Trends and concepts in fern classification

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INTRODUCTION

One can only understand current classifications of a particular group of organisms after a thorough review of past systems. Classification of ferns has been particularly unstable in the past, and to understand why this has been so problematic we review the excellent article by Tryon (1952), in which the history of classification was discussed up until that time. Since that date, many other classifications have been proposed, some of which were discussed by Pichi-Sermolli (1973). We build upon these and provide an overview of our understanding of fern relationships and fern classification from the early 1970s to the present, addressing the influence of molecular phylogenetic studies in redefining genera and families. We will highlight some of the major changes in classification and identify genera that need further study to come to a better understanding of fern phylogeny. Many of these problematic genera were already suggested by Holttum (1973) to need monographic study. We summarize current knowledge of fern relationships in an annotated classification and point out where foci for future studies should be directed.
In commemoration of the 250th birthday of Carolus Linnaeus, Pichi-Sermolli (1958) provided an overview of higher classification of pteridophyta (understood to mean at that time ferns and ‘fern allies’). He stated that classifications until then had reflected four trends: (1) accepting four major lineages usually treated as classes: Psilopsida, Lycopsida, Sphenopsida and Pteropsida, the last including ferns and seed plants (e.g. Jeffrey, 1903; Arnold, 1948); (2) treating ferns and ‘fern allies’ as a group but excluding Psilopsida, treating the latter as related to Bryophyta (Lam, 1948) or as an independent lineage together with the fossil Psilophytopsida (Rothmaler, 1951); (3) maintaining Pteridophyta in their classical sense, including all vascular cryptogams (e.g. Campbell, 1940; Reimers, 1954); and (4) recognizing the independent divisions Psilophyta, Lycophyta, Sphenophyta, Pteridophyta and Spermatophyta (Benson, 1957). Pichi-Sermolli effectively dismissed trends 1, 2 and 4, concluding that all should be treated in a single division in the classical sense, placing them next to Spermatophyta. His classification scheme is discussed further below, but we highlight these four approaches here so they can be compared with the more detailed discussion of trends below.

The concept of family has always been fluid in ferns and, up to now, no clear family concept has been established. Families are generally defined as one or more genera that share several morphological characteristics, and many angiosperm families have had a long history of recognition based on shared floral and fruit features. However, in ferns, it has never been clear from what part of the plant these features should be derived, and this has led to a shifting number of families being recognized and differing circumscriptions for many of the more frequently recognized families.

Characters and terminology of ferns

Most readers will have a general idea of what a fern is, but a review of their typical characters and life histories will help focus attention on their key characters (those typically used in fern classification); this section of course can be skipped by pteridologists. There are several profound differences from flowering plants and mosses that define ferns and allow different characters to be used in the classification of this group. We therefore thought it appropriate to include a short introduction to the terminology of ferns and highlight the features important in their classification.

Ferns are vascular plants that produce spores and undergo an alternation of generations (with separate gametophyte and sporophyte generations that exist as free-living plants). Lycopsids are similar to ferns in this regard, but ferns are the sister group of the seed plants (gymnosperms plus angiosperms), whereas the lycopsids are sister to all other vascular plants (ferns plus the seed plants). Mosses, hornworts and liverworts (bryophytes) are also spore-producing plants with alternation of generations, but their sporophytes are reduced in size and dependent on the dominant gametophyte stage.

Alternation of generations. The leafy fern plant we observe in the fields, marshes and forests is called a sporophyte (Fig. 1B), a vascular plant that through meiosis produces spores with half the number of chromosomes found in the mother plant. When a spore lands in a suitable place, it grows into a free-living gametophyte (Fig. 1A), a haploid plant. Typically gametophytes are photosynthetic, but there are mycoheterotrophic gametophytes in some fern genera and families (Merckx et al., 2012). Gametophytes, often called prothalli, produce gametes through mitosis: male gametes, formed in antheridia, swim to the archegonia where female gametes are produced. The mobile, free-swimming, flagellate male cells are the reason for the water dependence of ferns; water is required for their movement from one gametophyte to another. Gametophytes can be bisexual or unisexual, and some can also reproduce vegetatively (a few may have lost the sporophyte stage altogether). From the fertilized female gamete, a new sporophyte develops. This life-history pattern differs from that in mosses, in which sporophytes are parasitic on the photosynthetic gametophyte, and from seed plants. In the latter, male and female gametophytes develop inside the pollen and ovule, respectively, with ovules being retained on the parental sporophyte until after fertilization, when at some point it is released as a seed with the new sporophyte partly developed inside. Spore and gametophyte morphology and the number of sperm flagellae have been used for classification of ferns. Alternation of generations in ferns was first described by Lindsay in 1794 (Fig. 2).

Sporophyte structure. A fern sporophyte consists of a stem, which is often called a rhizome, even if it is above ground or forms a 20 m tall trunk, as in some tree ferns. The rhizome can have various forms of stelar structures and orientation and scales, all of which are commonly used characters for identification and classification. From the rhizomes, leaves (also called fronds) grow, which in ferns are megaphylls (in contrast to lycopsids, which have a similar life cycle but bear microphylls: simple leaves with single unbranched veins). Leaves emerge as fiddleheads (Fig. 1C; circinnately coiled), although some groups have different types of vernation; circinnate vernation is not exclusive to ferns and also occurs, for example, in the carnivorous sundews, Droseraceae, and in the gymnosperm genus Stangeria T.Moore (Zamiaceae). The petiole, or stipe, is used as a diagnostic character and in particular the number of vascular strands is important. Leaf blades can be simple or highly divided, and this has in the past been used for classification; however, in most cases, leaf division does not apply above the rank of species. Fertile leaves bear spore-producing sporangia, which are typically organized in groups, called sori. Usually sori are on the lower side of the leaf, and some groups have specialized structures associated with their sporangia and sori.

Sporangia. Spores are formed in sporangia. In eusporangiate ferns, sporangia are formed from a group of cells, which is the plesiomorphic state. Eusporangia are found in all other vascular plants, except in the leptosporangiate ferns, where the sporangium develops from a single cell into a structure with a stalk, wall and spores. Leptosporangiate ferns form a clade that includes the bulk of fern species, but eusporangiate ferns are composed of several independent groups. In some families, most notably Marattiales, eusporangia are fused together, forming a complex synangium. The numbers of cells in the stalk of a leptosporangium and the shape and orientation of the annulus, the structure that splits to release spores, have been used as diagnostic characters.
Sporangia can be organized in various ways. Sometimes they cover the entire lower lamina of the blade, especially towards its apex, which is called an *acrostichoid sorus* (Fig. 1F). In most taxa, sporangia are organized in discrete sori (Fig. 1E). The typical leptosporangiate sorus consists of a stalk to which sporangia are fixed; a sorus can be covered with a scale-like structure called the *indusium* (Fig. 1E). The shape and organization of the indusium has been an important character for higher classification of leptosporangiate ferns and is still employed today, although usually in combination with other
characters. It was clear, even before the advent of molecular systematics, that there is a great deal of convergence in sorus characters, so on their own they have not reliably revealed a great deal about relationships.

Indument. Fern leaves can be glabrous or densely covered in hairs or scales. The shape and anatomy of these hairs are often used to diagnose species, but this is rarely useful at higher levels of classification. Scales of the rhizomes have been demonstrated to be conservative and phylogenetically informative, especially if clathrate scales are present (these are translucent like a stained-glass window).

Spores. Spore characters are useful in classification and phylogenetically informative, and they have also allowed calibration of DNA-based phylogenetic trees with the ages of fossil spores. Most remarkable are perhaps the heterosporous ferns. Heterospory is a condition in which a single sporophyte produces a smaller number of large spores, megaspores, that develop into female gametophytes and, more typically, numerous microspores that develop into male gametophytes. In ferns, this condition is only found in two groups of aquatics: Marsileaceae and Salviniaeae; it is clearly an adaptation to their aquatic environments, where the female gametophyte develops inside the megaspore and is thus protected from the surrounding water. These heterosporous ferns were often placed among ‘fern allies’, together with the unrelated heterosporous lycopsids (Selaginellaceae, Isoëtaceae), but heterospory has evolved independently in these lineages (and again in seed plants, which are also heterosporous) and is not necessarily associated with aquatic habitats in those lineages. Spores in heterosporous ferns are formed in special structures called sporocarps (Figs 1D, 4I). All other ferns and Lycopodiaceae are homosporous, the plesiomorphic condition.

With this explanation of terms, we hope to have clarified the various characters used in fern classification, which should make a historical overview easier to follow.

HISTORICAL OVERVIEW

Difficulties with artificial classification and ‘natural’ systems

As with all botanical studies, the nomenclatural history of fern taxonomy starts with Linnaeus (1753, 1754), who first attempted
to organize ferns based on the shape and position of sori on leaves; this use of a single character produced an artificial but easily used classification. Fern diversity and anatomy had been addressed in earlier studies (e.g. Celsalpino, 1583; Grew, 1682; Malpighi, 1675; Morison, 1699; Van Rheede tot Drakestein, 1678–1703; Plumier, 1705; Petiver, 1712), but it is important to note that at the time little was known about the life cycle of ferns. Linnaeus’s sexual system, based on the number of stamens and pistils (the so-called ‘sex organs’ in flowering plants), functioned well for artificially classifying angiosperms, but it was impossible to place ferns in a system based on numbers of male and female parts. Ferns were therefore placed in Cryptogamae, among mosses, algae, fungi and even some animals (corals, sponges, etc). In his Cryptogamia filices Linnaeus (1753, Sp. Pl. 2: 1085–1100) recognized 16 genera and 174 species: Equisetum L. (6), Onoclea L. (1), Ophioglossum L. (6), Osmunda L. (17), Acrostichum L. (25), Pteris L. (19), Blechnum L. (2), Hemionitis L. (2), Lonchitis L. (3), Asplenium L. (20), Polypodium L. (58), Adiantum L. (15), Trichomanes L. (11), Marsilea L. (2), Pilularia L. (1) and Isoëtes L. (1). All but the last are still classified as ferns; Isoëtes is a lycopod and does not have a close relationship to the other genera. Linnaeus’s classification was artificial, with species of clearly distant relationships placed together. This was the case for Linnaeus’ classifications in general, but it served the purpose of organizing the numerous newly discovered species until a more ‘natural’ system would be available. Species had to be assigned to a genus as a requirement in Linnaeus’s binomial system, which is the reason why so many fern species were often placed seemingly randomly in a genus. This seems illogical now, but it made sense at a time when ferns were biologically so poorly understood. In many cases, the author may not have known in which genus to place species, Presl’s (1836, 1845, 1851) also added to his system vegetative characters (especially venation, habit, and rhizome and petiole anatomy) in addition to the traditional fertile characters. Spore characters were also discussed and illustrated, but these were not used to group taxa. In total, Presl recognized 176 genera. Independent of Presl, John Smith (1841, 1843) compiled a classification, which, although differing in generic delimitation, essentially used similar characters to distinguish the 138 genera recognized. Both should be credited for establishing modern pteridology in the sense that they eschewed simpler methods of relying only upon soral and indusial characters to define a genus and instead emphasized that key generic characters could be drawn from any part of a fern. This was certainly an improvement in terms of putting together species that were morphologically and anatomically similar (in many cases), but it was much more difficult to use; to many of their contemporaries, Presl’s and J. Smith’s shifting emphases on soral characters to define one genus, petiole anatomy for another and venation for a third made their systems look arbitrary and flew in the face of the standard practice of the time.

However these classifications were criticized by Hooker, who initially accepted many of Presl’s genera (Hooker and Bauer, 1842), but later changed his opinion and only recognized 63 genera based on classical characters of the sorus; nonetheless, he clearly saw the value of these other characters and treated many of Presl’s and J. Smith’s genera at subgeneric levels. It appears that he attempted to provide a consensus between the large artificially delimited but well-established genera of Swartz (1801, 1806) and the newer treatments based on intrinsic characters that were considered ‘more natural’, the definition of which meant different things to different authors and changed over time, particularly once the concept of evolution was proposed. Hooker emphasized that characters of ‘fructifying parts’ were more important than characters of ‘vegetative parts’ forclassifying all plants at the generic level. This worked better with angiosperms, for which the concept of family was long established; many families had a long history of recognition, with a consensus of ideas about which genera were to be included. This sort of consistency of circumscription of the major angiosperm families is without parallel in ferns, and the emphasis on ‘fructifying parts’ for ferns did not result in either a consistent or a widely accepted system. Hooker’s authority, as the director of the Royal Botanic Gardens, Kew, led to the
until the end of the 19th century.

Although sporangium and sorus characters remained predominant as the basic framework for fern classification thanks to Hooker’s prominence, some interest in applying other characters to classification continued; Fé, who published the elaborate series Mémoires sur la famille des fougères (Fé, 1844–1873), which included Genera filicum (6th and 7th Mémoire), emphasized an even wider range of characters for generic delimitation than Presl and recognized approx. 188 genera. The first Mémoire dealt entirely with characters of ferns useful for classification and focused on venation. It also reviewed other characters, seeking new ones in fertile structures. He applied, for instance, characters of the spores and number of cells in the annulus. The last character was not taken up by subsequent pteridologists, but it was reapplied nearly a century later in the classification of Copeland (1947).

John Smith, who had become curator of the living fern collection at Kew, knew his plants intimately, and his Historia filicum (Smith, 1875) presented his views on classification and incorporated the genera of Presl and Fé, accounting for 212 genera. Even though the classification of Fé found followers in the French Empire, his and the classifications of Smith and Presl remained in the shadow of the more influential one of Hooker until the end of the 19th century.

Evolution enters the scene: a diminished importance of sori and sporangia as pre-eminent characters for classification

The classification of Christ (1897) emphasized the importance of vegetative characters, and this was adopted and expanded by Diels (1898–1900) in his treatment for Die natürlichen Pflanzenfamilien (German for ‘the natural plant families’). Diels presented a phyletic classification, a discipline in its early stages, and he applied the modern usage of family as a formal category, previously often referred to at higher taxonomic ranks (e.g. suborders and orders). The classification of Diels was heavily criticized by most subsequent workers, even though, or maybe because, it was the first classification to take Darwinian evolution into consideration.

Christ’s (1897) evaluation of previous systematic works gives us an insight into the perception of that work at the time. He wrote, for instance, about Die Famnkräuter of Schkuhr (1809) that the ‘nomenclature is outdated but that it contains many good images’, and of Fé’s Mémoires (1844–1873) that ‘it contains many species that were later not recognized, but that it contains valuable material, in particular the images’. In contrast, he cited Mettenius (1856) as the ‘founding work for a natural systematics’ of ferns, but that classification was only natural in a pre-Darwinian sense and the work of Mettenius was largely ignored outside the German language sphere. Christ clearly attributed the shift from a classification based on the sorus to a classification based on the entire anatomy of the ferns to Mettenius, which is only partly true (Presl and Fé had already applied this concept earlier), but Christ and Mettenius shared a nationality, which may explain the preference. Christ accepted 99 genera of ferns in 13 families and divided them into Leptosporangiatae and Eusporangiatae, possibly the first use of the character of sorus development as a primary character.

At the turn of the century, phyletic classifications of ferns were also being compiled elsewhere, the first being a treatment by the British botanist Bower (1889), who based his ideas on his elaborate studies on anatomy, development and morphology. In his three-volume work, The ferns, Bower (1923–1928) recognized 12 families, and in Polypodiaceae, the first major use of this now ubiquitously applied name, he recognized six evolutionary lineages. This phyletic classification was, however, problematic, mainly due to an emphasis on the location of sori on the leaf blade.

Copeland (1929), who attempted to arrange the East Asian genera into a phyletic sequence, was the first to address this problem and suggested that a larger Polypodiaceae would be more natural. The tree-fern lineages were included, not making Polypodiaceae natural in the Bowerian sense, but this wider concept of Polypodiaceae resulted in the name being variously applied ever since.

Christensen’s influential Index filicum (1906) listed 147 genera and followed almost exactly the treatment of Diels, except for some name changes to conform with the rules and an increase in number to account for new genera described during the six years between these publications. In subsequent supplements, the number of genera grew to 213. Christensen (1938) advanced from a bibliographer to a researcher in his own right, taking up the advances in Bowerian ‘phylogeny’, and – like Bower – he recognized 12 families, but divided his Polypodiaceae into 15 subfamilies, about which he stated that these should maybe be treated at the family level. His approx. 247 genera were based on a variety of vegetative (mostly hairs and scales) and soral characters, but several genera were pointed out to be of uncertain delimitation or status. Even though his classification was highly regarded and widely followed, Christensen himself regarded it as a tentative scheme.

Ching (1940) divided ‘Polypodiaceae’ into 33 families, grouped into seven ‘lines of evolution’ corresponding more or less to Copeland’s families (1929), in which the essential unnaturalness of ‘Polypodiaceae’ was accepted. Holttum (1947), at that time the dominant personality in fern taxonomy, did not agree with certain relationships expressed by Bower, and he presented a revised classification of Polypodiaceae in which five families were recognized, but he admitted that some are difficult to define morphologically.

Microscopic characters added to the mix

Copeland (1947) was one of the first systematists to propose that ‘valid’ taxa should reflect ‘naturalness’ and ‘convenience’, which by this time meant that taxa must correspond to a single evolutionary lineage and be well circumscribed, supported by characteristics and easy to define. He based his Genera filicum (1947) on extensive fieldwork in tropical Asia, and altogether he recognized 305 genera, many of which included just a single species. He divided Trichomanes but treated Asplenium in the broad sense, which was prescient. On the other hand, Holttum (1913) stated that ‘Copeland’s limitation seems to be his lack of understanding of Christensen’s discovery of the great importance of the details of hairs and scales as evidence of relationships’, which is understandable from a field botanist’s point of view. Microscopic characters are often difficult to observe in the field when all you have is a hand lens, especially...
when other characters such as habit are more obvious to distinguish groups. Copeland also provided a good overview of the concepts of fern classification preceding his *magnum opus*. He presented a similar number of taxa to *Ching* (1940), which Christensen stated had much to do with the two having the same generic concept. However, Ching still recognized 28 genera more than Copeland, so Christensen’s criticism was not entirely justified.

*Holttum* (1947) revised the classification of *Christensen* (1938), which resulted from a detailed study of Malayan ferns. It had been customary to place all leptosporangiate ferns in a broad *Polypodiaceae*, apart from a few easily recognizable families such as *Osmundaceae* (considered intermediate between leptosporangiate and euphorbiaceous ferns), *Cyatheaceae* (the tree ferns with a helicogyrate annulus), *Gleicheniaceae* (with a strange leaf morphology) and *Hymenophyllaceae* (the filmy ferns with leaves one cell layer thick and cup-shaped sori). *Christensen* (1938) also followed this, but stated that the subfamilies he treated in *Polypodiaceae* ‘were perhaps better dealt with as families’, which *Holttum* (1947) did. He redefined *Polypodiaceae*, including *Dipteris* Reinw., but excluding *Grammitidaceae*. He also segregated *Thelypteris* Schmid. from *Christensen’s* Dryopteridioideae into Thelypteridaceae, and the remaining subfamilies except Gymnogrammoideae, Vittarioideae and Onocleoidae were placed by Holttum in a large and diverse Dennstaedtiaceae with 11 subfamilies. The other group of tree ferns, Dicksonia, were also segregated, even though they were previously thought to be closely related to *Dennstaedtia* Bernh. Gymnogrammoideae was renamed Adiantaceae, and Holttum included Vittarioideae because they are undoubtedly related. Onocleoidae remained unplaced. Many matters remained unsolved in Holttum’s scheme, and he did not claim that all major groupings had been solved. Although it was easier to recognize a broad circumscription of *Polypodiaceae* and only segregate a few clearly divergent families, such as *Osmundaceae*, *Cyatheaceae* and *Hymenophyllaceae*, all authors of the time believed that this was unsatisfactory and most split *Polypodiaceae* into several families. Which characters should be emphasized and which families should be recognized was far from uniform, and substantial disagreements among authors were the standard.

*Starting from the beginning (again), fossils included*

*Tryon* (1952) concluded his historical overview predicting that ‘the conflict between utility and naturalness will remain an issue’ in future classifications. He suggested that taxa based on evolutionarily circumscribed groups would not necessarily have characters that are useful for identification, and indeed for some modern fern families this is a major issue today. Current examples of continuing contentious issues are the numerous families in the tree-fern lineage, segregation of *Lonchitis*, Saccoloma Kauf. and *Cystodium* J.Sm. from Lindsaeaceae (*Lehtonen et al.*, 2010), merging of Hymenophyllopsidaceae with *Cyathaea* Sm. (*Christenhusz*, 2009b), family placement of *Nephelepis* Schott (*Smith et al.*, 2006; *Hennequin et al.*, 2010) and difficulties with morphological circumscription of Pteridaceae, to name a few.

*Reimers* (1954) produced a thorough overview of *Pteridophyta* and included fossil taxa in his classification, setting it apart from earlier classifications that dealt only with extant or fossil taxa. He first defined his concept of *Pteridophyta* as ‘chlorophyllous, autotrophic land plants with antithetic (heterophasic) change of generations, and both generations organically independent’. In this way he included the lycopsids with ferns in a single lineage. Reimers provided a developmental scheme in which fern lineages were set out on a time scale, all essentially originating from Psilophytales (in which he included the fossils *Rhynia* R.Kidston & W.H.Lang and *Psilophyton* J.W.Dawson) in the early Devonian (320 million years ago). Modern Psilotales are directly derived from them in his scheme, which was the general idea at the time and one that prevailed until this century. We will not go into the classification of his fossil taxa here, although we admit that to come to a good classification of ferns, fossil lineages will have to be included. It is, however, not within the remit of this paper to compare fossil classifications with those for extant ferns. Reimers concluded that there were four extant lineages (as classes) in *Pteridophyta*: Lycopsida (Lycopodiaceae, Selaginellales and Isoetales), Psilotopsida (Psilotales), Articulatae (Equisetales) and Filices (Ophioglossales, Marattiales, Filicales, Marsileales and Salviniales). In Psilotopsida, two extant families were accepted, Psilotaceae and Tmesipteridaceae, because the two extant genera ‘appear to be relict forms’. What is interesting is that Reimers mentioned that the lack of roots in these genera cannot be seen as proof for any relationship with Psilophytopsida (which he placed in a different class, in spite of the linearity shown in the scheme). Articulatae included a great number of fossil taxa, but within this lineage *Equisetum* was placed in its own order because he stated that ‘relationships between this and fossil Calamites are not yet clear’. The largest group, Filices, were divided into four subclasses, one extinct (Primofilicales) and three with extant members: Eusporangiateae and Leptosporangiateae with Osmundidae as a ‘transition group’. Eusporangiateae included the obvious non-leptosporangiate orders Ophioglossales and Marattiales, the latter divided into five families, of which one is the fossil Asterothecaceae (which includes the giant trees, *Psaronius* B.Cotta), and which should still be accepted as separate from extant Marattiacae (*Christenhusz*, 2007). Leptosporangiate ferns were divided into three orders, Filicales, Marsileales and Salviniales, on the basis of heterospory of the last two orders; Filicales included a dozen families, of which the largest, *Polypodiaceae*, was divided into 14 subfamilies. Reimers, however, stated that *Polypodiaceae* would undoubtedly be polyphyletic in this concept, noting that Dennstaedtioidoideae closely resembled Dicksonioideae, that the relationship of Dryopteridaceae to *Polypodiaceae* was unclear and that the origin of the obviously primitive Woodsioideae and Onocleoidae required further study. We now know that if the heterosporous ferns and Parkeriaceae are included, this would make a monophyletic group (and could in theory be treated as a single family), and most subfamilies treated here correspond to many of the currently accepted families or subfamilies. Differences lie mostly in the placement of single problematic genera, e.g. *Nephelepis* in Davallioideae, *Pteridium* Gled. and *Stenochlaena* J.Sm. in Pteridioideae, *Diacalpe* Blume and *Peranema* D.Don in Woodsioideae (now part of *Dryopteris* Adans., see *Zhang* and *Zhang*, 2012), *Cyclosorus* Link, *Lomariopsis* Fée and *Thelypteris* in Dryopteridoideae and *Chieropleuria* C.Presl in *Polypodiaceae*. In spite of obvious misplacement of some genera, this work actually sets a good example for how the fossil record can be used to aid in placement.
of extant taxa, and it has therefore been unfortunate that this classification did not find wider acceptance.

Alston (1956), while preparing an account for the Flora of West Tropical Africa, came to disagree with the classification of Holttum (1947). He segregated Lindsaeaceae from Dennstaedtiaceae and maintained Vittariaceae as separate from Adiantaceae [although this had already been done by Reimers (1954) at the subfamilial level]. He also segregated Aspleniaceae, Aspidiaceae, Athyriaceae and Blechnaceae from Holttum’s Dennstaedtiaceae, and he described Athyriaceae and Lomariopsidaceae. The movement of genera by various authors between Athyriaceae, Aspidiaceae and Dennstaedtiaceae shows, however, that relationships among these genera were far from understood.

Pichi-Sermolli (1958) subdivided Pteridophyta into six classes, which were mixtures of living and fossil taxa with the two exceptions noted: Lycopsida, Sphenopsida, Noeggerathiopsida (fossil taxa only), Psilotopsis, Psilophytopsid (fossil taxa only) and Filicopsida. He used Filicopsida because the name Pteropsida was then often used in a broader sense to include all vascular plants. Within Lycopsida, he placed Isoetaeeae amid Lepidodendridae. The two genera in Psilotales were placed in their own families, and he stated ‘that these show no close affinity with the other groups of Pteridophyta and should be placed near the Psilophytopsida purely as a matter of convenience, since their derivation from them is very questionable’, a return to artificial classification in the absence of evidence of relatedness. Moving on to his Filicopsida, he divided this class into seven subclasses, of which Primofilicidae are extinct, and all others have living members. For his classes Ophioglossidae and Marattiidae, by most earlier authors joined in a single group ‘Eusporangiatae’, Pichi-Sermolli agreed that these two lineages differ greatly in morphology, and he dropped the terms ‘Eusporangiatae’, Pichi-Sermolli agreed that these two lineages ‘because the origin of the sporangia does not offer us a sharply taxonomical distinction’. In his opinion, these terms ‘should be used as descriptive morphological terms, not as names of taxa’. Osmundidae were maintained in a separate subclass because of their ‘antiquity and their paleontological history’. The living heterosporous ferns are placed in two subclasses: Marsileidae and Salviniiidae. Formerly placed together, the two were not considered to be related and were treated by Nakai (1949) as classes, but ‘it is clear that they are related to homosporous Filiciidae’. Within Filiciidae, Pichi-Sermolli agreed that delimitation and classification had been difficult and that pteridologists are very far from agreement about their classification’. The number of families had at that time increased due to splitting of the traditional Polypodiaceae (e.g. Ching, 1940; Holttum, 1947; Alston, 1956; Pichi-Sermolli, 1957), and this led to the suggestion they should be organized in orders, which Pichi-Sermolli (1958) did in a tentative scheme.

**Cytology enters the picture**

Prior to 1950, cytological aspects of ferns were practically unknown, and what was known was mainly due to the pioneering work of Manton (1950, 1954, 1958; Manton and Sledge, 1954) and Brownlie (1957, 1958); as cytological information became more widely available, it began to be applied in classifications. As more chromosome counts accumulated over the next decades, the position of certain species and wider affinities of genera, families and higher ranks could be postulated. By the 1970s, chromosomes of around 15% of fern species had been counted, and Walker (1973) provided a good review of how cytological information could help in unravelling previous problems. Manton (1958) demonstrating, for instance, that Pteridaceae of Copeland (1947) did not form a uniform lineage, with Pteris, Cheilanthes Sw. and Adiantum having chromosome numbers based on 29 or 30, and the other group (corresponding to the ‘dennstaedtioid group’) showing a wide range of basic numbers, even within a single genus. Chromosome numbers continued to play a role in classification for several decades, only to be replaced by molecular phylogenetics in the 1990s, providing important insights into relationships of species, genera and sometimes families. Mehra (1961) finalized the application of chromosome numbers and described phylectic lineages in terms of their cytological evolution, but this information was not included in later fern classifications. Cytological data on ferns are well documented and may in the future provide good additional insights into fern relationships and evolution.

The beginning of a new approach to fern classification, before DNA

In 1961, Wagner wrote an essay discussing the problems related to fern classification. He correctly concluded that due to the problems of assumed homology of morphological traits and associated convergent evolution abundant in ferns, classification had been unstable and the concept of a fern family had become meaningless. A few years later, Wagner (1969) addressed the question of how a system of relationships is constructed. He discussed non-evolutionary classifications and the then-modern phenetic systems of numerical taxonomy. Numerical taxonomy in which all information is included, he believed, could lead to circumscription of evolutionary groupings, but he stated that he was concerned about parallelisms and convergences and that in the case of ferns he did not have enough data needed to make an objective classification. He also made a significant and relevant statement: ‘In botany it may be that purely phenetic systems, no matter how detailed, will not produce results that coincide with evolutionary relationships. This possibility very much needs to be tested’. Wagner illustrated the subjectivity of phnetics in assuming homology of similar forms, when in fact they have evolved independently. As an example, he used evolution of heterospory, which, although appearing highly similar, certainly evolved several times. Of course this argument over phenetic and evolutionary classifications — the latter in which certain characters are weighted and exhibit a trend from primitive to derived — is still ongoing, and therefore the classification given in this review below is as much biased and subjective as previous ones, were it not that it is based on far more data than were available in the 1960s.

Similarly, Nayar (1970) criticized the classifications proposed in the 1940s and 1950s because they diverged substantially, which he blamed on the differing importance various authors gave to morphological characters. He proposed a new classification based on the ‘significant additions to our knowledge relating to several morphological criteria of the pteridophytes’, which he enumerated, but neither justified nor cited sources for this ‘knowledge’. In spite of his argument that subjective classifications are bad practice, his classification was also based on subjective interpretation of morphological characters. Even though the genera were better known than before, this again
resulted in movement of genera into different families and changes between family and subfamily names. He did attempt to place his families and subfamilies into an evolutionary scheme, although also here there are some odd placements. Plagiogyriaceae, for instance, are in his scheme derived from Osmundaceae, which is a strange placement for a group that has sporangia bearing remarkable similarities to those of Cyathea (Hooker and Bauer, 1842). Blechnaceae, to which Plagiogyria (Kunze) Mett. generally was thought to be related due to superficial similarity, are the only unplaced lineage in Nayar’s scheme in which there had been no attempt to justify postulated relationships.

Because Pichi-Sermolli (1970b) published a detailed catalogue of fern family names, Holttum (1971) felt obliged to comment on it and pointed out that ‘in the case of family names we are very far from being able to make a good taxonomic judgement’. Holttum stated that it is difficult to decide which family names would be the correct ones to use because ‘most family names of ferns have had such different meanings’ and ‘that such names are only intelligible if we associate them with the names of particular authors’. Holttum therefore regarded ‘all family names of ferns as informal and tentative’, a comment that may still be relevant today. Against this background, genetic studies of ferns were proposed, helping us understand processes of speciation and inheritance in ferns (e.g. Klekowski, 1971); this sort of publication did not involve such genetic studies, but rather hinted at the advantages of using a fern system rather than an angiosperm system for genetic analyses (due to their alternation of generations producing haploid gametophytes).

The view in 1972: fern classification is still a jungle for the user, unless . . .

In April 1972, a symposium was held in London bringing pteridologists from around the world together to discuss phylogeny and classification of ferns. In the resulting conference volume, Holttum (1973) stated that it would be impossible to come to a new classification without additional good monographic studies. He was of the opinion that ‘phylogenetic understanding will develop as classification improves’, but that classification ‘is still in a very imperfect state’. Nevertheless, Holttum proposed a tentative scheme, showing possible relationships, setting his understanding of fern inter-relationships apart from the schemes of Nayar and Kaur (1971) and Bierhorst (1971). The groups that Holttum suggested should be revised taxonomically [e.g. Adiantum, Asplenium, Cheilanthes, Ctenitis (C.Ch.) C.Ch., Diplazium, Dryopteris and Tectaria Cav.] are often groups that even today are still in need of taxonomic study, and some still need an improved generic circumscription.

In the same volume, Pichi-Sermolli (1973) provided an historical review of fern classification, in which he showed how classifications had changed over time and how unstable they had been. He concluded that no real progress in fern classification based on morphology could be made without the collaboration of palaeobotanists. He placed his hopes in numerical approaches due to their ‘being more objective’, which ‘will placate the fight between splitters and lumpers’. We now know that this was a forlorn hope and has not turned out to be the case, and arguments about splitting and lumping are still a major problem in taxonomy of all groups despite large amounts of genetic (DNA) data having been collected. Knowing the phylogenetic tree does not, of course, tell us how to delineate families or genera.

Mickel (1974) criticized the development of phyletic classifications over the preceding years, stating that circumscription was often ‘ill-founded and the relationships were often based on speculation rather than solid evidence’. Mickel stressed the need for proper evaluation of homologous characters, and he discussed some of the major issues of homology, for instance the different types of scales and development of stelar structures. He also pointed out the usefulness of stomata morphology to infer phyletic relationships. He then presented a freely drawn phylogram, which is backed by a discussion of characters, but, even though the evidence was sound, the choice of characters used to develop this phyletic classification was still subjective.

Crabbe et al. (1975) discussed the rearrangement of pteridophytes at the British Museum of Natural History and New York Botanical Garden with a large number of pteridologists and provided a new generic sequence for fern herbaria. They provided a complete list of genera, which was helpful in making sense of the numerous names proposed and used in herbaria. Even though many groupings do not seem natural, it reflected knowledge of fern classification at the time. They did suggest an alphabetical system as an alternative, which is easy for filing, but they pointed out the limitation in implementation of generic changes without major reorganization of the entire herbarium in that case. Placement of the heterosporous ferns was still uncertain, and they were therefore placed at the end of the sequence. Some genera were placed where we would not expect them based on their morphology (e.g. Plagiogyria among eusporangiate ferns). Despite its ambitions, this sequence never found much of a following in herbaria.

Development of fern classification and further study of the structures of ferns were summarized in Tryon and Tryon (1982) and Kramer and Green (1990). Kramer and Green (1990) was the most influential classification until the arrival of molecular phylogenetics, although it should be stated that at that time Polypodiaceae was generally taken in a broad sense, including most leptosporangiate lineages, and within that it was preferred to place genera alphabetically, a practice that is still followed in some books and herbaria. It may make it easy to find a genus quickly (if you happen to know in which genus a species is placed), but it can be difficult and laborious to update the herbarium when generic changes take place or to find a species when it has previously been placed in more than one genus.

Molecular systematics (DNA) and fern classification

The advent of molecular phylogenetics has rapidly changed our understanding of fern relationships through phylogenetic analyses of DNA sequence data. Studies of plastid DNA data slowly started emerging in the mid 1990s. Plastid data became mostly used initially in seed plants, and maternal, biparental (Sears, 1980; Corriveau and Coleman, 1988; Harris and Ingram, 1991) and paternal (Neale and Sederoff, 1988) inheritance of plastids is known in seed plants, but ferns still had not been thoroughly studied. Gastony and Yatskievych (1992) studied a natural hybrid of Pellaea Link with known parentage,
which demonstrated that plastid DNA is maternally inherited in this cheilanthoid fern.

In an early phylogenetic study, Hasebe et al. (1993) sequenced four ferns to infer the relationships of ferns to other land plants. Sequencing was still a laborious process, but, due to the development and improvement of in vitro DNA amplification [polymerase chain reaction (PCR)], a much larger number of taxa could be studied, and rbcL data for ferns quickly accumulated. Hasebe et al. (1994) developed effective primers to sequence rbcL in ferns; their paper focused on leptosporangiate lineages, and it showed monophyly of heterosporous ferns within the leptosporangiate clade. They identified all main lineages in ferns and showed that Dennstaedtiaceae were not monophyletic when lindsaeoids were included; they also found Vittariaceae resolved close to Adiantum. Possible changes in classification were suggested, but not formally proposed. Wolf et al. (1995a) sequenced 45 species of dennstaedtioid ferns, which produced similar results to the work of Hasebe et al. (1994), but here too no formal recircumscriptions of groupings were proposed because a much wider sampling of taxa was needed to assess monophyly of such a diverse group of plants.

A large pteridophyte symposium was held at the Royal Botanic Gardens, Kew, in memory of Holttum in 1995. This resulted in a volume called Pteridology in perspective (Camus et al., 1995) in which a wide range of topics in fern research were addressed, including several early molecular phylogenetic studies, family, genus and species concepts, speciation and systematics. In this volume, Pahnke et al. (1995) discussed the utility of the internal transcribed spacer (ITS) in phylogenetics of ferns. Wolf (1995b), who earlier discussed the usage of plastid sequences in fern phylogenetics (1995a), evaluated the use of a combination of genes in molecular phylogenetic studies. Comparisons of morphology with molecular phylogenetic results for tree ferns (Conant et al., 1995) and filmy ferns (Dubuisson, 1995) were also provided, increasing our understanding of these lineages. A discussion of the use of phylogenetic hypotheses for formal scientific classifications was presented by Hennipman (1995), and he stated that ‘modern higher classifications of ferns are a jungle for the user’, showing the great need to simplify fern classification. Hennipman discussed criteria used in the past for family and generic classifications but did not provide a clear-cut way on how to define a family in ferns; he emphasized lineages that he considered monophyletic and which ones were in his opinion clearly paraphyletic. He divided ferns into four orders, Marattiales, Ophioglossales, Psilotales and Filicales, and his order Filicales, which corresponds to the leptosporangiate ferns, is divided into 14 families (Aspleniaceae, Cyatheaceae, Davalliaceae, Dennstaedtiaceae, Dryopteridaceae, Gleicheniaceae, Hymenophyllaceae, Lindsaeaceae, Marsileaceae, Osmundaceae, Polypodiaceae, Pteridaceae, Schizaceae and Thelypteridaceae). The classification of Hennipman formed the backbone on which more recent classifications have been based because this simplified organization of the ferns was much more accessible than previous schemes. Of course with the molecular analysis of previously unsequenced genera, this simplified classification was destabilized and new families were again added.

The number of taxa studied increased when Hasebe et al. (1995) analysed rbcL sequences of 107 species. Their results showed that Grammitidaceae are embedded in Polypodiaceae, and Davalliaceae are the closest relative of the Polypodiaceae clade. Tectaria was found to be closer to Oleandraceae rather than to Dryopteridaceae, with which it was usually associated. Again, Dennstaedtiaceae were found to be polyphyletic, but this time with three lineages emerging; Lonchitis and Orthopteris Copel. fell in a clade separate from Lindera Dryand. Vittariaceae were found deeply embedded in Pteridaceae and strongly supported as sister to Adiantum. Tree ferns (including Metaxyaceae, Loxsomataceae and Plagiogyriaceae) were monophyletic and diverged early from the rest of the leptosporangiate taxa, just after the divergence of heterosporous ferns. No clear placement was found for Psilotaceae, but in one of their analyses (neighbor joining), Psilotum Sw. and Tmesipteris Bernh. were sister to Ophioglossaceae, a portent of what would be later found.

Although the techniques were still laborious, data for the major groups began to accumulate rapidly, and certain problematic fern groups were logically targeted using rbcL sequences, e.g. cheilanthoid ferns (Gastony and Yatskievych, 1992; Gastony and Rollo, 1995), dennstaedtioid ferns (Wolf, 1995a) and vittarioid ferns (Crane et al., 1995), helping to better understand relationships of genera in these groups.

Despite the molecular revolution in fern phylogenetics, morphology was not forgotten, and Smith (1995) summarized non-molecular ideas of fern relationships. He hypothesized that the eusporangiate families Ophioglossaceae and Marattiaceae constituted isolated lineages and that Osmundaceae, Schizaceae, tree ferns and a few other small families represent early branches off the main lineage leading to other leptosporangiates. This was based on fossil and living plant morphology and became the most broadly accepted classification. Morphologically most leptosporangiate families were recognizable, but inter-relationships among these were still unknown and widely disputed. Fern systematics had been diverted from producing phyletic schemes based on morphological characters, and thus these ideas were never robustly tested. Smith (1995) posed 16 challenging questions and invited readers to answer as many as possible. Many of these questions related to placement of genera with difficult to interpret morphology – Ceratopteris Brongn., Hymenophyllopsis K.I. Goebel, Monachosorum Kunze, Plagiogyria, Pleurosiopsis Fomin, Psilotum, Saccoloma Kauff. – or to relationships of entire large groups – Dennstaedtiaceae, Grammitidaceae, Hymeno-phyl laxaceae, Lindsaeaceae, Schizaceae, Thelypteridaceae, Vittariaceae and heterosporous ferns. Delimitation of Dryopteridaceae and other higher leptosporangiate fern families also needed sorting out because at the time their relationships were hotly debated. Hasebe et al. (1994) had already resolved some of these issues in their phylogenetic analysis of rbcL, showing Adiantum to be sister to Vittariaceae, Monachosorum sister to Dennstaedtiaceae, Lindsaeaceae sister to all higher leptosporangiates separate from other dennstaedtioids, Plagiogyria and Metaxy C.Presl in the treefern lineage, and the heterosporous ferns forming a lineage in leptosporangiate ferns (see also Fig. 3). The base of their tree did not have good support, but it was after all a first attempt at using DNA sequences to examine broader relationships within ferns; many relationships had to be left for resolution in subsequent studies.

Smith’s review (1995) soon led to analysis of combined molecular and morphological data sets. Pryer et al. (1995) presented the first cladistic analysis based on morphology combined with
Fig. 3. Summary phylogenetic tree showing relationships of a representative selection of fern genera based on molecular (DNA) data, modified from Schuettpelz and Pryer (2007), Lehtonen (2011), Rothfels et al. (2012) and Schneider et al. (2013).
Plastid 16S ribosomal DNA sequences were then applied to address the placement of ‘fern allies’. Manhart (1995) found *Psilotum* as sister to *Tmesipteris*, together forming a well-supported sister group to Ophioglossaceae. This relationship seemed odd at first, but it is now supported by morphological characters (Schneider, 2013). When the problematic lycophyceae *Selaginella* P. Beauv. was removed from their matrix (which perhaps had been causing some sort of spurious attraction of *Equisetum* to it), *Equisetum* also fell within the fern lineage but without strong support. Hasebe et al. (1995) further addressed the problems posed by Smith (1995) in an expanded rbcL phylogenetic analysis, which supported the sister relationship of Psilotaceae and Ophioglossaceae and showed Dennstaedtiaceae, Polypodiaceae, Pteridaceae and Dryopteridaceae in their traditional senses to be polyphylectic.

Placement of Hymenophyllopsis, a peculiar genus from the tepuis in the Guayana Shield, has always been problematic, and it was therefore often placed in its own family, Hymenophyllopsidaceae. The sorus structure is unique in ferns, although superficially similar to filmy ferns. Wolf et al. (1999) had shown that these are related to the tree fern genus *Cyathea*, with which they share scale and spore morphology. More recent studies have shown that *Hymenophyllopsis* is embedded in *Cyathea* (Korall et al., 2006, 2007), and thus it should not even be treated as a genus, let alone as a family or order. Christenhusz (2009b) and others now treat it as part of *Cyathea*.

*Lophosoria quadripinnata* (J.F. Gmel.) C. Ch., a common montane species in the Neotropics, was placed in its own family by Pichi-Sermolli (1970b), who removed it from the mosly extinct Protocatheceae, where it had been placed by Bower (1923–1928) and Reimers (1954). The relationship of this trunkless species to the tree ferns was never disputed, but Wolf et al. (1999) showed it to be most closely related to Dicksoniaceae.

Placement of *Equisetum* was evaluated again by Pryer et al. (2001a), who showed with strong support that it belongs to the fern lineage. In the process, they refuted the prevailing theory that horsetails and ferns form a transitional grade between lycophytes and seed plants. As a replacement name for the ferns including Equisetaceae, Kenrick and Crane (1997) had proposed the infradivision Moniliformes, an invalid name under the current nomenclatural code (McNeill et al., 2012). The term ‘monilophytes’, loosely based on this infradivision, became established as a result of the paper of Pryer et al. (2004), in which phylogenetic relationships of ferns were inferred. Why this new term not based on a generic name is useful in the context of a cladistic study is unclear to us. Nevertheless, it was quickly picked up by the community and rendered into the vernacular as ‘monilophyte’ (e.g. Pryer et al., 2001a, 2004; Schneider et al., 2004; Schneider, 2013). ‘Moniliform’ means ‘bead-shaped’, but it was never stated to what part of a fern, or to which fern, this refers, although it was suggested that this might refer to the stelae structure of some extinct Devonian taxa. It seems that a new colloquial name has been coined for no particular reason other than providing a scientific-sounding synonym for the word ‘fern’; in fact, it is often used in parentheses after the word fern, so ‘monilophyte’ is superfluous. It is an alternative to the word ‘pteridophyte’, which traditionally included all ferns plus lycophytes; thus, an alternative, similar sounding word was coined, without taking notice of its etymology: ‘bead-plant’ being as uninformative as ‘wolf-plant’ or ‘lycophyte’, which should correctly be called clubmoss (in the English vernacular) or lycopod (based on *Lycopodium* and referring to the vernacular ‘wolf-claw’ of many languages).

Psilotaceae were not included in the study of Kenrick and Crane (1997), but Pryer et al. (2004) included these to show they also belong to the fern clade. Critical taxa were included in this data set and additional plastid (rbcL, atpB and rps4) and nuclear (18S rDNA) regions were added to those previously used. As expected, Osmundaceae were sister to the leptosporangiate ferns. Hymenophyllaceae – which had recently been shown to consist of two monophyletic groups (Pryer et al., 2001b) – was found to be sister to the gleichenioid ferns, and tree ferns and heterosporous ferns were shown to belong to ‘core leptosporangiates’.

In a high-profile paper, Schneider et al. (2004) studied the age of fern diversification and concluded that the higher leptosporangiates, to which the majority of extant ferns belong, diversified in the Cretaceous, most probably in response to diversification of the angiosperms. The hypothesis is that, because of the increase in diversity of angiosperms, numerous new niches appeared in which ferns could diversify. Schneider et al. (2004) also introduced the term ‘eupolypods’, defined as the polyads excluding ‘basal’ polyads (e.g. Dennstaedtiaceae and Lindsaeaceae) and pteridoids (Pteridaceae). The eupolypods were found to be represented by two clades (numbered I and II), on which we will elaborate below. This showed that many fern lineages are of similar age to or younger than many angiosperm families, refuting the usage of clade age as a useful character for defining a fern family.

Classification appeared to be more stable after publication of the fern classification by Smith et al. (2006), which summarized molecular findings to that date and provided synapomorphies for the accepted families. Schuettpelz et al. (2006) added plastid atpA to their earlier fern data set to gain better support at deeper nodes. They continued their work on fern phylogeny and performed a global analysis to serve as the basis for addressing large-scale evolutionary questions (Schuettpelz and Pryer, 2007). They assembled and analysed the most inclusive molecular data set to date with three plastid regions (rbcL, atpB and atpA) including 400 leptosporangiate fern species. Their findings were of great value for the placement of genera that were previously unavailable for molecular study and were dependent on the generous contributions of many fern collectors.

Meanwhile, a number of families were studied in greater phylogenetic detail, resolving many taxonomic problems (e.g. Korall et al., 2006, 2007; Nagalingum et al., 2006, 2007; Christenhusz et al., 2008; Hennequin et al., 2008; Rothfels et al., 2008). New findings were incorporated in the updated classification of Smith et al. (2008), although this did not substantially differ from his classification published two years earlier.

Schneider (2007) tested the congruence of molecular and morphological data with the idea of incorporating fossil taxa in phylogenetic studies, which is currently being further developed.
Morphological data have since been combined with molecular data in analyses with some important insights (e.g. Schneider, 2007; Lehert et al., 2009; Lehtonen et al., 2010; Sundue, 2010; Sundue et al., 2010).

In a new edition of A. Engler’s Syllabus of Plant Families, Fischer (2009) adopted findings of recent molecular analyses and treated Lycophytina as separate from Euphyllophyta. Within the latter, he accepted super-class ‘Moniliformopses’ of Kenrick and Crane (1997), comprising all ferns, Equisetaceae and Psilotaceae. A synopsis was provided on classification of this group, and it included all fossil taxa, which in many cases were placed in families consisting of exclusively fossil taxa. We did not check all fossil families, but we noticed that nomenclatural priority of family names was not always taken into account. For the extant taxa, it simply followed the classification of Smith et al. (2006), although here and there with some reservations and occasionally following some subclassifications that had already been shown to be incorrect. The acceptance of subfamily Stenochlaenioidea as separate from Blechnoidea, for instance, is inappropriate if Woodwardia Sm. is included in either of them.

The classifications of Smith (2006, 2008) initiated a discussion about morphology being incompatible with the molecular results, and indeed some groups such as Hypodematiaceae, Lomariopsidaceae, Pteridaceae and Woodsiaceae (sensu Smith) were difficult to define morphologically. Therefore, an analysis with morphological data alone was carried out (Schneider et al., 2009), and this recovered the same deep phylogenetic relationships (recognizing the four major clades in ferns: Equisetidae, Ophioglossaceae, Marattiidae and Polypodiidae) found in previous studies of DNA sequence data. It would appear that molecular data and morphology are compatible, provided they are treated in similar ways.

To provide a guide for herbaria to organize their specimens in a phylogenetic way, a linear classification for all vascular plants was composed of which here the lycopods and ferns are of consequence (Christenhusz et al., 2011). This linear sequence was not intended as a new classification and followed in many ways Smith et al. (2006, 2008), but major changes were made in the treatment of some ‘eupolypods II’, with the disintegration of Smith’s polyphyletic Woodsiaceae and with descriptions of Diplaziopsidaceae, Rhachidosoraceae and Hemidictyaceae (Christenhusz and Schneider, 2011; Christenhusz et al., 2011). Also, placement of Equisetaceae as sister to all ferns seems to be better than as sister to Marattiidae (as in Smith et al., 2006). The position of Equisetum is poorly supported but always among the deeper nodes of the fern clade, and therefore it could end up in various places, depending on the method of analysis or data set used (Lehtonen, 2011). Placement as sister to all other ferns is also in agreement with palaeontological studies that show its early appearance in the fossil record (Taylor et al., 2009).

The classifications of Smith et al. (2006, 2008) and Christenhusz et al. (2011) mostly reduced the number of genera, resulting in an expansion of several (e.g. Asplenium, Blechnum, Cyathea and Hymenophyllum Sm.), but also resulted in the acceptance of narrower generic concepts in other groups (e.g. Hymenophyllaceae, Poly podiaceae and Pteridaceae). These classifications are, however, remarkable in that they combine morphological and molecular data to come to a consensus classification, aiming to define monophyletic families and genera with synapomorphies.

Some of the notable discrepancies are due to the traditionally broader generic concepts used by workers in the Neotropics compared with those in tropical Asia. Now we can combine the data on these taxa and come to a consensus classification resulting in evolutionarily defined natural genera and families. With the wealth of data currently available (e.g. Eiserhardt et al., 2011; Lehtonen, 2011; Lehtonen et al., 2012; Rothfels et al., 2012; Zhang and Zhang, 2012; Liu et al., 2013; Schneider, 2013; Schneider et al., 2013) it is now possible to compare generic concepts across continents, applying a global concept of family and genus.

DISCUSSION AND SOME HIGHLIGHTED GROUPS

Concepts of higher classification of ferns and lycopods have changed considerably throughout history, and we therefore discuss a few groups with regard to their previous and current delimitation.

Fern allies have previously been loosely defined as vascular spore-bearing plants that are not ferns. They have variously included the lycopods (Hyperzia Bernh., Lycopodiella Holub, Lycopodium, Isoëtes and Selaginella), horsetails (Equisetum, Fig. 4A), whisk ferns (Psilotum and Tmesipteris, Fig. 4G), Ophioglossaceae (Fig. 4B), water ferns (Marsileaceae, Fig. 4I, Salviniaceae and Ceratopteris) and sometimes even members of Schizaceae and Gleicheniaceae. Of course this was never considered a natural group but just included genera that according to subsequent authors could not be placed among the ferns proper for various reasons. Because of its vague circumscription and now evident non-monophyly, the term ‘fern ally’ should be avoided. In a similar way, the term ‘eusporangiate fern’ is to be avoided because this includes ferns that are not leptosporangiate; it just describes the plesiomorphic state (eusporangia are also present in seed plants and lycopods) and the taxa included do not form a clade.

The water ferns (Salviniaceae, Fig. 4I) were previously placed among ‘fern allies’ because they are heterosporous like Selaginella and Isoëtes. They have, however, been found to be sister to the polypods, so they are deeply embedded in the homosporous fern clade. Heterospory in general evolved several times independently in lycopods, ferns and seed plants, but this evolution is not necessarily the result of adaptation to an aquatic life form, although it appears as a plausible association for heterosporous ferns.

Even though it is not a new concept (Linnaeus, 1753, already included Equisetum in his ‘Cryptogamae filices’), the term ‘monilophyte’ was coined (Pryer et al., 2001a) for the fern clade including Equisetaceae and Psilotaceae, in contrast to the ‘lycophytes’, both of which are linguistically erroneous and superfluous terms. This clade has been traditionally called pteridophytes, but it has been suggested that ‘pteridophyte’ cannot be used because in some classifications it has included the lycopods.

However, the origin of the term ‘monilophyte’ is unclear and has never been published as a formal taxon and, moreover, its etymology is obscure. ‘Sphenophytes’ (a term commonly used for extant horsetails and their fossil relatives) share a common ancestor with ferns, even though their exact placement within the fern lineage is still uncertain. It is therefore preferable to use
the term ‘fern’, which as noted above has in the past often included Equisetaceae and Psilotaceae, although alternatively the early branching lineages of the fern clade could be treated as independent lineages (‘sphenophytes’, ‘psilotophytes’, ‘marattiophytes’), although this does not reflect their membership in a clade with the rest of the ferns.
To illustrate the changing ideas of how ferns should be classified, we focus below on a number of groups and genera and describe the variation in opinions that have existed about how best to handle their taxonomy.

**Leptosporangiate ferns** are a natural group, forming the crown group of ferns including the vast majority of species. In the past, most of these have at one time been included in a single family, *Polypodiaceae* (or variants such as Filices or Dennstaedtiaceae, variously including Osmundaceae, tree ferns and gleichenioid ferns), resulting in great variability in application and circumscription of this family. The concept changed from including the majority of the leptosporangiate ferns to the most recent concept of those having scaly creeping rhizomes with abaxial (rarely marginal), rounded to elliptic, elongate or acrostichoid concept of those having scaly creeping rhizomes with abaxial expanding the description above to include taxa occasionally having green spores and a petiole not always cleanly abscising (as is the case with most other polypods; *Smith et al., 2008*).

Generic classification of graminitid ferns has been unstable during the last 20 years, and they are currently divided in a great number of genera, even though they form a subclade within Polypodiaceae. It would be better for general users if these are treated as a single genus (with approx. 700 species, making it a large genus, the size of *Asplenium*, but smaller than *Elaphoglossum* or *Thelypteris*) and treat most currently recognized genera at the subgeneric level. Polypodiaceae as a family should probably be expanded again to include all families that have been treated as subfamilies, allowing movement of genera between them without altering their family placement. Not all genera in *Cyatheaceae* are tree like; they include trunkless *Plagiogyria*, *Metaxya* and *Loxsoma* R.Br. They also include *Hymenophyllopsis* (Lellinger, 1984), a group of small ferns from the Guayana Highlands in South America. This genus was difficult to place on a morphological basis and was thus placed in its own family or even its own order. It is now known that these are embedded in *Cyatheaea* (*Korall et al., 2007*), and combinations in *Cyatheae* have been made for all eight species (*Christenhusz, 2009*).

**Lonchitis** has also been variously treated in the past, the genus previously including many species, but now reduced to two (most of the remaining species were moved to *Bliotilia* Tryon, *Dennstaedtiaceae*). Placement of *Lonchitis* has been contentious in recent times. Earlier it was placed in *Polypodiaceae* subfamily *Dennstaedtioideae* or *Dennstaedtiaceae*, but *Smith* (2006) placed it in *Lindsaeaceae*. This makes the last difficult to define morphologically; hence, it was placed in its own family *Lonchitidaceae* (Schomburgk, 1848; Christenhusz, 2009a; *Lehtonen et al., 2010*; *Christenhusz et al., 2011*), a family placed near *Saccolomataceae* and *Cystodinaeae* among the early branching polyponds, even though the deep nodes of the polyponds are not well supported in most studies (*Lehtonen et al., 2012*).

**Lomaria** Willd. has, since its publication, been widely accepted, so much so that *Smith* (1875) stated that it is a natural genus containing a considerable number of species of great uniformity of habit. *Hooker* (in *Hooker and Bauer, 1842*) agreed with this to a certain extent, but he pointed out that a number of species, including *L. spicant* (L.) Desv. ‘vacillate between Lomaria and Blechnum . . .’, but such or similar gradations are common to other genera of ferns, and if made an excuse for abolishing a long-established one, the number of genera would undergo a great reduction*. *Hooker* further discussed inclusion of *Plagiogyria* in *Lomaria*, and he pointed out that there is an ‘affinity of those species with *Cyatheae*’ (to which we now know *Plagiogyria* is related), especially in characters of the ‘capsules’ (the annulus of the sporangium is oblique, e.g. helicogryrate), ‘yet the habit and sori are so entirely in accordance with Lomaria that’ the genus *Plagiogyria* is difficult to identify. He further discussed differences in the number of genera accepted in this group according to other authors. *Mettenius* (1857), who excluded *Plagiogyria*, stated that several species depart from the pteridoid character and completely merge into *Blechnum*, such that *Blechnum* and *Lomaria* are no longer separate, whereas *Presl* (1851) kept them separate and divided them into a multitude of genera based on minor characters. Some of these genera are still in use today, but recent phylogenetic studies (*Lehtonen, 2011*) have shown *Salpichlaena* J.Sm. and *Stenochlaena* (which had been separated from other genera on the basis of their climbing habit) to be
embedded in Blechnum (specifically related to B. serrulatum Rich.), which should also include Brainea J.Sm., Doodia R.Br., Pteridoblechnum Hennipm. and Sadleria Kauff. Lomaria was based on the Australian L. nuda (Labill.) Willd., and Blechnum is based on the Neotropical B. occidentale L.; the European species that has characters of the two was placed in Spicantia C.Presl, an intermediate genus created to solve the problem of intergrading morphology. The current consensus is to apply a broad concept of Blechnum, but further studies on the evolution of Blechnaceae are needed to assess which, if any, of the segregate genera should be maintained.

Acrostichum was a genus applied initially to all ferns with sori that were acrostichoid (Fig. 1F), i.e. distributed as a solid mass across the back of the frond rather than organized in separate, discrete structures. We know today that an acrostichoid fertile lamina can be formed in a variety of ways, and thus it is not useful for classification, although it is definitely a distinctive structure. Therefore, ontology of this sorus type is more important for the placement of these ferns than its product. Initially genera such as Blechnum, Ceratopteris, Danacea Sm., several species of Asplenium, Bolbitis Schott, Tectaria, etc. were included, but it was soon realized that these confluent sori were covered by indusia at least at the youngest stages, or, as in the case of Danacea, they were fused sporangia with pores on the apex (Christenhusz, 2010). It even included genera as different from each other as Actiniopteris Link, Anetium Spligt., Anogramma Link, Cheiropleuria, Hemionitis, Platycerium Desv., Polybotrya Willd., Schizaea Sm. and Todea B. Rich. It was Schott (1834) who separated Elaphoglossum Schott from Acrostichum, which was followed by Moore (1857) who provided numerous new combinations. This process was completed by Christensen (1906), after which Elaphoglossum was widely accepted and Acrostichum (with A. aureum L. as the type species) achieved its modern sense and was restricted to a few species of mangrove ferns.

The name Thelypteris was coined by Schmidel (1763) based on Acrostichum thelypteris L., but the name Thelypteris had already been published in the same year by Adanson (his name is a synonym of Pteris L.), making Schmidel’s name illegitimate. This caused confusion, and the name was not accepted by subsequent authors. Pichi-Sermolli (1953) discussed this nomenclature, and Thelypteris of Schmidel was subsequently conserved (Holttum, 1968), resulting in the transferral of numerous species, mainly from Aspidium Sw., Dryopteris and Nephrodium Rich., to Thelypteris. It was soon recognized as a natural group and broadly accepted. Thelypteridaceae as a family were published [including related genera such as Cyclosorus, Macrothelypteris (H.Ito) Ching, Meniscium Schreb., Phegopteris (C.Presl) Fée, Oreopteris Holub, etc.] by Pichi-Sermolli (1970c), and molecular studies soon showed five subgroups in this family, resulting in five genera being recognized as comprising it, although it may be better to accept only three at the generic level (see classification below).

Eupolypods form two subclades often called eupolypods I and II. The first includes the families Davalliaceae, Dryopteridaceae, Hypodemataceae, Lomariopsidaceae, Nephrolepidaceae, Ölandraceae, Polypodiaceae and Tectariaceae (sensu Christenhusz et al., 2011). These families are morphologically divergent, and it is difficult to see them as a single taxonomic entity, especially if Dryopteridaceae and Polypodiaceae sensu stricto are compared; nevertheless, they share characters of soral structure, which are microscopic, and in the vascular structure of petioles (generally an arch of vascular strands in a U-shape, except in Hypodemataceae). The second lineage (eupolypods II) includes Aspleniacaeae sensu stricto, Athyriaceae, Cystopteridaceae, Diplaziosidaceae, Hemidictyiaceae, Onocleaceae, Rhachidosoraceae, Thelypteridaceae and Woodsiaceae (sensu Christenhusz et al., 2011), which share the character of a laterally attached indusium (except in some Thelypteridaceae) and two vascular bundles in their petioles. Even though synapomorphies are few and not visible to the naked eye, it is possible to treat these as two broad families instead of the current 18 (or 20) based on their shared characters. Within these two lineages, a plethora of families has recently been proposed (including the recent segregation of Arthropteridaceae from Tectariaceae; Liu et al., 2013), and many genera have changed family (e.g. Christenhusz et al., 2013) when new phylogenetic analyses have become available. Such a broad familial circumscription, if accepted, would make the taxonomy of these groups more stable, allowing further changes at subfamilial levels without affecting the rank of family, which is the one above the genus most frequently used by non-specialists.

A summary of published studies showing relationships of major groups of ferns as well as some problematic genera discussed above is given in Fig. 3. This tree is further discussed below.

CONCLUSIONS
As often observed in taxonomy of many organisms, there has been a trend from an initial artificial classification towards more natural ones over time, although the definition of the latter has changed several times. Artificial classifications are useful for identification purposes, which were often the only intention of early authors (such as Linneaus with his sexual system of plant classification), and these were not intended to reflect phylogenetic relationships between taxa; of course, phylogenetic classification was not a conceptual framework available to taxonomists at that time. These artificial and early ‘natural’ groupings often resulted in species that are superficially similar being placed together, which does not take convergent evolution into account; some such cases in ferns required the application of DNA phylogenetics to demonstrate that they were not reflective of relationships. On the other hand, natural classifications may separate species that are highly similar, particularly in terms of ‘key characters’ that were judged (often subjectively) to be important, in completely unrelated families, which makes these families difficult to tell apart despite their distant relationship. Fern classification is particularly full of such problems, and now that we can use DNA studies to obtain information about where problematic species and genera should be placed, we often find that we lose the capacity to diagnose many families with clear sets of morphological characteristics. As has been observed previously, the more natural a classification becomes, the more difficult it typically is to use, especially for researchers not intimately familiar with the details of fern taxonomy. This is of course not unique to ferns, and it can also be observed in recent classifications of some angiosperm families such as Asparagaceae, Liliaceae and Xanthorrhoeaceae in monocots and Acanthaceae, Gesneriaceae, Lamiaeae and Plantaginaceae among eudicots in APG III (2009). There are few characters that can be easily used...
to distinguish these families, and their history has been as complicated and controversial as those for most fern families.

A second common trend has been the use of minute characters to circumscribe narrowly circumscribed genera, and some authors have used phylogenetic trees as the basis for placing every pair of sister species into separate genera. Following such episodes, another trend typically emerges, which expands generic limits and subsumes such small genera into larger ones, often redefining the original generic concept to fit this greater number of species with more divergent morphology. Even though this waxing and waning of generic limits may eventually produce a consensus taxonomy, it destabilizes nomenclature and confuses the users. Ultimately, such instability hinders research on ferns by non-taxonomists, thereby retarding progress on understanding their biology, conservation, ecology and population dynamics. Of course, in an ideal world, complete revisions at a generic level should be a prerequisite before familial revisions, but monographic study is time consuming, material for study is not always available, species sampling is often incomplete and funding for such research is scant, resulting in many groups not having been studied in their entirety. Nevertheless, when type species of genera are sequenced and analysed, phylogenetic positions of genera can be inferred and, even though a genus may be found to be polyphyletic, the type species and its associated original generic concept will remain, merely resulting in the transfer of species to other genera, but not genera to different families. Ferns have been notorious in the past for having an unstable higher level classification, but molecular phylogenetic techniques have helped to better understand relationships between genera, and this has resulted in the three most recent classifications (Smith et al., 2006, 2008; Christenhusz et al., 2011) being highly similar, especially when compared with most earlier classifications. There are still likely to be changes in the treatment of families recognized in future classifications. Numerous small families in the eupolypods could be merged with a larger family to which they are sister, such as Davalliaceae with Polypodiaceae, Nephrolepiaceae with Lomariopsidaceae, Hypodematidae with Dryopteridaceae, and Hemidictyaceae with Aspleniaceae. We would actually argue that to stabilize fern familial classification, the two clades informally called eupolypods I and II should be treated at the family level, Polypodiaceae and Aspleniaceae, respectively, in which the corresponding families of Christenhusz et al. (2011) would be treated at the subfamilial level. This may now seem like it would be further destabilizing the fern classification, but it will in the longer term stabilize matters, especially when taking into account the poor knowledge of several currently recognized large families, such as Woodsiaceae and Tectariaceae. Within a broader family concept, the genera may be moved between subfamilies but will remain in the same larger family, which is the only higher taxonomic level frequently used by non-taxonomists.

Current consensus classification of extant Lycopsidophyta (lycopsids) and Polypodiophyta (ferns)

Numbers in parentheses after families correspond to estimated numbers of genera and species (genera/species). Currently accepted genera are listed; for a full synonymy of genera see Christenhusz et al. (2011). Estimated species numbers for some of the larger genera are also provided. Sori of all major lineages are illustrated in Fig. 4.

**Lycopodiales DC.**
- **Lycopodiaceae P.Beauv.** (3/approx. 400) (Huperzia Bernh., Lycopodiella Holub, Lycopodium L.)
- **Selaginellaceae Willk.** (1/approx. 750) (Selaginella P.Beauv.)
- **Isoetales Prantl**
- **Isoetaceae Richenb.** (1/approx. 140) (Isoëtes L.)

**Ferns;** 21 families, approx. 212 genera, approx. 10 535 species

**Equisetales DC.**
- **Equisetaceae Michx.** (1/20) (Equisetum L.)

**Ophioglossidae K ling e**
- **Ophioglossales Link**
  - **Ophioglossaceae Martinov** (4/approx. 80) (Botrychium Sw., Helminthostachys Kauf., Mankyuu B.Y.Sun, M.H.Kim & C.H.Kim, Ophioglossum L.)
- **Psilotales Prantl**
  - **Psilotaceae J.W.Griff. & Henfr.** (2/12) (Psilotum Sw., Tmesipteris Bernh.)

**Marattidae K ling e**
- **Marattaceae Kauf.** (6/approx. 130) subfamily **Danaeoidae J.Williams** (Danaea Sm.), subfamily **Marattioideae C.Presl** (Angiopteris Hoffm., Christensenia Maxon, Eu podium J.Sm., Maratia Sw., Ptisana Murdock)

**Equisetaceae**

1The number of genera in Lycopodiaceae is not yet certain, but molecular phylogenetic studies (Wikström and Kenrick, 1997) have clearly identified three clades, corresponding to the genera Huperzia, Lycopodiella and Lycopodium. A treatment of three genera will result in the loss of some well-established names such as Diphasiostrum Holub as a synonym of Lycopodium and Phylloglossum Kunze in Huperzia. Use of these names has resulted in the recognition of a great number of genera in Lycopodiaceae, and as a result some authors have treated the three broader clades as subfamilies (e.g. Ölligaard, 2012). We see no benefit in destabilizing the taxonomy of these genera to accommodate these additional groups at the generic level, and these should therefore be treated at the subgeneric level if some form of formal classification of these groups is desired.

2Delimitation of genera in Ophioglossaceae is generally accepted, although many segregate genera have been proposed, subdividing Botrychium and Ophioglossum. The latter includes epiphytic Cheiroglossa C.Presl and Ophioderma (Blume) Endl., which can be maintained as separate, but, since they form a clade with core Ophioglossum, we have included them there.

3Marattiacae can be divided into two subfamilies, although one will be monogenic, which calls into question the need for subfamilies, especially when there are only six genera recognized in total. Danaea is nevertheless sufficiently genetically and morphologically distinct from other extant genera in the family that it could be recognized at the subfamilial level. Archangiopteris Christ & Giesenh., is embedded in Angiopteris, and, to prevent Angiopteris from being synonymized with Marattia, the last has been divided into three genera (Murdock, 2008).
subfamily Loxsomatoideae Christenh.\(^7\)
(Loxsoma R.Br., Loxsomopsis Christ)
subfamily Culcitoideae Christenh.\(^8\)
(Culcia C.Presl)
subfamily Plagiogyrioidae Christenh.\(^9\)
[Plagiogryia (Kunze) Mett.]
subfamily Cibiotoideae Nayer
(Cibiotium Kaulf.)
subfamily Cyatheoideae Endl.\(^10\)
(Alsophila R.Br., Cyathea Sm., Gymnosphaera Blume, Sphaeropteris Bernh.)
subfamily Dicksonioideae Link
[Calochlaena (Maxon) R.A. White & M.D. Turner, Dicksonia L’Hér., Lophosoria C.Presl]
subfamily Metaxyoidoideae B.K. Nayar
(Metaxy C.Presl)

**Polyplodies Link**

**Cystidiaceae J.R. Croft** (1/1)
(Cystodium J.Sm.)

**Lonchitidaceae C.Presl** (1/2)
(Lonchitis L.)

**Lindsaeaceae C.Presl** (6/approx. 220)

**Saccolomataceae Doweld** (2/approx. 12)
(Orthiopteris Copel., Saccoloma Kaulf.)

**Dennstaedtiaceae Lotsy** (10/approx. 240)
[Blotiella Tryon, Dennstaedtiia Bernh., Histiopteris (J. Agardh) J.Sm., Hypolepis Bernh., Leptolepia Prantl, Microlepia C.Presl, Monachosorum Kunze, Onotrichia Copel., Paesi St.-Hil., Pteridium Gled.]

12 Several issues remain in the variable subfamily Pteridoideae. Anogramma is polyphyletic with regard to Pityrogramma, Cerosora and Cosentinia (Schneider et al., 2013). Perhaps these are better treated under a single genus or maybe as two genera, merging Cerosora and Anogramma chaerophylla (Desv.) Link with Pityrogramma and Cosentinia with core Anogramma. The status of Aspleniopsis, Austrogramma and Syngramma has not yet been studied, and some of these may not remain after phylogenetic revision of this group. Pteris may be polyphyletic, and it is possible that up to three genera should be accepted or some genera may have to be merged with it to achieve monophyly. Pteris in the broad sense includes Afropoteris Alston, Anopteris Prantl, Camptopteris C. Presl, Litothrix C. Presl, Neurocallis Fée and Platycyana R. Br. (Schnüppel and Pryer, 2007; Christenhusz et al., 2011).

13 The genera in this subfamily are in great need of recircumscription. Most previously recognized genera, such as Chelanioidae, Doryopteris and Pellaea, are not monophyletic, although about 14 clades can be recognized (Eiserhardt et al., 2011). Perhaps these are better treated under a single genus or maybe as two genera, merging Chelanioidae and Anogramma chaerophylla (Desv.) Link with Pityrogramma and Cosentinia with core Anogramma. The status of Aspleniopsis, Austrogramma and Syngramma has not yet been studied, and some of these may not remain after phylogenetic revision of this group. Pteris may be polyphyletic, and it is possible that up to three genera should be accepted or some genera may have to be merged with it to achieve monophyly. Pteris in the broad sense includes Afropoteris Alston, Anopteris Prantl, Camptopteris C. Presl, Litothrix C. Presl, Neurocallis Fée and Platycyana R. Br. (Schnüppel and Pryer, 2007; Christenhusz et al., 2011).

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15 Subfamily Cystopteridoideae Link, Cystopteris Nakai, Deparia Hook. & Grev., Diplazium Sw. (approx. 350) subfamily Blechnoideae Hook. [Blechnum L. (approx. 250), Onoclea L.18, Stenochlaena J. Sm., Woodwardia Sm.]
Polyodiaceae J.Presl & C.Presl = eulopods I
(subfamily Didymochlaenoideae Christenh. 21
(Didymochlaena Desv.)
(subfamily Hypodemioidae Christenh. 22
(subfamily Dryopteridoideae Link 23
(subfamily Lomariopsidoideae Crabbe, Jermy & Mickel 24
rest of Blechnum sensu lato contains the vining taxa Stenochlaena, Salpigichaena J.Sm. and a few non-vining Blechnum species with long-ceping rhizomes, which may have to be accepted at the generic level pending further studies. 

D. canariensis (L.) Sm., has only recently

Lomariopsis is placed here following recent findings of Christenhusz et al. (2013). Nephrolepis probably also belongs here (as stated by Smith et al., 2006). The unsampled Thysanocostia Gepp from New Guinea is here tentatively merged with Lomariopsis on morphological grounds: Lomariopsis dimorphophylla (Gepp) Christenh., comb. nov. Basionym: Thysanocostia dimorphophylla Gepp, Contr. Phytogeog. Fl. Arfak Mts. 193. 1917. Lomariopsis pteridiformis (Ces.) Christenh., comb. nov. Basionym: Gymnogramma pteridiformis Ces., Rendiconti della Reale Accademia delle Scienze Fisiche e Matematiche di Napoli 16: 30. 1877. A number of genera previously placed in the Tectaria group by Holtum (1991) appear to belong to either the Cenitis assembly (Dryopteridoideae) or Lomariopsidoideae. Further studies should show the relationships within this subfamily, Psammiosorus C.Chr. belongs to Arthropteris (e.g. Lehtonen, 2011; Lü et al., 2013).

Some controversy remains about the generic classification of Davallioideae, due to the work of Kato and Tsutsui (2009), Tsutsui and Kato (2006) and Tsutsui et al. (2008). Even though clades can be recognized as genera in the Davallia alliance, the type of the genus, D. canariensis (L.) Sm., has only recently been included in a study (Lü & Schneider, 2013), supporting the concept of two genera. Davallia is better treated in its broad sense including Araiaestegiella M.Kato & Tsutsui and Humata Cav. because characters proposed to define these genera do not hold up to closer scrutiny, which does not warrant splitting them from Davallia. Araiaestegiella Copel. is tentatively included in Davallidae, but it may warrant recognition.

Currently being studied, tribe Drynariæae probably only consist of two genera (Drynaria and Selliguea), but how these are to be defined is still being investigated (H. Schneider, pers. comm.).


Tribe Microsoreae need major study to determine generic delimitation, especially Microsorium. Colysis C.Presl and Kontumia S.K.Wu & P.K. Löc belong to Leptochilus (Kim et al., 2013). Leptochilus heterophyllus (S.K.Wu & P.K. Löc) Christenh., comb. nov. Basionym: Kontumia heterophyllus S.K.Wu & P.K. Löc, Novon 15: 245. 2005. Many small genera are still commonly applied, and larger genera are poorly defined, but with phylogenetic analyses it should be possible to apply a broader generic concept and redefine these as larger more natural entities.
tribe Polypodiaceae Hook. & Lindl. ex Duby

[Campyloneurum C.Presl, Grammitis Sw. (approx. 700), Microgramma C.Presl, Pecluma M.G.Price, Phlebodium (R.Br.) J.Sm., Pleopeltis Humb. & Bonpl., Pleurosoiripsis Fomin, Polypodium L. (approx. 70), Serpocalton A.R.Sm., Syngnamma C.Presl]

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LITERATURE CITED


30As stated earlier in this paper, the grammitid ferns are all treated as a single genus in this classification. Few of the numerous relatively new genera have been properly evaluated for monophyly, and some of those that have been studied were found to be polyphyletic (Ranker et al., 2004; Sundue et al., 2010; Hirai et al., 2011), resulting in the description of yet more genera. The broad approach taken for Asplenium and Hypnomenophyllum seems to be a better fit here. Of course, when this is done, the remaining genera in tribe Polypodiaceae also need to be addressed. Niphidium J.Sm. and Campyloceras are difficult to separate and should therefore be merged, better to be treated at the subgeneric level, with the latter name having nomenclatural priority. Polypodium and its close relatives should also be reassessed, and generic delimitation may have to be addressed in the future.


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