions congenitally fused with each other, leaving five holes between them. The two carpels free. (F) Level of corona and stamens forming a ring around the gynoeceum, at the base of the five holes. Carpels at the upper portion of the ovary. Scale bars: all = 500 µm.

pentamerous outer region of the flower and the dimerous gynoeceum has been overcome by the early postgenital fusion of the carpels and by complete conformation of the gynoeceum symmetry to that of the androeceum. Thus, the upper part of the gynoeceum becomes secondarily functionally pentamerous during development (Fig. 6B). This is nicely seen in cases in which the two carpels have a random, irregular position with respect to the outer floral whorls. This is an impressive example of an imprinted shape, a shape superimposed (moulded) by the five-angled shape of the immediate neighbourhood (Endress, 2008). The gynoeceum remains clearly dimerous at the base, but looks pentamerous on top at anthesis. Only the two vascular bundles in the style head at anthesis give testimony to its dimerous origin (Fig. 5A–C). The disposition of a pentamerous androeceum and dimerous gynoeceum is most common in Gentianales (and in early branching Apocynaceae), and thus is most probably ancestral in the family and was present in the evolutionary history before the synorganization of androeceum and gynoeceum.

Among asclepiads, diversity is expressed in particular in the shape of the corolla and corona. The corona is highly plastic

with regard to nectar holder function (Kunze, 1997). Units of the corona may be simple in small flowers (e.g. *Vincetoxicum*) or complex in larger flowers (Fig. 7) (e.g. *Asclepias*, Galil and Zeroni, 1965, or *Hoya*, Kunze and Wanntorp, 2008), and convoluted in *Calotropis* (Puri and Shiam, 1966). Stamen shape in Asclepiadoideae is strongly influenced by the integration of all parts of the gynostegium (Liede, 1994; Kunze, 1996). The shape of the style head (Simões *et al.*, 2007a) and the depth of the guide rails are evolutionarily plastic, correlated with the size of the pollinators (Fig. 8). Diversity in size of the entire flowers is addressed below. The synorganization of flowers in Asclepiadoideae underlying the diversity as described here is a constant feature through the subfamily, as several detailed morphological and developmental studies have shown (Corry, 1884; Demeter, 1922; Kunze, 1981, 1990, 1991, 1994, 1995, 2005; Demarco, 2014).

A pinnacle of complexity is exhibited by the flowers of *Ceropegia*, a genus of over 200 species (Huber, 1957; Vogel, 1961; The Plant List, <http://www.theplantlist.org/>; Figs 9 and 10). Here, not only are the corona, androeceum and gynoeceum

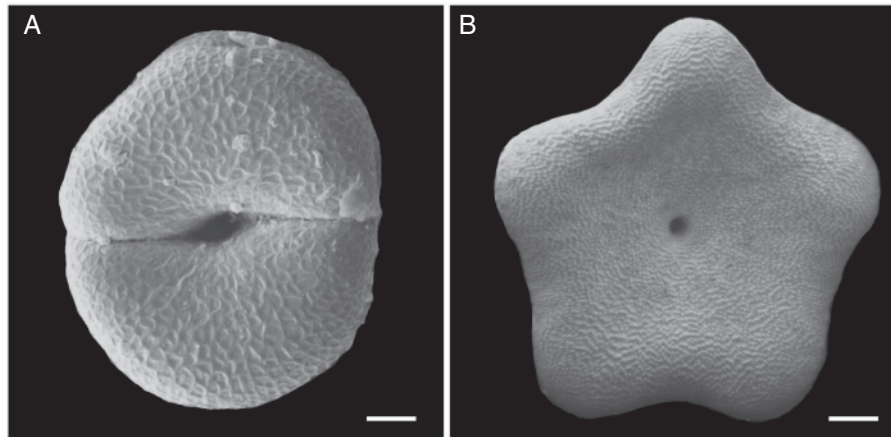


FIG. 6. Style head moulding from disymmetry to pentasymmetry in two developmental stages, from above, in *Gomphocarpus fruticosus* (Apocynaceae-Asclepiadoideae), scanning electron micrographs (modified from Endress, 2006). (A) Very young stage, with the two carpels still distinct. (B) Older stage, with the two carpels postgenitally united and outline changed to five-angled. Scale bars: A = 50 μm , B = 500 μm .

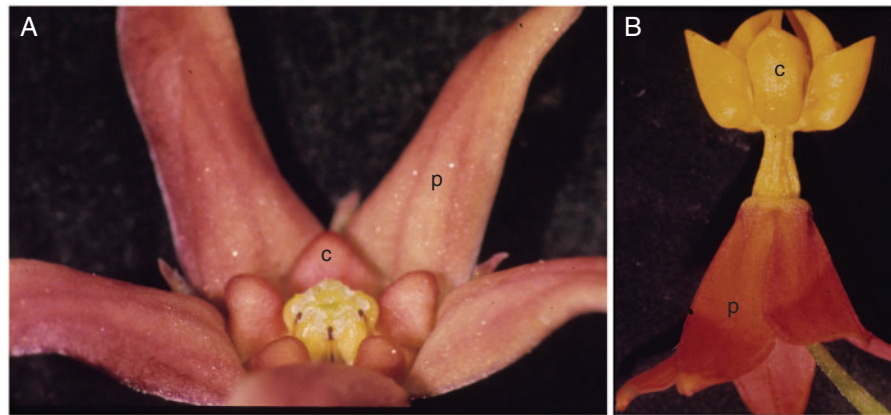


FIG. 7. Flowers of Apocynaceae-Asclepiadoideae to show diversity in proportions of petals (p) and corona (c). (A) *Vincetoxicum nigrum*, with relatively small and simple corona. (B) *Asclepias curassavica*, with relatively large and complex corona.

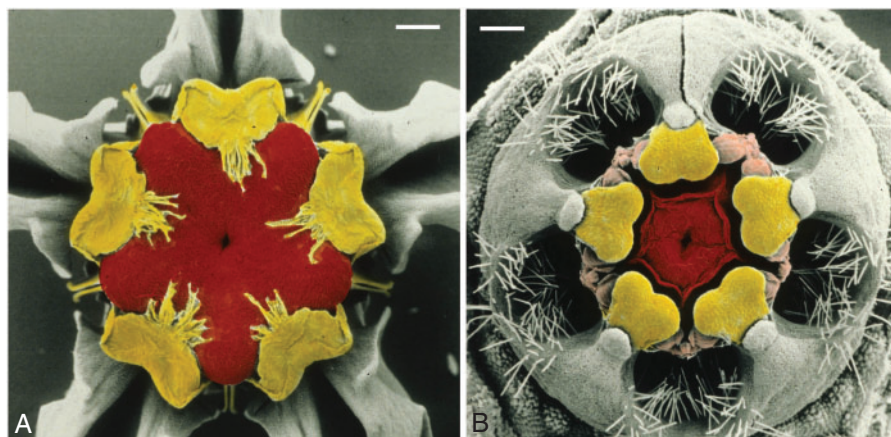


FIG. 8. Diversity in extension of the edges of the style head (marked with red), depth of the guide rails (marked with yellow, together with uppermost part of the stamens) and exposition of the pollinaria (marked with pink), correlated with pollinator size, in Apocynaceae-Asclepiadoideae. (A) *Gomphocarpus fruticosus* (style head edges not extended, guide rails deep, pollinaria hidden – for large pollinators). (B) *Caralluma penicillata* (style head edges long extended, guide rails shallow, pollinaria exposed – for small pollinators). Scale bars: both = 500 μm .

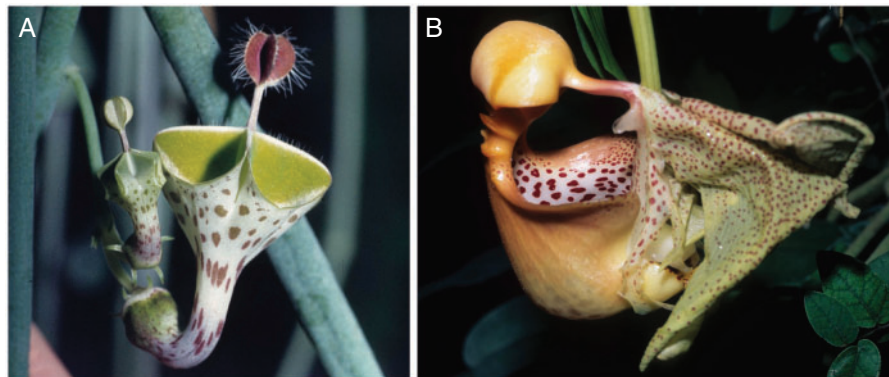


FIG. 9. Flowers with extremely complex corolla. (A) *Ceropegia distincta* (Apocynaceae-Asclepiadoideae). (B) *Coryanthes macrantha* (Orchidaceae-Epidendroideae).

intimately synorganized, but also the corolla is involved in the complexity in a unique way. The flowers are trap flowers with a long and slender tube formed by congenital fusion of the petals. The complex pollination apparatus is small and hidden in the base of the corolla tube. Thus, not only are parts of the androecium and gynoecium hidden, but the entire pollination apparatus. In the uppermost zone of the corolla the petals are free but are postgenitally fused in bud. At anthesis this postgenitally fused zone opens differentially. In the most complex species, such as *C. distincta*, at the base it opens to provide five separate entrances into the corolla tube, each with a wax-covered gliding zone. Then follows a zone where the petals remain fused and form a kind of central stalk. Above this, a second zone of opening presents a flag with the inner petal surfaces turned outward and acting as an osmophore to attract flies (Ollerton *et al.*, 2009). The tips of the petals remain closed, forming a firm top of the flag. Some species have flickering hairs directed outwards at anthesis in the opening zones, which add to the attractiveness for flies. Thus, the corolla exhibits five zones with regard to differential fusion, from base to top: (1) congenitally fused, (2) open, (3) postgenitally fused, (4) open and (5) postgenitally fused (Fig. 10).

Stepwise synorganization of the flowers in the phylogeny and evolution of Apocynaceae sensu lato (s.l.)

The phylogeny of Apocynaceae s.l. has been intensively studied in the past 20 years, and for some subclades aspects of floral evolution have been analysed (Nilsson *et al.*, 1993; M. E. Endress *et al.*, 1996, 2007a, b, 2014; Sennblad and Bremer, 1996, 2002; Civeyrel *et al.*, 1998; Sennblad *et al.*, 1998; M. E. Endress and Bruyns, 2000; M. E. Endress, 2003, 2004; M. E. Endress and Stevens, 2001; Rapini *et al.*, 2003, 2006, 2007; Verhoeven *et al.*, 2003; Simões *et al.*, 2004, 2007b, 2010; Liede-Schumann *et al.*, 2005; Ionta and Judd, 2007; Livshultz *et al.*, 2007; Livshultz, 2010; Rapini, 2012; Nazar *et al.*, 2013; M. E. Endress *et al.*, 2014; Straub *et al.*, 2014). In the basal subclades of Apocynaceae (grade of Rauvolfioideae), the degree of floral synorganization is relatively low. There are no pollinaria, no pollinia and no gynostegium. The very top of the gynoecium is commonly not secretory, but the secretory part somewhat lower down is without functional differentiation into receptive

and non-receptive zones in some groups (e.g. *Condylocarpon*, *Aspidosperma*, M. E. Endress, pers. comm.). An incipient corona may be present as alternipetalous lobes on the corolla (corolline corona) as also in the sister family Gentianaceae; in some genera of Rauvolfioideae these corona lobes are conspicuous. The corolla is sympetalous, and the stamens are fused with the corolla and have two pollen sacs per theca. However, the flowers are already revolver flowers, as also in some other polysymmetric asterids.

There are trends in the evolutionary pathway of the pollinia and pollinaria and of style head differentiation. In Rauvolfioideae and Apocynoideae, pollen is normally dispersed as single grains, but in both subfamilies there are some taxa with tetrads. The proboscis of pollinating insects becomes sticky by touching the style head and takes up pollen, which may be deposited on the stigmas of recipient flowers. Schick (1982a) distinguished two types and Fallen (1986) found four levels of increasing complexity of the style head in Apocynaceae s.s.: (1) the entire style head is secretory and (probably) stigmatic in its entirety; (2) the stigma is restricted to the lowermost part of the style head below a downward directed collar, which collects incoming pollen, whereas the upper part of the style head secretes the adhesive for pollen transport, and an upper ring of hairs presents the pollen shed from the anthers; (3) in addition, the anthers are postgenitally fused with the style head, and thus the gynostegium emerged (occurs only in Apocynoideae); (4) the downward directed collar and the upper ring of hairs disappeared and, instead, hairs are present on the ventral side of the anthers, which scrape incoming pollen from the proboscis of insects. The most interesting innovation in Apocynoideae, found in *Apocynum* and *Forsteronia*, is a precursor of a translator. It consists of five small platelets of a gummy consistency secreted between the anthers on the style head, which take up pollen from the adjacent thecae of two anthers and are removed and transported by pollinators (Demeter, 1922; Nilsson *et al.*, 1993; M. E. Endress, 2003); pollen still occurs as single grains (as tetrads in *Apocynum*), and the translator is without attachment device to pollinators, except for sticky secretion.

In Periplocoideae, the translator is spoon-shaped. Pollen is transported in the form of free tetrads or tetrads packed in soft pollinia (without pollinium wall) (Verhoeven and Venter, 2001), and is deposited in the sticky concave part of the spoon.

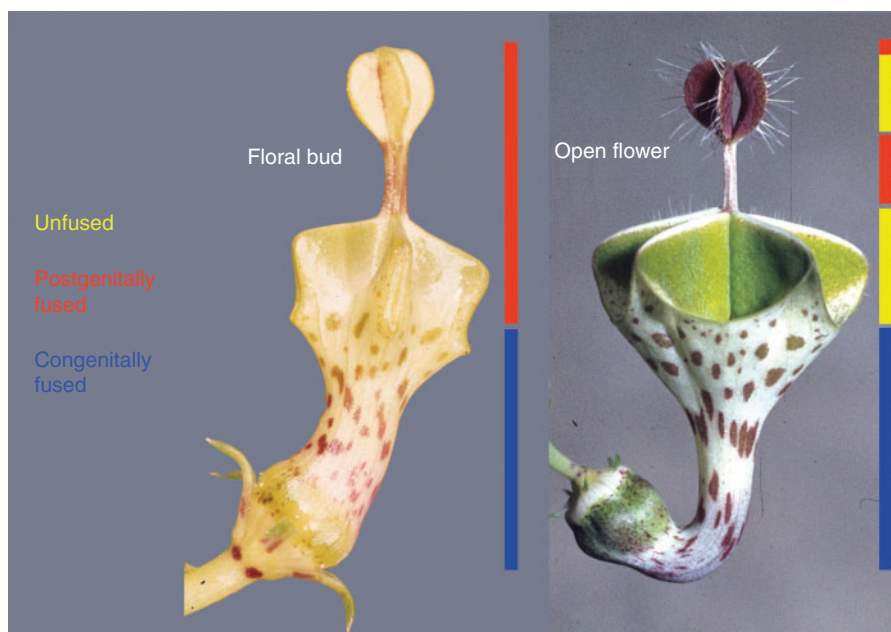


FIG. 10. Longitudinal differentiation of corolla and differential opening of zone of postgenital fusion of petals at anthesis in *Ceropegia distincta* (Apocynaceae-Asclepiadoideae). Left: floral bud. Corolla congenitally fused in the lower half, postgenitally fused in the upper half. Right: open flower. Corolla with five zones from base to top: (1) congenitally fused (floral tube), (2) open (five entrances with gliding zones for pollinators), (3) postgenitally fused (stalk of flag), (4) open (osmophoric flag), (5) postgenitally fused (upper end of flag).

The other end of the spoon, the ‘handle’, has a sticky pad underneath, which is attached to pollinators (Demeter, 1922; M. E. Endress, 2003). It has been assumed that pollinia evolved twice in Periplocoideae (Verhoeven and Venter, 2001) or at least three or four times (Ionta and Judd, 2007).

In Secamonoideae and Asclepiadoideae the pollinaria are attached to pollinators with a clip. In Secamonoideae, the translator consists only of this clip, and pollen is in tetrads within the soft pollinia (without pollinium wall). In Asclepiadoideae the translator has, in addition, two arms, and the pollinia are hard, having a pollinium wall, and pollen is no longer in recognizable tetrads within the pollinium (Verhoeven and Venter, 2001). Within Asclepiadoideae, only *Fockea* still has tetrads in the pollinia, which are soft and lack a pollinium wall (Verhoeven and Venter, 2001).

Another important innovation in Asclepiadoideae is a reorganization of the anthers. The dorsal pollen sacs disappear, and instead the now sterile anther flanks form the rigid (lignified) guide rails that direct pollinator body parts precisely into the clip of the translators (Kunze, 1996). In addition to this enhanced precision, the diversity of the depth of the guide rails adds to enhanced pollinator specificity (Fig. 8). Another consequence is that each pollinarium here consists of only two pollinia (in contrast to the four pollinia from the ancestral two pollen sacs per theca in Secamonoideae; Safwat, 1962).

The corona is ancestrally corolline, and thus develops on the corolla in alternipetalous positions (Fishbein, 2001; Kunze, 2005). This is not only the case in Apocynaceae but also in other Gentianales (Gentianaceae). In addition, there is a staminal corona in some Periplocoideae and in Secamonoideae and Asclepiadoideae (Rudjiman, 1982; Kunze, 2005). This also develops in alternipetalous position but is closely associated

morphologically and functionally with the androecium. A novelty here is that corona elements also develop in alternistaminal position. All elements together may form a complex nectar holder (and a holding device for nectar-seeking pollinators) of these revolver flowers (Liede and Kunze, 1993; *Asclepias*, Galil and Zeroni, 1965; *Hoya*, Kunze and Wanntorp, 2008). The consistency of the corona is then conspicuously firm. This is also true for the corolla in many Asclepiadoideae (Fig. 2A). Associated with such firm consistency is also a diversity of surface sculptures of corolla and corona (Ehler, 1975; Bruyns *et al.*, 2005). If the staminal corona is large and the five units are bulging, as in *Asclepias*, each of these convex, smooth and slippery units leads a pollinator leg exactly towards a guide rail (Fig. 7B).

Flowers of the most elaborate Orchidaceae

As in Apocynaceae *s.l.*, an impressive increase in synorganization can be observed following evolutionary trends through the family Orchidaceae. I will also begin with the most extremely synorganized clade, which constitutes the largest subfamily Epidendroideae (with 20 000 species; Chase *et al.*, 2015) and then show aspects of the evolution of this complexity. As in the derived subfamilies of Apocynaceae, in Epidendroideae the number of floral organs is also completely fixed, always with 3 + 3 perianth organs, 1 stamen and 3 carpels. The same is true in Orchidoideae and Vanilloideae. Orchidaceae with a single stamen are often called monandrous orchids (but they do not form a clade). In the two smallest subfamilies stamen number is higher: 2–3 in Apostasioideae and 2 in Cyrtipedioidae. In many of the derived Orchidaceae

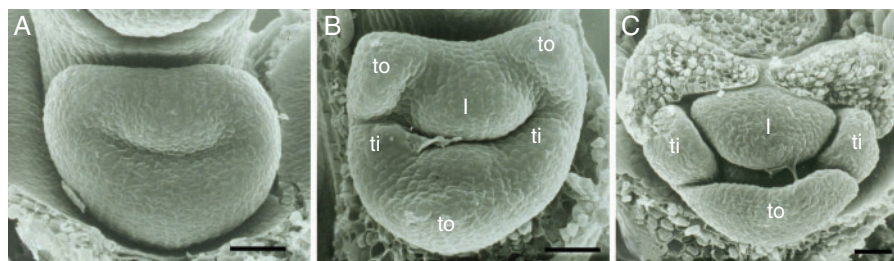


FIG. 11. Congenital fusion of all organs in very young flowers of *Oncidium ornithorhynchum* (Orchidoideae-Epidendroideae), scanning electron micrographs, flowers not resupinated, and thus lip located in upper part (modified from Endress 1994). (A) All six perianth organs congenitally fused, forming a ring wall, individual organs not yet visible. (B) Individual perianth organs visible: outer tepals (to), inner tepals (ti), lip (l). (C) More advanced stage, with lateral outer tepals removed, gynostemium hidden by median outer tepal and lip; abbreviations as in (B). Scale bars: all = 50 µm.

(Orchidoideae and Epidendroideae) with a single stamen, two developmentally early formed lateral outgrowths of the gynostemium are interpreted as staminodia; they may canalize the movements of the pollinators (Burns-Balogh and Bernhardt, 1985; Kurzweil and Kocyan, 2002). However, lateral appendages in some Orchidoideae may not be remnants of staminodia but late elaborations of the fertile stamen (reviewed by Kurzweil and Kocyan, 2002). In contrast to the polysymmetric Apocynaceae, the flowers of Orchidoideae are monosymmetric. The single stamen is in the single symmetry plane (Fig. 2B, D). All Orchidoideae have an inferior ovary.

The organs (tepals) of the two perianth whorls are congenitally fused, at least at the base, and also between the whorls. The three carpels are congenitally fused up to the top. Also the single stamen is congenitally fused with the gynostemium up to the top, androecium and gynostemium together forming a gynostemium (also called column). Note the difference between Orchidoideae and Apocynaceae: gynostemium (congenitally fused) vs. gynostegium (postgenitally fused). The perianth organ on the opposite side of the flower to the stamen, the lip, is commonly more elaborate and larger than the other five. It is part of the inner perianth whorl. It mainly functions as a landing platform for pollinators. In all Orchidoideae early floral development is remarkably similar. The flowers are pronouncedly monosymmetric from the beginning, and all six tepals originate as a conspicuous medianly compressed ring wall, without distinction of the single organs, and thus they appear congenitally fused before their tips become visible (Bletia, Kurzweil, 1987a, fig. 2C; Malaxis, Kurzweil, 1987a, fig. 4A, B; Dactylorhiza, Kurzweil, 1987b, fig. 1A–C; Prescottia, Kurzweil, 1988, fig. 4A, B; Listera, Kurzweil, 1988, fig. 6A; Phragmipedium, Kurzweil, 1993, fig. 4B, C; Oncidium, Endress, 1994, fig. 8.67.1–3; this study, Fig. 11; Satyrium, Kurzweil, 1996, fig. 3a; Pholidota, Kurzweil, 1998, fig. 6A; Hemipilia, Luo and Chen, 2000, fig. 1B, C; Amitostigma, Luo and Chen, 2000, fig. 3B, C; Gymnadenia, Luo and Chen, 2000, fig. 5B, C; Platanthera, Luo and Chen, 2000, fig. 5S, T; Telipogon, Pabón-Mora and González, 2008, figs 2D, 4D–F, 5B, E). Only in Apostasioideae is this early fusion less pronounced, but also present (Kocyan and Endress, 2001, figs 2B, C, H, I, N, O, 11B, G, L). Floral development of orchids has been reviewed by Kurzweil (1998) and Kurzweil and Kocyan (2002).

The complete congenital fusion between the stamen and the gynostemium ensures the immediate proximity of the androecial and gynoecial parts that contribute to the formation of the

pollinarium. Pollen of the anther is basically organized into four pollinia, one per pollen sac. In many groups the two pollen masses of the two pollen sacs in a theca form a single pollinium so that the pollinarium has only two pollinia (Rasmussen, 1986a), comparable to Asclepiadoideae (but where dimery results from transformation of the dorsal pollen sacs into sterile but mechanically reinforced parts). The stamen is in the symmetry plane of the flower and one of the three carpels is also in the symmetry plane and is adjacent to the stamen. The tip of this carpel is differentiated into a proximal stigmatic zone and a terminal secretory zone (viscidium) that comes into contact with the pollinia, and the secreted material acts as a glue to attach the pollinarium to a pollinator (Yeung, 1987a; Prutsch and Schill, 2000). The joint between the pollinia (coming from the androecium) and the viscidium (coming from the gynostemium) is a piece of disintegrated tissue from the anther (elastoviscin) with elastic properties (Dressler, 1986; Schill and Wolter, 1986; Wolter and Schill, 1986) in most Orchidoideae. However, in Cymbidoideae and Vandeae of Epidendroideae, this joint is a piece of tissue from the short tip of the median carpel. This carpel tip is called a rostellum. Depending on the site of origin the joint is called a caudicle (caudicula) (if from the androecium) or a stipe (stipes) (if from the gynostemium); there are even additional terms used in the literature (Rasmussen, 1982, 1985, 1986a, b). Pollinaria with a stipe also have a short portion of a caudicle adjacent to the pollinia. Thus, there is a combination of caudicle and stipe, whereby the largest part of the joint is made up by the stipe. The joint functionally corresponds to the arms of the translator in Apocynaceae. Conspicuous diversity is also exhibited by the proportions and directions of curvature of the gynostemium, which aid in attaching pollinaria to different body parts of different pollinators (Vogel, 1959; Garay, 1972; Rasmussen, 1982, 1986a, b; Burns-Balogh and Bernhardt, 1985; Manning and Linder, 1992). Dressler (1981) mentioned 13 areas for pollinarium attachment in euglossine bees.

Because of their often firm structure, the perianth organs, especially the lip (labellum), are extremely plastic in shape and diverse in details of surface differentiations (holding structures, secretory structures, three-dimensional surface sculptures of optical significance) (e.g. Davies *et al.*, 2002; Davies and Stipczyńska, 2006; Bradshaw *et al.*, 2010). The firm structure also allows the differentiation of a spur (another means of diversification is by spur diversity, e.g. Kurzweil and Weber, 1992; Micheneau *et al.*, 2009). Rarely, two collateral spurs are present (*Satyrium*, *Satyridium*, Vogel, 1959; Kurzweil, 1996) or

a complex spur in which the lip and neighbouring tepals participate (Vogel, 1969). Spurs on tepals of the outer whorl also occur but are less common than lip spurs (Vogel, 1959, 1969; Kurzweil, 1998). Optical, olfactorial and tactile devices on the lip are involved in different kinds of deception (food, brood site, sexual deceit, floral mimicry) of pollinators (e.g. Ackerman, 1986; Davies *et al.*, 2002; Cozzolino and Widmer, 2005; Schiestl, 2005; Jersáková *et al.*, 2006). Sexual deceit appears to be only known from orchids among angiosperms (Cozzolino and Widmer, 2005). The spur is involved in nectar production in many orchids. However, also here there is deception: many spurs do not secrete nectar. Nectar may also be produced from floral parts other than spurs. Loss and gain of nectar is plastic and may have occurred many times even within a genus (e.g. *Disa*, Hobbhahn *et al.*, 2013). A number of orchids have flowers that produce oil, either exposed on the lip (Vogel, 1974) or in a spur (Steiner, 2010).

Because of its often highly three-dimensional structure and sculpture, the lip of the orchids has sometimes been interpreted as a complex organ, a tepal fused with staminodia, beginning with Brown (1833). At present the tendency is rather to assume it to simply represent a tepal (Endress, 1994; Rudall and Bateman, 2002, 2004; Rudall *et al.*, 2013). However, a decisive answer is elusive because all outer floral organs are completely congenitally fused from the beginning (Fig. 11). In another, unrelated, clade of monocots, Zingiberaceae plus Costaceae, which also have a lip, the lip consists of two fused staminodia in the former, and of five fused staminodia in the latter (Kirchoff, 1988, 1997; Endress, 1995; 2–4 in Zingiberaceae, according to Specht *et al.*, 2012). As there is such divergence of lip formation within a clade of two families in Zingiberales, the question of whether the lip is homologous in all Orchidaceae should also be explored. Although molecular developmental studies are being performed in orchids, the specific question of lip homology has been difficult to tackle because in monocots perianth determination is somewhat different from eudicots (to which *Arabidopsis* and *Antirrhinum* belong) and because in orchids androecium and gynoecium are completely congenitally united into a gynostemium (Mondragón-Palomino and Theissen, 2008, 2009, 2011; Pan *et al.*, 2011; Mondragón-Palomino, 2013).

A speciality in some derived clades of Orchidaceae are extremely high ovule and seed numbers (689 000 ovules per ovary estimated for *Coryanthes senhasiana*; Nazarov and Gerlach, 1997). This is enabled by non-synchronous ovule development on large, convoluted surfaces. It is associated with an enormously high pollen number in each pollinium. In a study of eight species of eight genera, a range of 40 000–4 000 000 pollen per pollinium was calculated (the maximum among them for *Cochleanthus discolor*) (Schill *et al.*, 1992).

Coryanthes exhibits an extreme three-dimensional differentiation of the large, hanging flowers, in which the labellum takes part prominently; it is a genus of approx. 60 species (Gerlach and Schill, 1993; Fig. 9B). The lip is longitudinally differentiated into three conspicuous parts. The distal part has the shape of a hanging bucket. It tapers into a part bearing a helmet-shaped structure. The helmet is fastened with a horizontally directed stalk at the pendant floral base. The stalk is associated with two protrusions. The protrusions look like twin water taps (faucets), and secrete water drops, which at anthesis

continuously drip into the bucket, filling it with water. The helmet is a gland, secreting a perfume, which is collected by male euglossine bees for attracting females (Dodson, 1965; Vogel, 1966). The bees flying around the helmet and collecting perfume will *nolens volens* touch the water drops hanging from the taps with their wings and, when these get wet, immediately fall into the water-filled bucket. From there they find only one narrow exit at the morphological tip of the labellum, which is more or less obstructed by the tip of the rostellum. In struggling on their way out they either become the pollinarium of this flower attached to their back or, if they already have one attached to their body from another flower, they pollinate the stigma with it (Gerlach, 2011).

Stepwise synorganization of the flowers in Orchidaceae

As in Apocynaceae, phylogeny across Orchidaceae and some aspects of flower evolution have been intensively studied in recent decades (for a phylogeny of the family or subfamilies: Burns-Balogh and Funk, 1986; Cameron *et al.*, 1999; Cameron and Chase, 2000; Freudenstein *et al.*, 2004; Kocyan *et al.*, 2004; van den Berg *et al.*, 2005; Cameron, 2006; Carlswald *et al.*, 2006; Górniak *et al.*, 2010; Chase *et al.*, 2015; Freudenstein and Chase, 2015).

In the basal clade of Orchidaceae, Apostasioideae, there are no pollinaria, no pollinia, only an incipient gynostemium, floral monosymmetry is already present but not conspicuous, and flowers are only moderately complex. In some *Apostasia* species with superficially almost polysymmetric flowers, this feature appears to be secondary, as they are probably buzz-pollinated, and thus a special lip is not needed, whereas *Neuwiedia* has a lip and is not buzz-pollinated (Vogel, 1998; Kocyan and Endress, 2001). In addition, the androecium is monosymmetric in all Apostasioideae, also in early development; stamens are only formed on the developmentally abaxial side of the flower. In some cases stamen primordia also appear to be present on the adaxial side but do not develop into stamens (Kocyan and Endress, 2001).

In Apostasioideae the stamens (and staminode) are congenitally fused with the style for about half its length, and the style is shorter or longer than the stamens (Kocyan and Endress, 2001). In the other subfamilies (except for Cypridipodioideae), only the median stamen is formed. It is developmentally abaxial but becomes secondarily adaxial in almost all orchids, either by torsion (resupination) of the flower or, in epiphytic groups, by the hanging position of the inflorescence. By complete congenital fusion of the remaining stamen with the gynoecium, and adjustment of the length of these two components, the position of the anther and the median carpel tip become closely associated. This is a precondition for the evolution of the pollinarium.

Pollen is present as separate grains (monads) surrounded by pollen kitt or elastoviscin in Apostasioideae and Cypridipodioideae (Schill and Wolter, 1986; Pacini and Hesse, 2002; Pacini, 2009), rarely soft pollinia in Cypridipodioideae (Johnson and Edwards, 2000). Vanilloideae have monads or tetrads, rarely pollinia. Orchidoideae and Epidendroideae predominantly have pollinia. In some Orchidoideae the pollinia are 'sectile' and portioned into numerous massulae, which are loosely connected by elastoviscin. In a flower visit, part of the

massulae can be deposited on the stigma, and in this way a pollinium may be used for several pollinations (Freudenstein and Rasmussen, 1997; Johnson and Edwards, 2000; Pacini and Hesse, 2002; Pacini, 2009). Anthers with pollen not organized into pollinia or with pollinia with massulae occur in taxa with fewer ovules per flower.

The normal number of two or four pollinia per anther may be increased by septation of archesporial areas into eight, or this septation may be incomplete so that pollinia are incompletely divided (Hirmer, 1920; Freudenstein and Rasmussen, 1996). In Epidendroideae (and Vanilloideae), the rostellum with the stigma is bent down towards the lip and concomitantly the anther is also curved downwards ('incumbent') (Dressler, 1981; Freudenstein and Chase, 2015). In non-vandoids, curvature is at the base of the anther, whereas in vandoids it is in the sporogenous zone (Freudenstein *et al.*, 2002). A consequence is that in vandoids the sporogenous tissue acquires a convolute shape. This increase of the surface area of the sporogenous tissue per volume may be advantageous for the synchronous process of meiosis of the large and pollen-rich microsporangia. However, because of the curvature it is not always easy to determine the number of microsporangia in an anther from serial microtome sections (Hirmer, 1920; Freudenstein and Rasmussen, 1996).

Viscidia are not present in the three small subfamilies, Apostasioideae, Cyrtipedioidae (Johnson and Edwards, 2000) and Vanilloideae (Cameron, 2003b), but are present throughout in Orchidoideae and Epidendroideae. Generally there is one viscidium, but in some Orchidoideae and some Vandaeae there are two viscidia associated with two separate pollinaria, which are removed together or separately (Schill and Pfeiffer, 1977; Johnson and Edwards, 2000). In groups without viscidia, contact of the pollinator with secretion from the median stigmatic lobe may help in attaching pollen to the pollinator (Schick, 1989).

In some Orchidoideae, the viscidium is covered by a bursicula, a 'small purse', formed by the rostellum (Fig. 3B). The bursicula is pushed backwards by the pollinator, exposing the viscidium so that the pollinarium adheres to the pollinator. The viscidium is sometimes associated with a small plate (scutellum), which makes it more robust, and it is then called a retinaculum (Fig. 3B). This occurs especially in Orchidoideae and Epidendroideae. Elaborations of this region were studied in more detail by Schick (1988, 1989).

There is yet more complexity in the detailed structure of the pollinarium in Epidendroideae. In Cymbidieae and Vandaeae, in particular, the joint between pollinia and viscidium consists not only of caudicles but, in addition and for its main part, of a stipe. In flowers with elaborate pollinarium application mechanisms, stipes can perform forceful movements by precise deformation (Catasetinae, Romero, 1990), which would perhaps not be possible for simple caudicles. This is an enhanced way of synorganization between androecium and gynoecium. It requires a developmentally early bending of the anther, in contrast to pollinarium development with simple caudicles (Kurzweil, 1987a; Freudenstein *et al.*, 2002). This anther bending in flowers with stipes is congenital, i.e. the anther is incumbent from the beginning of development, whereas in other epidendroids bending occurs only late in development (Kurzweil, 1987a; Freudenstein *et al.*, 2002; Freudenstein and Chase, 2015). Stipes and early anther incumbence probably

evolved at least twice (Freudenstein and Chase, 2015). Additional synorganization in Epidendroideae also occurs between the lip and the gynostemium by extensive congenital fusion (especially in *Epidendrum*; with some 1400 species this is one of the largest genera of Orchidoideae).

A trilocular or almost trilocular ovary occurs in Apostasioideae, Vanilloideae and Cyrtipedioidae (Cameron, 2003a). However, it is unilocular and thus the carpels are more synorganized in Orchidoideae and Epidendroideae, as the presence of one unified locule allows more space than three separate locules for the development of an excessive number of ovules in ovaries of the same size.

Convergences in flowers of Apocynaceae and Orchidoideae: Fixed features

The advanced clades within Apocynaceae and Orchidoideae share many prominent features that are all connected with extreme synorganization, either as preconditions for or as results of synorganization or both:

1. stability of floral organ number,
2. highly regular floral symmetry (pentasymmetry in Asclepiadoideae, monosymmetry in Orchidoideae),
3. thick and firm consistency of floral organs (often plastic-like),
4. fusion of androecium and gynoecium (postgenital in Asclepiadoideae, congenital in Orchidoideae: gynostemium vs. gynostemium),
5. pollen aggregation into pollinia (Pacini and Hesse, 2002; Harder and Johnson, 2008),
6. two (or more) pollinia organized into pollinaria (with components from androecium and gynoecium),
7. pollinaria with a translator (at least partly secreted or of transformed tissue, i.e. elastoviscin) with an efficient apparatus to attach the pollinarium to a pollinator (glue or clip in Apocynaceae, glue in Orchidoideae),
8. attachment of pollinia to the translator with elastoviscin (Wolter and Schill, 1986; Dannenbaum and Schill, 1991; Pacini and Hesse, 2002),
9. number of pollinia per pollinarium two in all Asclepiadoideae and in many orchids (by a decrease of number of microsporangia per theca to one in Asclepiadoideae, and by synorganization of the two pollen sacs per theca into one in many Orchidoideae),
10. forward movement of pollinia by bending of the translator by desiccation after extraction from the flower, to reach optimal positioning of pollinia for application to a stigma, which is different from the position at extraction,
11. presence of floral guiding parts for exact positioning of pollinators (guide rails formed by anther flanks and shape of corona in Asclepiadoideae, floral monosymmetry and shape of lip and lateral outgrowths of gynostemium in Orchidoideae),
12. a consequence of the presence of pollinia is that the stigma becomes hidden in both families (in Asclepiadoideae the stigma is hidden in the guide rail, in some Orchidoideae it is a concavity below the rostellum; e.g. Dannenbaum *et al.*, 1989),

13. strong sectorial differentiation of the flowers: in Apocynaceae, this has led to the functional differentiation of one flower into five meranthia; these flowers can be visited and pollinated from five different sides; an exceptional pattern also occurs in an orchid: *Huttonaea pulchra* has two meranthia, and thus the two-spurred flowers can be pollinated from two sides; each side has a separate pollinarium (Steiner, 2010).
14. Very prominent are novel floral parts that converge in both families: pollinia, pollinaria with translator, corona (corona only in Apocynaceae).

Convergences in flowers of Apocynaceae and Orchidaceae: new flexible features: sources of diversification

Orchids and ascleps with their highly synorganized flowers have attained a high diversity within the confinements of their bauplan. Although this bauplan is remarkably fixed with regard to the number, position and fusion of the conventional floral organs, flexibility has arisen superposed on it. Pollinaria are extremely diverse in size and proportions in both families, and in asclepiads also in restrictions of the site in the pollinium where pollen can germinate, which differs from species to species (Schill and Pfeiffer, 1977; Schill and Jäkel, 1978; Johnson and Edwards, 2000). The guiding organs for pollinators, guide rails in asclepiads and spurs in orchids, vary greatly in depth. The firm texture of the corolla and corona in asclepiads and of the lip in orchids allows a multitude of surface elaborations for diverse tactile, optical and scent properties (Ehler, 1975, 1976; Schiestl, 2005). Pollinia, although they are key innovations in both families, are by no means uniform but are diverse in some groups, especially in Orchidoideae, where they may become portioned (sectile) or lose coherence in other ways (Freudenstein and Rasmussen, 1997; Pacini, 2009). Pollinaria (translators) are also diverse in asclepiads (e.g. Schill and Jäkel, 1978; Cocucci *et al.*, 2014) and orchids (Freudenstein and Chase, 2015).

Because of the fixed, precise position of the floral organs, progressive latitudinal and longitudinal floral differentiation with local separation of functions becomes possible. In basal Apocynaceae, there is no clear differentiation of the style head into a receptive area and an area with mere sticky secretion, but in more advanced clades there is spatial separation of stigma and translator secretion. Likewise in basal Orchidaceae, the receptive area and sticky part are identical and later in evolution there is separation of stigma and viscidium.

A highly specialized case of longitudinal elaboration in Apocynaceae is differential postgenital fusion in the corolla as described above for *Ceropegia* (Fig. 10), which has led to different functional zones of the petals and diversity within the genus. In Orchidaceae with longitudinal differentiation in shape and function, an impressive case is the lip of *Coryanthes* (Fig. 9B). In the two extreme cases, *Ceropegia* and *Coryanthes*, the perianth has become most diverse and inventive, after the formation of pollinaria was established and thus evolution proceeded to produce additional diversification at new levels.

In both families the novel floral organs or elaborated parts of old organs led to a confusing plethora of new terms (some of them redundant). In Apocynaceae this is especially the case for

corona elements and parts of the pollinarium (e.g. Bhatnagar, 1975; Bookman, 1981), and in Orchidaceae also for parts of the pollinarium and for the rostellum (Rasmussen, 1986a). A source of confusion may also be that the term *retinaculum* is used in both families, but with different meanings. In Apocynaceae it has been used to refer to the postgenital attachment area of the stamens to the style head, whereas in Orchidaceae it means a viscidium that is associated with a scutellum.

Changes in floral size in Apocynaceae and Orchidaceae, compared with less synorganized angiosperms

A general means of diversification in angiosperms is evolutionary change in flower size. It is interesting to see how flowers become miniaturized. The pathways are different depending on the degree of synorganization. In clades with a lower degree of synorganization, miniaturization occurs by a decrease of floral organ number and decrease of organ size. In highly synorganized flowers a decrease of organ number is not possible but there is more potential for a decrease of organ size in flowers with strongly fused organs. Congenital fusion of organs leads to a reinforcement of the architecture of the entire flower so that less tissue is necessary and may result in a more economical construction. An extreme case of each group, Asclepiadoideae and Orchidaceae, and an extreme case of an angiosperm clade with low degree of synorganization, Nymphaeales, may show this.

Among the smallest orchids are species of *Oberonia* with flowers 1 mm in diameter (Pridgeon *et al.*, 2005). In Asclepiadoideae, flowers of some *Tassadia* species are also 1 mm in diameter (Medeiros *et al.*, 2008). In both cases these flowers have the full set of floral organs (as described above for the two clades). The same set is also present in the largest flowers in the two families (*Stapelia gigantea*, up to 40 cm in diameter, Meve and Liede, 1994; and *Phragmipedium caudatum*, with inner tepals up to 75 cm long, Vogel, 1963). Also of interest is that in these cases what is increased are not the most synorganized parts (gynostemium or gynostegium) but the perianth organs, which are more independent and less functionally burdened.

In contrast, in Nymphaeales, although they exhibit about the same range of floral sizes, the extremes are reached differently. The smallest flowers are in *Trithuria* (Hydatellaceae), which are around 1 mm long (Rudall *et al.*, 2007), and the largest are in *Victoria* (Nymphaeaceae), which are up to 50 cm in diameter (Schneider and Williamson, 1993). To reach the miniature size of flowers of *Trithuria* appears only possible by stripping the flowers of all organs but one. *Trithuria* species are wetland plants with unisexual flowers (for a discussion of evolution, see Endress and Doyle, 2009). Even within the genus *Nymphaea*, a change in flower size is conspicuously linked to a decrease or increase in floral organ number. An example is *Nymphaea micrantha*, which produces two kinds of flowers. Flowers from dwarf plants, grown from bulbils, are much smaller and have many fewer organs than flowers from plants grown from seeds (Schmucker, 1932). The length of outer tepals is 72 mm in plants grown from seeds (21 mm in plants grown from bulbils). The difference in mean number of floral organs is as follows:

outer (sepaloid) tepals 4 (4), inner (petaloid) tepals 18 (11), stamens 111 (21), carpels 23 (8) (Schmucker, 1932).

Differences between Apocynaceae and Orchidaceae

The main differences in the evolutionary behaviour of flowers of Apocynaceae and Orchidaceae are due to the different symmetry: polysymmetry (pentasymmetry) in the former, and monosymmetry in the latter. However, because Apocynaceae are so highly differentiated into five sectors they could also be viewed as consisting of five monosymmetric modules in some sense and representing five meranthia. In both families the pollinators, predominantly insects, are forced into fixed positions for the pollination process by the specific floral architecture. These positions are exactly in the symmetry planes, five in Apocynaceae and one in Orchidaceae. In the elaborate revolver flowers of the Apocynaceae the body parts are guided by the corona elements and anthers into the five guide rails; in Orchidaceae the body is guided by the shape of the lip and the specific position of the attractive site, be it nectar or oil, often in a spur, or a dummy in deceptive flowers, and perhaps also by the two lateral appendages of the gynostemium.

Regarding diversity, according to Johnson and Edwards (2000, p. 243), 'The bilateral symmetry of orchids has allowed a greater degree of specialization in pollination systems and a much greater diversity in the morphology of pollinaria [than in ascleps].' Although orchids are uniform in having an attachment mechanism with glue, whereas in Apocynaceae there was an evolutionary change from glue to a clip, there is great diversity in the differentiation of the sticky part in orchids (see above, Schick, 1988, 1989). However, it may be added that the diversity of the translator shape of the pollinaria is higher in Asclepiadoideae than in Orchidaceae in some respects (Schill and Jäkel, 1978; Wiemer *et al.*, 2012; Cocucci *et al.*, 2014). In contrast to Orchidaceae, there are no spurs in Apocynaceae. In the latter the access to the flowers for different-sized pollinators is mechanically regulated by the general flower size, the size and shape of the corona, and specifically the depth and robustness of the guide rails.

Early flower development is different in the two families. In the highly synorganized clades of Apocynaceae, the perianth appears as separate sepals and petals (although the petals are congenitally fused; late sympetaly, for term, see Erbar and Leins, 2011), the androecium as separate stamens, and the gynoecium as separate carpels (Endress, 1994). Synorganization begins largely only after all individual organs are present. In contrast, in the highly synorganized clades of Orchidaceae, synorganization appears at the beginning of flower development by early congenital fusion of the organs.

Aspects of synorganization take place early in development, but the corona as a new organ appears late

The evolution of floral synorganization in Apocynaceae and Orchidaceae took place in the crown group of each family. Aspects of synorganization also take place at different times during floral ontogeny. Major episodes of ontogenetic synorganization occur early in development. There appears to be a tendency that they occur earlier in more derived clades than in

basal clades in both families. In contrast to the early expression of synorganization in floral development, the appearance of the corona as a new organ is late in ontogeny (Fig. 4).

In Asclepiadoideae, postgenital intercarpellary fusion takes place very early in flower development (Endress *et al.*, 1983, figs 9–12), earlier than in other Apocynaceae with a lower degree of synorganization (Walker, 1975, figs 4–8; Gomes *et al.*, 2008, figs 12–15). It is remarkable how the style head in Asclepiadoideae is transformed early in development from the genuine dimerous structure (the two carpels) to a pentangular structure, moulded by the neighbouring five stamens (Figs 2C and 6B). These five angles will be the five sectors where later the five translators are secreted. Thus, there is a change from two to five units in early ontogeny. Because of the very early postgenital fusion of the two carpels, this dimery is later no longer visible, except for the two main vascular bundles. Complexity obliterates the original morphological organ boundaries.

Particularly impressive is the early congenital fusion of all floral organs within whorls and between whorls in Orchidaceae (Figs 2D and 11A). In the basal Orchidaceae (Apostasioideae) fusion of the organs occurs later in the course of floral development (Kocyan and Endress, 2001). In some epidendroid orchids with particularly intimate synorganization of pollinaria by the addition of a stipe, the incumbence of the anthers is congenital (and not postgenital as in other orchids; Freudenstein and Chase, 2015). Thus, the connection between anther and rostellum is protracted in development.

Evolutionary sequence of synorganization and innovation in Apocynaceae and Orchidaceae

Several key innovations evolved sequentially from ancestral flowers to the highly synorganized flowers of orchids or asclepiads. These innovations did not necessarily appear in a straight line. It is to be expected that there was zig-zag evolution with gain and sometimes loss of features. Details are poorly known. However, there was a first time of evolutionary appearance for each of these innovations. The following are the putative sequences (of the first occurrence) of innovations.

Apocynaceae: fixation of pentamery → fixation of syncarpy (at least at the base of the gynoecium) → fixation of a single whorl in each organ category → sympetaly → postgenital fusion of carpel tips and of anthers with gynoecium → ontogenetic moulding of style head from two to five symmetry planes (→ translator number, nectary number, functional stigma number) → 4 or 2 pollinia per stamen → 5 pollinaria per flower (translators, including glue or clips, by secretion of 5 style head sectors).

Orchidaceae: fixation of trimery with two perianth whorls and two androecial whorls → congenital fusion of all tepals → lip differentiation → reduction of stamen number to 2 or 3 → complete syncarpy → reduction of stamen number to 1 → 4 or 2 pollinia per stamen → 1 or 2 pollinaria per flower (translator by tissue decay in the anther and – in some clades – by detachment of part of rostellum; viscidium by rostellum secretion).

Synorganization of organs by fusion also resulted in further stabilization of the position of organs and by the organs becoming more sturdy. In turn, it allowed further differentiation of the

organs along their length (longitudinally) or along their width (latitudinally), and also differentiation of a simple basic structure into two substructures, which assume different functions (progressive separation of functions, division of labour). For instance, in both families the former stigma differentiates longitudinally into a subapical receptive part and an apical part that produces the translator (among orchids in Cymbidieae and Vandeeae at least the viscidium of the translator; see, for example, Yeung, 1987b). In both families, the translator differentiates either longitudinally or latitudinally into a part for pollinator attachment (by glue or clip) and arms or joint connecting pollinia and part of pollinator attachment. In Apocynaceae (Asclepiadoideae), each theca with two fertile pollen sacs differentiates latitudinally into a fertile pollen sac and a sterile guide rail element. The corona differentiates longitudinally and latitudinally into a complex apparatus with several functions.

Orchidaceae with approx. 25 000 species are one of the two most diversified angiosperm families, and species and genus richness are especially concentrated in those subclades with the highest flower synorganization (Orchidoideae and Epidendroideae, approx. 21 500 species) (Chase *et al.*, 2015). Apocynaceae have almost 5000 species. Also here, the clade with the most synorganized flowers (Asclepiadoideae and Secamonoideae) have more species (3180) than all other subclades of the family together (Meve, 2002). It may be expected that flower synorganization (in addition to other traits) played an important role in the high diversification in both families, although different ages and different habitats of the families make a detailed direct comparison difficult.

OUTLOOK

Comparison of the flowers of the unrelated families Apocynaceae and Orchidaceae, which share a conspicuous convergence by having pollinia and pollinaria (unique in angiosperms), highlights the developmental and evolutionary preconditions for these traits. The flowers of the two families also show a large number of other convergences in detail. These are largely a precondition for, or a result of, synorganization. Whereas the synorganization of androecium and gynoecium and the structure of pollinaria have been studied in many representatives of the two families, the structure and diversification of the corona in Apocynaceae has been explored less well in spite of its extreme diversity.

There are additional effects of the synorganization of the flowers in Apocynaceae and Orchidaceae: because of their firm consistency, the flowers lend themselves to three-dimensional studies using scanning electron microscopy or tomography for morphometrics or morphospace studies because these flowers are expected to undergo less distortion than flowers with more delicate organs (van der Niet *et al.*, 2010; Gamisch *et al.*, 2013; Chartier *et al.*, 2014; Sedeek *et al.*, 2014). Thus far, tomography has been applied to orchids but not yet to Apocynaceae. Molecular developmental genetics has provided results for orchids as mentioned in the section on Flowers of the most elaborate Orchidaceae. Given the prominence of synorganization in both families and the emergence of novel organs (corona) in Apocynaceae, molecular developmental research regarding

organ fusion and boundary formation as studied in model organisms could be promising (e.g. Aida and Tasaka, 2006; Vandenbussche *et al.*, 2009; Lampugnani *et al.*, 2012; Žádníková and Simon, 2014; Zhong and Preston, 2015). There is great potential for interesting research topics.

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APPENDIX

Material used in this study is based on the following collections. The collection date is only mentioned if there is no collection number.

Acineta densa Lindl. (Orchidaceae-Epidendroideae), P.K. Endress s.n., 15.vii.1983 (not collected, only photographed), Botanic Garden, University of Zurich.

Asclepias curassavica L. (Apocynaceae-Asclepiadoideae), P.K. Endress 7368, Botanic Garden, University of Zurich.

Calotropis procera (Aiton) Dryand. (Apocynaceae-Asclepiadoideae), P.K. Endress s.n. (not collected, only photographed), Botanic Garden, University of Zurich.

Ceropegia distincta N.E.Br. (Apocynaceae-Asclepiadoideae), P.K. Endress 5210, Botanic Garden, University of Zurich.

Coryanthes macrantha (Hook.) Hook. (Orchidaceae-Epidendroideae), P.K. Endress s.n., 22.i.1996 (not collected, only photographed), Botanic Garden, University of Zurich.

Gomphocarpus fruticosus (L.) W.T.Aiton (Apocynaceae-Asclepiadoideae), P.K. Endress 7534, Botanic Garden, University of Zurich.

Oncidium ornithorhynchum Kunth (Orchidaceae-Epidendroideae), P.K. Endress 9759, Botanic Garden, University of Zurich.

Ophrys fusca Link (Orchidaceae-Orchidoideae), P. Voser s.n., 6.iii.1980, Corsica, France.

Vincetoxicum nigrum (L.) Moench (Apocynaceae-Asclepiadoideae), P.K. Endress 4690, Botanic Garden, University of Zurich.

Caralluma penicillata (Deflers) N.E.Br. (Apocynaceae-Asclepiadoideae), P.K. Endress 7516, Städtische Sukkulentsammlung, Zurich 81/1685.

Collected material was fixed and stored in 70 % ethanol. Material studied with a scanning electron microscope was critical point dried, sputter coated with gold and studied at 20 kV with a Hitachi S-4000 microscope. Material used for microtome section series was dehydrated and embedded in paraplast. Section series, 10 µm thick, were produced with a Leitz rotary microtome, stained with safranin and Astrablue, and embedded in Eukitt. Vouchers and permanent slides of the microtome sections are deposited at the Institute of Systematic Botany, University of Zurich (Z).