

Multiple origins of crassulacean acid metabolism and the epiphytic habit in the Neotropical family Bromeliaceae

Darren M. Crayn^{†‡}, Klaus Winter[†], and J. Andrew C. Smith^{§¶}

[†]Smithsonian Tropical Research Institute, Box 2072, Balboa, Ancon, Republic of Panama; [‡]Royal Botanic Gardens, Mrs. Macquaries Road, Sydney, NSW 2000, Australia; and [§]Department of Plant Sciences, University of Oxford, South Parks Road, Oxford OX1 3RB, United Kingdom

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The large Neotropical family Bromeliaceae presents an outstanding example of adaptive radiation in plants, containing a wide range of terrestrial and epiphytic life-forms occupying many distinct habitats. Diversification in bromeliads has been linked to several key innovations, including water- and nutrient-impounding phytotelmata, absorptive epidermal trichomes, and the water-conserving mode of photosynthesis known as crassulacean acid metabolism (CAM). To clarify the origins of CAM and the epiphytic habit, we conducted a phylogenetic analysis of nucleotide sequences for 51 bromeliad taxa by using the plastid loci *matK* and the *rps16* intron, combined with a survey of photosynthetic pathway determined by carbon-isotope ratios for 1,873 species representing 65% of the family. Optimization of character-states onto the strict consensus tree indicated that the last common ancestor of Bromeliaceae was a terrestrial C₃ mesophyte, probably adapted to moist, exposed, nutrient-poor habitats. Both CAM photosynthesis and the epiphytic habit evolved a minimum of three times in the family, most likely in response to geological and climatic changes in the late Tertiary. The great majority of epiphytic forms are now found in two lineages: in subfamily Tillandsioideae, in which C₃ photosynthesis was the ancestral state and CAM developed later in the most extreme epiphytes, and in subfamily Bromelioideae, in which CAM photosynthesis predated the appearance of epiphytism. Subsequent radiation of the bromelioid line into less xeric habitats has led to reversion to C₃ photosynthesis in some taxa, showing that both gain and loss of CAM have occurred in the complex evolutionary history of this family.

The Bromeliaceae are frequently celebrated as an outstanding example of adaptive radiation in vascular plants (1, 2). They represent one of the largest families with a Neotropical distribution (3), comprising 2,885 species in 56 genera (4, 5), with an ecological range that encompasses extremes of moisture availability (from rain forests to hyperarid coastal sands), elevation (from sea level to >4,000 m), and exposure (fully exposed sites to shaded forest understories). The family contains a correspondingly rich diversity of life-forms, from soil-rooted terrestrial plants, through rosulate “tank” epiphytes with water- and nutrient-impounding phytotelmata, to extreme epiphytes completely independent of their substratum for nutrition. The evolutionary transition from terrestrial to epiphytic life-forms appears to have been closely linked to elaboration of the absorptive epidermal trichomes characteristic of the family (1, 6, 7). Indeed, about half of all bromeliads are epiphytic, and they constitute one of the most distinctive components of the Neotropical forest canopy (8, 9).

In addition to morphological specializations, another key innovation associated with the success of bromeliads in more arid habitats is the form of photosynthesis known as crassulacean acid metabolism (CAM) (10–13). In typical CAM plants, CO₂ is taken up at night and temporarily stored, via a pathway involving fixation by phosphoenolpyruvate carboxylase (PEPC), in the form of malic acid in the cell vacuole. In the following light period the stomata close, malic acid is released from the vacuole

and decarboxylated, and the CO₂ liberated is photosynthetically reduced in the Calvin cycle (13). By restricting gas exchange with the atmosphere during the daytime, CAM plants use their available water more efficiently than C₃ plants, and consequently characterize many tropical and subtropical environments with intermittent or strongly seasonal water supply. This includes epiphytic niches in the forest canopy, which can be microclimatically arid, to the extent that, amongst the 6% of flowering plants estimated to show CAM photosynthesis, there may be almost as many epiphytic as terrestrial species (13, 14).

Despite being one of the best-understood metabolic examples of an ecological adaptation in plants, relatively little is known of the evolutionary origins of the CAM pathway. Its occurrence in >30 diverse families suggests that CAM has arisen many times, but only limited work has been undertaken from a phylogenetic perspective at a finer taxonomic scale. Because of their diversity, the Bromeliaceae provide an excellent model of adaptive radiation on which to trace the origins of CAM (and the epiphytic habit) in closely related taxa. A prerequisite for such evolutionary reconstruction is a sufficiently robust phylogeny for the family based on molecular and morphological characters, given the almost complete lack of a fossil record (15). Taxonomically, Bromeliaceae have long been regarded as an isolated and natural group (3, 16), a view supported by cladistic analyses of molecular data that resolve a monophyletic Bromeliaceae within the large order Poales (17–19). The three traditionally recognized subfamilies (20, 21) all contain a mixture of C₃ species and CAM species (12, 22), but the first molecular-phylogenetic analyses shed little light on CAM evolution because of limited taxon sampling and poor resolution on the trees (23–26). Moreover, CAM is common in both terrestrial and epiphytic bromeliads, so the underlying relationships between photosynthetic pathway, plant life-form, and phylogenetic lineage in the family may be complex.

To obtain further insight into the origins of CAM photosynthesis and its relationship to the evolution of epiphytism, we have derived a more detailed phylogeny for the Bromeliaceae based on nucleotide sequences of two rapidly evolving plastid loci, *matK* and the *rps16* intron. This is combined with an extensive survey of photosynthetic pathway at the species level to determine the minimum number of times CAM may have arisen in this family. Together with evidence from present-day biogeography and ecology, this permits a reconciliation of previously conflicting hypotheses for the origins of CAM photosynthesis and the epiphytic life-form within this exceptionally diverse Neotropical family.

Abbreviations: CAM, crassulacean acid metabolism; RTA, random-taxon-addition-order; SW, successive weights.

Data deposition: The sequences reported in this paper have been deposited in the GenBank database (accession nos. can be found in Table 2, which is published as supporting information on the PNAS web site).

[¶]To whom correspondence should be addressed. E-mail: andrew.smith@plants.ox.ac.uk.

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Materials and Methods

Molecular Systematics. Fifty-one species of Bromeliaceae from 27 genera were chosen to represent known diversity within and between previously recognized taxonomic groups. Species selection was biased toward subfamily Pitcairnioideae (20, 21), because this group may contain the earliest diverging lineages in the family (1, 24, 26, 27) and includes both C₃ and CAM taxa (10, 12, 22). Species of seven genera of Tillandsioideae (of nine currently recognized) and eight (of 30) genera of Bromelioideae were chosen to reflect the ecological and taxonomic diversity in those groups. Three species of Rapateaceae were used as outgroups, because most molecular studies place Rapateaceae amongst the closest relatives of Bromeliaceae (18, 28).

Sequences of the *matK* gene (29) and *rps16* intron (30) were obtained by using standard protocols for total DNA extraction, PCR amplification, and sequencing (25). These are among the most rapidly evolving plastid loci and are well suited to phylogenetic reconstruction within many angiosperm families (29–32). Details of PCR primers used are given in Table 1, and DNA sequence information, GenBank accession numbers, and voucher details are provided in Table 2, both of which are published as supporting information on the PNAS web site.

Manual alignment of sequences and all subsequent analysis was performed in PAUP* 4.0b6 (33). Inferred indels, where informative, were scored as additional characters following the “simple” method of Simmons and Ochoterena (34).

The *matK* and *rps16* intron data were first analyzed separately and then simultaneously. Phylogenetic resolution can be improved by combining independent molecular data sets (32), and analysis of *matK* sequences for a subset of these species suggested that this locus alone would not provide sufficient resolution on the resulting trees (25, 35). Parsimony analysis used tree bisection–reconnection branch swapping and successive weights (SW) analysis (36), with iterative rounds of search followed by reweighting until tree length stabilized. To search for multiple islands of optimal trees, a random-taxon-addition-order (RTA) procedure (37) was used with 1,000 replicates, saving 30 trees per replicate. Weights were determined by using the maximum rescaled consistency index for each character on the best trees. Trees found in the final round of SW analysis were swapped to completion (using 1,000 RTA replicates) or until 80,000 trees were found. Before proceeding with the simultaneous analysis, we tested the combinability of the *matK* and *rps16* intron data by using the incongruence length difference (ILD) test (38) to assess the statistical significance of congruence between phylogenies based on the two data partitions. The test was performed by 200 replicate parsimony analyses, each consisting of 100 RTA searches, with 100 trees saved per search. Final SW values were used.

Branch support was evaluated by 1,000 bootstrap replicates (using the SW values), saving 100 trees per replicate. This “reduced effort” procedure gives an unbiased estimate of the true bootstrap proportions (39). Support for alternative evolutionary hypotheses was evaluated by using the Kishino–Hasegawa test (40). One of the best trees consistent with each hypothesis (found by a constraint parsimony analysis with 1,000 replicate RTA searches, SW values, 30 trees saved per replicate) was compared with one of the trees found by SW analysis of the combined data set.

Photosynthetic Pathway. Photosynthetic pathway was determined from tissue carbon-isotope ratio, $\delta^{13}\text{C}$. This can distinguish plants that use the C₃ pathway, in which the primary carboxylating enzyme is ribulose-1,5-bisphosphate carboxylase-oxygenase, from C₄ or CAM plants, in which the primary carboxylating enzyme is PEPC, because of a kinetic isotope effect (41). C₄ photosynthesis is not known in Bromeliaceae (1,

11), so the $\delta^{13}\text{C}$ values could be used to distinguish CAM and C₃ species. Relative natural abundance of ¹²C and ¹³C ($\delta^{13}\text{C}$) was determined for samples of dried shoot tissue taken from herbarium specimens as described (41, 42).

Species were classified as CAM or C₃ if the $\delta^{13}\text{C}$ value was less negative or more negative than -20.0‰ , respectively. [Values more negative than -20.0‰ do not preclude the possibility of some dark CO₂ fixation, but indicate that this did not make a major contribution to photosynthetic carbon gain (43).] The character states “C₃” and “CAM,” and “terrestrial” and “epiphytic” were mapped onto the strict consensus tree by using MACCLADE 3.08a (44).

Results

Molecular Systematics. For the 51 species of Bromeliaceae and 3 Rapateaceae, the aligned *matK* data set comprised 858 positions, 142 of which were potentially parsimony-informative, with one indel (shared by the three *Puya* spp.). Ingroup pairwise sequence divergence (uncorrected for multiple hits) reached a maximum of 5.3% (between *Fosterella penduliflora* and *Guzmania monostachia*). Alignment of the *rps16* intron sequences required 63 gaps, which were scored as 22 indels (seven being restricted to the outgroup). This data set comprised 1135 positions, 177 of which were parsimony-informative; ingroup pairwise sequence divergence reached a maximum of 4.6% (between *Ayensua uaipanensis* and *Pitcairnia nuda*).

Estimates of phylogenetic relationships derived from the *matK* and *rps16* intron data sets did not differ significantly ($P = 0.175$) based on the ILD test, so we proceeded with a simultaneous analysis. Initial (unit weight) parsimony analysis of the combined data set of 1,993 characters, including the indel characters, found optimal trees of length 753 steps (consistency index excluding uninformative characters, CI = 0.749, retention index, RI = 0.819). SW analysis converged on trees of length 495 steps (CI = 0.928, RI = 0.947) after three rounds. Swapping to completion found 756 trees in one island. The strict consensus of these trees (which is identical to that from an equal weights analysis) is shown in Fig. 1.

This analysis strongly supports the monophyly of Bromeliaceae (bootstrap value of 100%). Other well supported groups (bootstrap values >80%) include the two subfamilies Bromelioideae and Tillandsioideae, and the genera *Brocchinia*, *Catopsis*, *Dyckia*, *Encholirium*, *Fosterella*, *Hechtia*, and *Puya*. *Deuterocohnia meziana* does not group with other *Deuterocohnia* species but is part of a robust clade including *Dyckia* and *Encholirium*, confirming that this genus is not monophyletic (24–26). *Pitcairnia* species are placed in two well supported groups, *Pitcairnia* 1 (3 spp.) and *Pitcairnia* 2 (8 spp.). *Navia igneosicola* is sister to *Pitcairnia* 2, whereas *N. phelpsi* groups with *Cottendorfia*. In addition, there is support for *Brocchinia* as sister to the rest of the family, *Puya* as sister to Bromelioideae, and a derived clade comprising Bromelioideae and all of the Pitcairnioideae sampled except for *Brocchinia*, *Hechtia*, *Cottendorfia*, and *Navia phelpsi* (the “DFPPB” clade). Within Tillandsioideae, *Catopsis* and then *Mezobromelia* plus *Werauhia* are strongly supported as successive sister groups to a core clade comprising members of *Guzmania*, *Racinaea*, *Tillandsia*, and *Vriesea*.

Photosynthetic Pathway. Carbon-isotope ratios for the 51 species used to construct the molecular phylogeny are optimized by using parsimony onto the strict consensus tree in Fig. 2 together with plant life-form. A Kishino–Hasegawa test rejects the hypothesis of a single origin for CAM ($P < 0.001$). A sister-group relationship between *Hechtia* and Tillandsioideae receives only weak bootstrap support, and indeed the present data do not rule out a placement of *Hechtia* as sister to the DFPPB clade ($P = 0.05$), but *Hechtia* as sister to Tillandsioideae is corroborated by maximum likelihood and neighbor-joining analyses of *trnL* in-

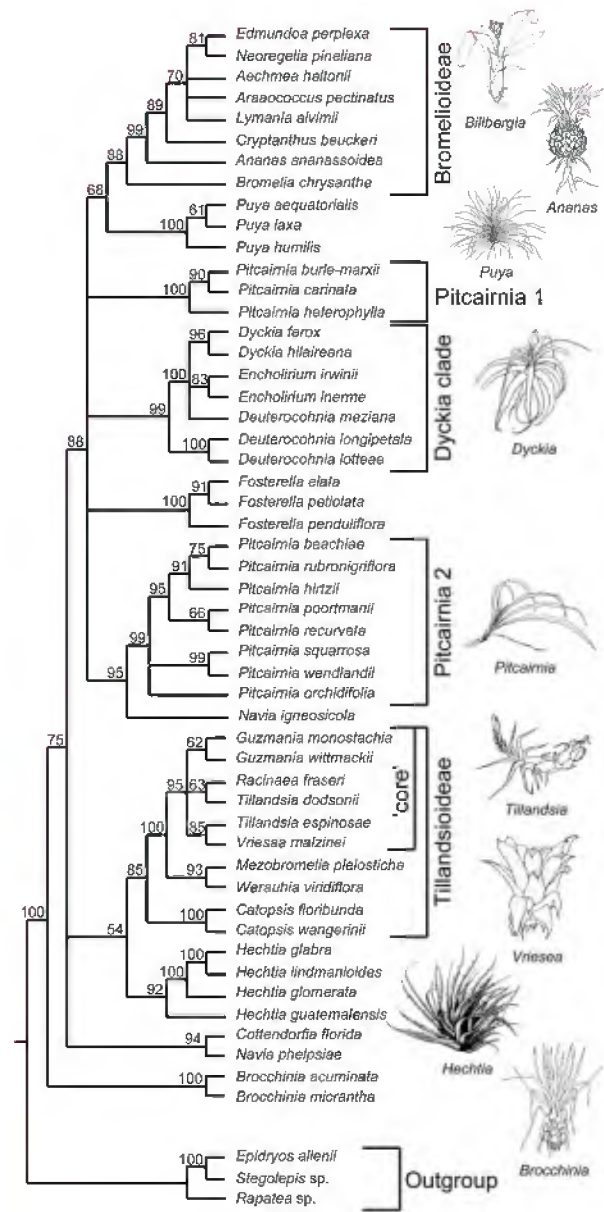


Fig. 1. Strict consensus of 756 trees found during the final round of 5W analysis of the combined *matK* plus *rps16* intron data set for 51 species of Bromeliaceae. The tree was rooted on the branch separating Rapateaceae and Bromeliaceae. Bootstrap values are indicated above the relevant branches; clades referred to in the text are bracketed alongside representative life-forms. [Illustrations reproduced with permission from ref. 1 (Copyright 2000, Cambridge Univ. Press) and ref. 45 (Copyright 1997, Missouri Botanical Garden).]

tron sequence (26). A sister-group relationship between *Fosterella* and the *Dyckia* clade also receives weak bootstrap support but is not present in the strict consensus tree. Thus, CAM photosynthesis occurs in four clades (Fig. 2): *Hechtia*, the “core” Tillandsioideae, the *Dyckia* clade, and *Puya* + Bromelioideae. In contrast, epiphytes are distributed in two main clades, Tillandsioideae (virtually all of which are epiphytic or lithophytic) and Bromelioideae [≈50% of which are epiphytic (1, 20, 45)]. Two of ≈20 species of *Brocchinia* can grow epiphytically, although this is only one of several life-forms found in this ecologically diverse genus (1, 2, 46).

Because taxon sampling for sequencing of the plastid loci was

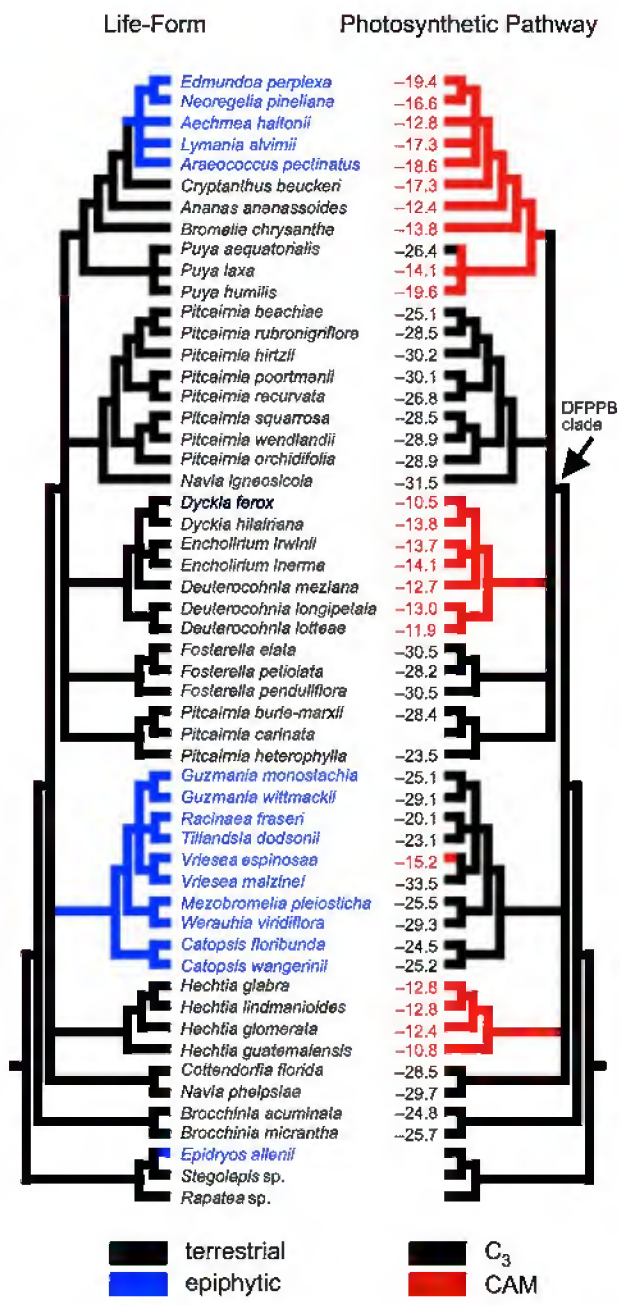


Fig. 2. Most-parsimonious reconstruction of the evolution of life-form and photosynthetic pathway in Bromeliaceae, based on relationships supported by bootstrap analysis of the combined *matK* plus *rps16* intron data set. Carbon-isotope ratios ($\delta^{13}\text{C}$ values in ‰) are shown for the taxa analyzed. The derived character-states “epiphytic” and “CAM” are highlighted in blue and red, respectively. Two species of *Brocchinia* can grow epiphytically (not shown), probably representing a further independent origin of epiphytism in Bromeliaceae.

necessarily restricted, a more complete survey of $\delta^{13}\text{C}$ values in the family was undertaken that included 55 genera (i.e., all except the monotypic bromelioid genus *Pseudaechmea* L.B.Sm. & Read) and 1,873 species (65% of the estimated total in the family). Of these species, 826 (44%) were found to be CAM plants, all of which were in genera that can be ascribed to the four CAM lineages in Fig. 2. Taking into account the detailed ecological information available for the family and the range of species sampled, it is very unlikely that other lineages containing CAM plants exist in the Bromeliaceae.

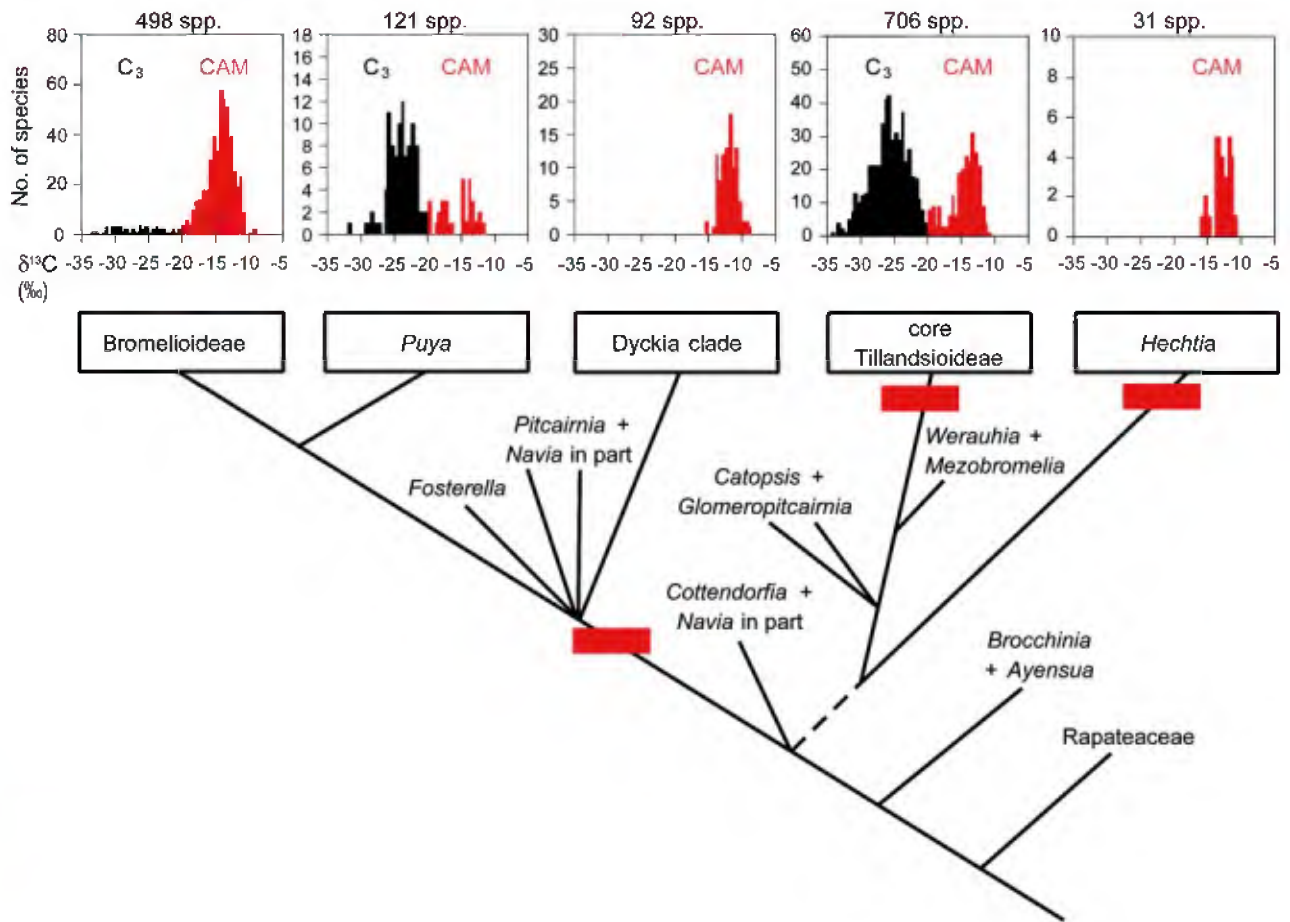


Fig. 3. Summary of numbers of C_3 and CAM species in the five lineages within Bromeliaceae found to contain CAM taxa. Phylogenetic inferences are based on Fig. 1, together with information on *Ayensua* (26) and *Glomeropitcairnia* (26, 53). Horizontal red bars indicate the minimum number of independent origins of CAM. The closest relative of *Hechtia* was not resolved by the present analysis; the dashed line indicates its affinities based on analysis of *trnL* intron sequence data (26).

The relative abundance of CAM species in the major phylogenetic lineages deduced from this study are summarized in Fig. 3. Both *Hechtia* and the Dyckia clade [*Deuterocohnia* (including *Abromeitiella*), *Dyckia*, and *Encholirium*] are comprised solely of CAM species. Amongst the Tillandsioideae, 28% (223 of 788 species sampled) were identified as CAM plants: these are effectively all restricted to the genus *Tillandsia*, because the four species of *Vriesea* (of 135 sampled) identified as CAM plants are taxa that should probably be realigned into *Tillandsia* (47). CAM is also well represented in *Puya* and Bromelioideae, which were resolved as sister groups (Fig. 1), as found in the *ndhF* (24) and *trnL* (26) phylogenies. In *Puya*, a minority of species sampled (24%) are CAM plants, whereas in Bromelioideae the great majority are CAM (91%).

The position of *Brocchinia* (sister to the rest of the family) supports previous suggestions that C_3 photosynthesis is plesiomorphic in Bromeliaceae (10, 12). The monotypic *Ayensua uaipanensis* formed a robust clade with *Brocchinia* (94% bootstrap support) based on *rps16* intron sequence data (data not shown), a relationship supported by *trnL* intron sequence analysis (26); however, *matK* sequence could not be obtained. With *Brocchinia + Ayensua* at the base, the phylogeny indicates a minimum of three independent origins of CAM within the family (Fig. 3): one ancestral to *Hechtia*, one in the core Tillandsioideae, and a third in the DFPPB clade. On the basis of the *matK + rps16* intron data set alone, it is not possible to reject the hypothesis that CAM had only a single origin in the DFPPB

clade, but an *ndhF* phylogeny also resolved *Dyckia* in a pitcairnioid clade distinct from *Puya + Bromelioideae* (24) (Fig. 2), suggesting that CAM evolved independently in these two lineages.

Discussion

Plants showing CAM photosynthesis are widely believed to have evolved from C_3 ancestors, but the exact circumstances under which the major CAM lineages arose are not well understood. The highly dispersed taxonomic distribution of CAM photosynthesis, which occurs in 33 families and an estimated 16,000 species of vascular plants (13), suggests it has arisen on multiple occasions. In families such as the Agavaceae, Cactaceae, and Didiereaceae, almost all species have the capacity for CAM and thus exhibit the presumed apomorphic character-state (13, 48). Other families such as the Aizoaceae, Bromeliaceae, Crassulaceae, and Orchidaceae contain large numbers of both C_3 and CAM species, so these may be more informative for reconstructing the origins of the CAM pathway, providing such an analysis can be supported by an appropriately resolved phylogeny.

Although the Bromeliaceae have been much studied with respect to their ecological diversity and life-forms, taxonomic relationships within the family have remained controversial (1, 3, 20, 24, 45). Previous molecular-phylogenetic studies have suffered from relatively poor resolution because of low sequence divergence within the family, but the *matK* and *rps16* loci used in the present study gave a well-resolved phylogeny when ana-

lyzed as a combined data set. Our results support a basal separation of *Brocchinia* + *Ayensua* (24, 26), distinctive C₃ taxa that are geographically restricted to the escarpment of the Guayana Shield (1, 2, 46, 49). Further, the family Rapateaceae, which appears the most likely sister group and was used to root the bromeliad tree, also has a distribution centered on the Guayana Shield and consists largely of terrestrial C₃ herbs of wet, infertile soils, frequently cooccurring with *Brocchinia* (2, 28, 42). Given the absence of CAM and rarity of epiphytism in the other 17 families making up the order Poales (9, 13, 19), this strongly suggests that both C₃ photosynthesis and a terrestrial growth habit are plesiomorphic in Bromeliaceae.

Within Bromeliaceae, the epiphytic life-form and CAM photosynthesis have clearly arisen multiple times independently, so their origins must be sought in the evolutionary history of separate lineages. In the family as a whole, there is a strong correlation between habitat aridity and the occurrence of CAM (10–12, 50), but CAM is widespread in both terrestrial and epiphytic species, and in all three major subfamilies. Our survey of carbon-isotope ratios suggests that most genera of the largely terrestrial Pitcairnioideae are exclusively either CAM (*Hechtia*, *Dyckia*, *Encholirium*, *Deuterocohnia*) or C₃ (*Brocchinia*, *Navia*, *Steyerbromelia*, *Brewcaria*, *Cottendorfia*, *Lindmania*, *Connellia*, *Pitcairnia*, *Fosterella*), and that only *Puya* contains both C₃ and CAM species. The *matK* plus *rps16* phylogeny confirms earlier suggestions (23–26) that Pitcairnioideae as traditionally circumscribed are paraphyletic, although formal taxonomic revision should await clarification of the phylogenetic relationships of four other rare genera of Guayana Shield endemics, *Steyerbromelia*, *Brewcaria*, *Lindmania*, and *Connellia*. Among the CAM taxa, *Hechtia* had been considered closely related to the other xeromorphic pitcairnioids with succulent, spiny leaves (21, 27), but the present analysis suggests that CAM and the associated vegetative characters are independently derived in *Hechtia* and the DFPPB clade (Fig. 2). *Hechtia* has a notably disjunct distribution, its 51 species being restricted to northern Central America, Mexico, and southern Texas (20). CAM very likely arose in this taxon in the same arid-zone habitats that fostered evolution of the Agavaceae and Cactaceae, two of the most distinctive Neotropical families of terrestrial CAM plants (48, 51). In contrast, the CAM taxa in the *Dyckia* clade (Figs. 2 and 3), comprising *Deuterocohnia*, *Dyckia*, and *Encholirium*, are all centered on xeric habitats in the southern Andes, Argentina, and south and eastern Brazil (20).

The *matK* plus *rps16* phylogeny confirms the monophyly of the two other bromeliad subfamilies, Tillandsioideae and Bromelioideae, consistent with phylogenies derived from *ndhF* (24) and *trnL* intron (26) sequences with somewhat different taxon sampling. Tillandsioideae are almost wholly epiphytic or lithophytic, but C₃ photosynthesis is clearly plesiomorphic in the subfamily (Figs. 2 and 3). CAM photosynthesis is restricted to *Tillandsia* s.l., a very large (≈540 spp.) and diverse genus containing both C₃ and CAM species (12, 20, 22, 52, 53). The epiphytic habit also reaches its most extreme form in this genus, approximately half of which (the so-called “atmospheric” species) lack water-impounding phytotelmata, have root systems reduced to hold-fasts, and are entirely dependent for water and nutrient uptake on absorptive trichomes that cover the shoot (1). All of the atmospheric species of *Tillandsia* are CAM plants, suggesting that CAM may have been a key innovation enabling the adaptive radiation of this genus into more xeric habitats.

The other monophyletic subfamily, Bromelioideae, was resolved as sister group to the genus *Puya*. This relationship was also found in the *ndhF* (24) and *trnL* (26) phylogenies, and is supported by putative synapomorphies such as leaf morphology and trichome structure (1). A sister-group relationship of *Puya* and Bromelioideae has important implications for the origins of CAM photosynthesis. *Puya* is a large genus (195 spp.) of

terrestrial, often xeromorphic plants commonly found on open slopes of the Andean cordillera, but only a minority (24%) appear to be CAM plants. Whether the last common ancestor of the *Puya* + Bromelioideae lineage already possessed CAM photosynthesis, or whether CAM arose more than once in this clade, should be testable by a phylogenetic analysis with greater sampling density in these taxa. Within Bromelioideae, epiphytism is clearly the derived condition, but >90% of the subfamily are CAM species (and the basally diverging genera *Bromelia* and *Ananas* entirely so). Nevertheless, there has been considerable ecological diversification within the subfamily. Several CAM species of *Aechmea* are found in relatively shaded, humid habitats (11, 54, 55); epiphytic genera such as *Nidularium*, *Ronnbergia*, and *Wittrockia* contain both C₃ and CAM species; and four small genera that are phylogenetically more derived, *Fascicularia*, *Greigia*, *Fernseea*, and *Ochagavia* (26), contain exclusively C₃ species. Thus, although CAM is ancestral in the subfamily, there is evidence for reversion from CAM to C₃ photosynthesis as certain lineages radiated into more mesic habitats.

Many earlier authors have speculated on the evolutionary origins of epiphytism in Bromeliaceae, and the *matK* plus *rps16* phylogeny, together with other molecular studies (24, 26), helps to reconcile some previously conflicting views. Schimper (6) originally proposed that tropical epiphytes evolved from terrestrial ancestors in the relatively moist, shaded forest understory, with some forms migrating up the forest profile and eventually colonizing the canopy. Although this model is applicable to other families of rainforest epiphytes, Tietze (56) and Pittendrigh (45) proposed a radically different explanation for the origin of epiphytic bromeliads, suggesting they may have entered the forest as relatively light-demanding forms derived from terrestrial ancestors adapted to open habitats, subsequently diversifying into a variety of microhabitats including the shaded understory. The present results support the notion that the ancestral bromeliad was a terrestrial C₃ plant of exposed but relatively moist environments (3, 10, 45), perhaps similar to those occupied by present-day *Brocchinia* (1, 2). These habitats may also have been nutrient-poor, possibly providing a strong selective pressure for the evolution of absorptive trichomes (1, 10, 57). Apart from occasional examples of epiphytism in *Brocchinia* (2), the epiphytic life-form has become widespread in two main lineages. The last common ancestor of Tillandsioideae, although a C₃ plant, had already acquired an epiphytic growth habit, and this subfamily has remained wholly epiphytic (or lithophytic), with the appearance of CAM being limited to more xeric forms in the genus *Tillandsia*. In contrast, the last common ancestor of Bromelioideae already possessed CAM, and epiphytism, as well as reversion to C₃ photosynthesis, has been a later development in certain taxa. The phylogenetic reconstruction does not reveal any evidence for shade-tolerant terrestrial forms amongst the immediate ancestors of these two subfamilies. This lends support to the Tietze–Pittendrigh model, suggesting that the progenitors of epiphytic bromeliads were plants adapted to relatively exposed habitats, and that the species now found in the forest understory are secondarily shade-adapted.

The low degree of nucleotide sequence divergence found for four loci are consistent with the Bromeliaceae being relatively young. But in the almost complete absence of a fossil record, with the exception of a single report of *Tillandsia*-type pollen from the Upper Eocene (15), it is not yet possible to assign a precise chronology to the family's evolutionary history. The Neotropical distribution of bromeliads suggests an origin some time after the break-up of West Gondwana and the reduction of biological exchange between Africa and South America ≈85 million years ago (Ma) (58). An emergence of Bromeliaceae by the early Tertiary is also suggested by the appearance of other Poales in the fossil record by 70–55 Ma (59). Some major events

in the family's evolution may have occurred much more recently. For example, the mainly Andean distribution of *Puya* (20), and the abundance of Tillandsioideae in northern Peru, Ecuador, and Colombia, suggests that diversification may have been associated with the emergence of new habitats during periods of Andean orogeny in the Miocene and Pliocene (60, 61), as proposed for epiphytic Lycopodiaceae (62). Progressive aridification and declining CO₂ concentrations during the Tertiary (60, 63, 64) would have gradually favored the emergence of CAM photosynthesis in Bromeliaceae, perhaps in a manner similar to the Miocene expansion of grasses showing C₄ photosynthesis (65, 66). Firmer conclusions about the chronology of events will only be possible once the molecular phylogeny can be calibrated against other evidence. However, this study suggests that com-

bined information from a variety of sources may also be valuable in tracing the origins of CAM photosynthesis in other groups of vascular plants.

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