PHYLOGENY AND BIOGEOGRAPHY OF THE SARRACENIACEAE

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The carnivorous plant family Sarraceniaceae in the order Ericales consists of three genera: *Darlingtonia*, *Heliamphora*, and *Sarracenia*. *Darlingtonia* is represented by one species that is found in northern California and western Oregon. The genus *Heliamphora* currently has 23 recognized species all of which are native to the Guiana Highlands primarily in Venezuela with some spillover across the borders into Brazil and Guyana. *Sarracenia* has 15 species and subspecies, all but one of which are located in the southeastern USA. The range of *Sarracenia purpurea* extends into the northern USA and Canada. Closely related families in the plant order Ericales include the Roridulaceae consisting of two sticky-leaved carnivorous plant species, Actinidiaceae, the Chinese gooseberry family, Cyrillaceae, which includes the common wetland plant *Cyrilla racemiflora*, and the family Clethraceae, which also has wetland plants including *Clethra alnifolia*.

The rather charismatic plants of the Sarraceniaceae have drawn attention since the mid 19th century from botanists trying to understand how they came into being, how the genera are related to each other, and how they came to have such disjunct distributions. Before the advent of DNA sequencing it was very difficult to determine their relationships. Macfarlane (1889, 1893) proposed a phylogeny of the Sarraceniaceae based on his judgment of the overlap in features of the adult pitchers and his assumption that *Nepenthes* is a member of the family (Fig. 1a). He based his phylogeny on the idea that the pitchers are produced from the fusion of two to five leaflets. Goebel (1891) strenuously objected to the leaflet idea but Macfarlane did not accept or understand Goebel's arguments (Macfarlane 1893). Macfarlane placed *Heliamphora* more ancestral to *Darlingtonia* and *Sarracenia* because of the pitchers of *Heliamphora* are intermediate between the *Darlingtonia* and *Sarracenia* pair and *Nepenthes*. Both *Nepenthes* and *Heliamphora* have double "dorsal flaps" with those of *Nepenthes* being farther apart. Of course we know today that *Nepenthes* is in a totally different plant order and the pitchers of all four genera develop as if from one simple folded leaf (*cf.* Lloyd 1942).

Goebel (1891), Lloyd (1942), and Franck (1976) offered additional views on pitcher development and morphology. But unlike Macfarlane they were not champions for Darwinism so did not feel compelled to discuss their observations in an evolutionary context. And unlike Macfarlane they had access to more plant samples and studied the juvenile plants in much more detail. Goebel actually started *Darlingtonia* and *Sarracenia* from seed, studied the seedlings in detail, and lamented the fact he could not get any *Heliamphora* seeds. Lloyd, who studied under Goebel, did extensive field studies and saw variations in pitcher structure that are not typically seen in cultivated plants. Lloyd also confirmed and expanded on Goebel's observations of developing leaves. Except for removing *Nepenthes* from the phylogeny none of this work helped improve our understanding of the relationships between the genera.

Franck (1976) did the most important morphological work with respect to evolutionary biology. He showed that the juvenile pitchers of *Darlingtonia* are not immature versions of adult pitchers. The juvenile pitchers have significantly different developmental patterns from the adult pitchers

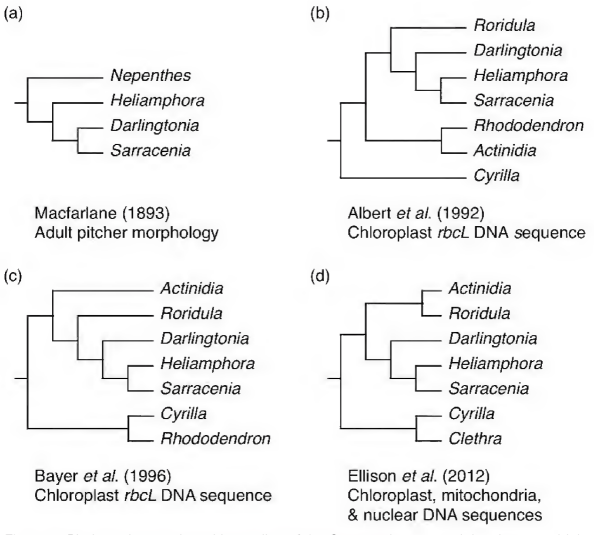


Figure 1: Phylogenies produced in studies of the Sarraceniaceae and the data on which each phylogeny was based. The length of the lines is arbitrary.

when compared to what is typically found in other plants that change leaf morphology as they grow. All the previous authors had assumed the juvenile pitchers were developmentally immature forms of the adult pitchers and thus not that interesting. I expect a study of *Heliamphora* adult and juvenile pitcher development would give similar results. However even today the juvenile pitchers of *Heliamphora* are pretty much ignored (*e.g.* McPherson *et al.* 2011) if not misunderstood. Juvenile *Heliamphora* pitchers may not provide much taxonomic value but like *Darlingtonia* juvenile pitchers are exquisitely adapted for trapping in a different way from the adult pitchers.

If the Sarraceniaceae is not related to *Nepenthes*, what is the family related to? The plant families closest to the Sarraceniaceae could not be determined from morphological details even as late as 1975. DeBuhr (1975) did a comparative study of the Sarraceniaceae in relation to other families in the Ericales to assign it to a suborder. Based on stamen morphology and number he placed the Sarraceniaceae in the suborder Theineae that contained the tea family, Theaceae, and the Actinidiaceae and not in the suborder Clethrineae with the families Clethraceae and Cyrillaceae. In retrospect the definition of Theineae was overly broad and almost a catchall suborder while the Clethrineae was too narrow based on incomplete information and the use of pollination-associated adaptations. The curious genus *Purdiaea* now placed in the Clethraceae based on DNA sequencing is from Central America, northern South America, and Cuba and matches the Sarraceniaceae in number of stamens (Anderberg & Zhang 2002) but did not fit the earlier definition of the Clethrineae. Species in the Clethraceae (as well as some in the Actinidiaceae) have poricidal anthers, another character used to define the suborder Clethrineae. Poricidal anthers fold in making a tube to protect the pollen from rain and fog. There are pores at the downward end of the anthers where the pollen falls out when the anthers are vibrated by bees. *Heliamphora* has this feature but the pore is not as fully defined as it is in typical poricidal anthers (Renner 1989) so DeBuhr did not observe it in his dried material. Taxonomists like characters like poricidal anthers because they can be seen in dried flowers. Unfortunately they are not really appropriate for use in taxonomy above species level because they are climate and pollinator adaptations that have arisen separately in many plant groups (Renner 1989).

A full understanding of the phylogenetic context of the Sarraceniaceae was not approached until a DNA sequence study done by Albert *et al.* (1992). This study used the DNA sequences of the chloroplast *rbcL* gene to place the carnivorous plant genera among 72 plant families. They used this sequence because it changes slowly and helps resolve relationships well at the genus and family level. The study showed unequivocally that *Nepenthes* is related to *Drosera*, *Dionaea*, *Drosophyllum*, and *Plumbago* while the genera in the Sarraceniaceae are related to *Roridula* as well as *Rhododendron* and *Actinidia* (Fig. 1b). The results of the study showed that gross physical characteristics like those used by Macfarlane cannot be used to determine relationships among carnivorous plant genera. There are too many cases where unrelated carnivores have superficially identical adaptations. They also showed that *Darlingtonia* is the most basal species in the Sarraceniaceae. This result was so surprising that the next major study by Bayer *et al.* (1996) repeated the sequencing of *Darlingtonia* chloroplast *rbcL*. The Bayer *et al.* (1996) study confirmed the Albert *et al.* (1992) study placement of *Darlingtonia* but shuffled around the related genera (Fig. 1c).

The reshuffling in the phylogenies reflects the minimal amount of useful data that can be obtained from studying only one or a few genes. DNA sequences that code for proteins or other cellular components are useful in phylogenetic studies of families and orders because the sequences change slowly over time. The kinds of changes seen are limited by the need of the plant to maintain a functional gene product. The DNA sequences of non-coding regions are useful for studying closely related species. They display more and different kinds of changes resulting in proportionally more useful phylogenetic information. But as species become more distantly related there can be so many changes in non-coding regions that analysis becomes difficult. These tradeoffs mean the sequence regions used in a study need to match the expected relatedness of the taxa and that enough sequence needs to be recovered to make sure there are enough meaningful changes in the data to draw valid conclusions. In addition, studies using nuclear sequences should include more than one individual per species because of the expected polymorphisms within species. As studies get more data and include additional sequence regions it is not surprising the trees change.

Owing to advances in technology Ellison *et al.* (2012) analyzed more than six times the sequence data than previous studies and tweaked the phylogeny further (Fig. 1d). The biggest change is *Roridula* being a sister group to *Actinidia*. The important aspect of this change is it puts *Roridula* and the Sarraceniaceae into an evolutionary speed perspective. *Actinidia* is a member of the Actinidiaceae, a family of about 360 species in three genera. The genera and species of the Actinidiaceae had to evolve after the split with *Roridula* and that had to happen after both split from the Sarraceniaceae. However there is no way of knowing at this point if the Ellison *et al.* (2012) phylogeny will have the last word in the families related to the Sarraceniaceae and their relationships. Future studies with even more data may result in a shuffle of the related families again.

The primary focus of the Bayer *et al.* (1996) study was species level relationships. They used the faster evolving nuclear DNA sequence: ribosomal RNA internal transcribed spacer, or ITS for short. This sequence is usually useful in separating taxa at the species level. Using ITS appeared to work for the three *Heliamphora* species tested but not for the nine *Sarracenia* species in the study. They were not able to recover enough sequence data to uniquely identify species. Neyland and Merchant (2006) made a second attempt at determining the phylogenetic relationships among the *Sarracenia* species using part of the 26S rRNA sequence plus adjacent ITS sequences. With the increased amount of sequence they were able to separate the clade consisting of *S. purpurea* and *S. rosea* from the other species in the genus and that was about it. Ellison *et al.* (2012) acquired even more nuclear sequences adding more 26S rRNA sequence as well as *PHYC*. Taken separately the Ellison *et al.* (2012) data give some hints at further species associations in *Sarracenia* but I do not consider the results significant. The major reason is not enough differences were found between species and they sampled only one individual per sequence per species.

Neyland and Merchant (2006) showed that at least for nuclear sequences it is necessary to sample more than one individual of each species when the species are as closely related as they are in Sarracenia. They reported on six individuals of Sarracenia alata and three of S. leucophylla as well as one individual for each of the other species. Based on the sequence data they posted to NCBI Genbank, they did not find any single site that unambiguously differentiated S. alata and S. leucophylla from the species other than S. purpurea and S. rosea. However three of the S. alata and one of S. leucophylla individuals sampled had unique base substitutions or insertions. This implies you can get a different answer by picking the right, or wrong, individual for DNA sequencing. Schnell and Krider (1976) and McPherson and Schnell (2011) comment on this state of affairs indirectly. They established sets of physical characters that define *Sarracenia* species but cautioned that the character state sets are not appropriate for determining relationships between species. Even more importantly, although they can establish an ideal for each species as they define them, when one goes out into the field, masses of hybrids are found unless a site is remote from other sites and contains only one species. And even in isolated single species sites within the extended Sarracenia *rubra* complex there are individuals that do not necessarily match the type of that particular species or subspecies.

It is now possible to do the kinds of DNA sequencing studies necessary to understand genera like *Sarracenia*. Zellmer *et al.* (2012) performed an ecological genetics demonstration study using "Next-generation" sequencing technologies designed for whole genome sequencing in a very clever way to generate a massive amount of sequence data for *Sarracenia alata*. They sequenced 86 individuals from 10 sites across the species range and compared the genetic divergence between populations to physical barriers such as the Mississippi River basin and correlated the divergence to specific local environmental factors. They estimated the eastern and western populations of *S. alata* have been separated by 60,000 generations (or a minimum of 300,000 years). However in their analysis Zellmer *et al.* (2012) made the assumption that *Sarracenia* species are reproductively isolated from each other. Four other *Sarracenia* species co-occur with *S. alata* in the eastern part of its range. Hybrids involving *S. alata* are commonly found today in Mississippi and Alabama (Sheridan 1991; McPherson & Schnell 2011). It is quite possible past hybridization resulted in introgression of genetic material into *S. alata* from other species. The presence of genetic material from other species in the eastern population would make the length of time the eastern and western populations were separated to appear longer than actually happened.

DNA sequence data provide a molecular clock according to base substitution rates. With proper scaling and reference points one can estimate dates of divergence of taxa. Ellison *et al.* (2012) used

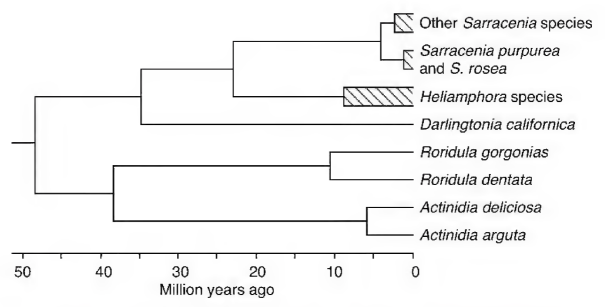


Figure 2: Phylogeny of the Sarraceniaceae calibrated to a time scale. Hatched areas indicate multiple species. Figure adapted from Ellison *et al.* (2012). Estimates of errors in the dates are not shown.

their data to put dates on events within the Sarraceniaceae to help understand the biogeography of the family (Fig. 2). To add dates they used techniques and fossil calibrations from Bell *et al.* (2010). Bell *et al.* (2010) place the first Ericales at about 120 million years ago. Coincidentally that is approximately the age of the possible pitcher plant *Archaeamphora longicervia* of Li (2005). This fossil cannot be a member of the Sarraceniaceae because it is 80 million years too old. Ellison *et al.* (2012) estimated the Sarraceniaceae came into existence between 35 and 48 million years ago. There are large uncertainties in these numbers, some of which result from scaling uncertainty and some from sampling error. Taken at face value the split between the Sarraceniaceae and *Roridula*/ Actinidiaceae clade was about 48 million years ago. The plants most likely did not have pitcher leaves at that point. Unless *Darlingtonia* developed pitchers separately from the other members of its family, by about 35 million years ago the Sarraceniaceae existed as pitcher plants. That is a time span of 13 million years to develop pitchers. The split between *Sarracenia* and *Heliamphora* is placed at about 23 million years ago, 12 million years after the *Darlingtonia* split.

The families related to the Sarraceniaceae generally have a South American focus but are by no means restricted there. *Roridula* is only found in South Africa and the species in the Actinidiaceae extant today are found in eastern Asia as well as Central and South America. During this time, al-though there were no confirmed land bridges between North and South America, based on fossils there was movement of mammals from South America into the Antilles and from North America into South America. These were singular events that happened well before North and South America became connected via Panama 4 million years ago. Ellison *et al.* (2012) proposed that the Sarraceniaceae arose in South America, spread to North America via the Antilles, and then fragmented first between western and eastern North America and then between eastern North America and South America. *Darlingtonia* is what is left of the western North American fragment, *Sarracenia* from the eastern North American fragment, and *Heliamphora* from South American fragment. On the other hand, Bayer *et al.* (1996) argued there is no way to discern where the Sarraceniaceae originated with any confidence. The current distribution of genera could easily be the result of two long distance dispersal events with the first event originating either in North or South America. There are plenty of

examples of plants moving between North and South America during this time without land bridges presumably by migrating birds. Although none of the species in the family have typical adaptations for attracting birds to their seeds, birds would be expected to eat the seeds on the ground. *Darling-tonia* seeds are adapted for dispersal by mammals but that could be a relatively recent development.

Of course there is no way to know the actual history of the Sarraceniaceae without more data than we have today. The results of the studies show long lines of 10 to 35 million years without any branches. That does not mean there was only one species during that time. It means with the data we have there is no evidence of the species that undoubtedly existed but went extinct. Also the branch points are not literally where a new genus or species came into being. The branch points are estimates of when two genetic lines diverged. Those separate lines may not become distinct species until much later. What the data from the Ellison *et al.* (2012) study do tell us are the approximate times of the major splits that led to the genera and species and how long the genera and species as we know them have been separate genetic entities. For *Heliamphora* Ellison *et al.* (2012) studied six species and the oldest branch point is 9 million years ago. Presuming that date holds after a study of the whole genus is published, the genus has diversified from effectively one species into 23 across 800 km in 9 million years. The *Sarracenia* species oldest branch point is 4 million years ago and the genus developed from effectively one species into 15 species and subspecies across 2000 km since that time.

In spite of these results showing very fast diversification in Heliamphora and Sarracenia, Ellison et al. (2012) proposed that a species was able to sufficiently exchange genetic material across the 3500 km between North and South America to remain one species over 12 million years. And what about the 14 and 19 million year intervals between the split between the genera and the recent diversification? If Heliamphora was really in the Guiana Highlands before the diversification was there an extinction event that reduced the genus to one species? Or were they somewhere else where they went extinct and we now only see the results of a long distance dispersal event 9 million years ago? The only evidence I see weighing on this issue is all *Heliamphora* species have an adaptation found in high elevation equatorial plants: the leaves totally wrap the apical meristem and protect it and emerging leaves from nighttime frosts. McPherson (2008) explains it is unlikely the tepuis experienced extensive freezes or frosts as freezing would have accelerated the demise of the tepuis. So why would *Heliamphora* have the feature unless it lived in an area that did get frosts at random times during the year? Darlingtonia and Sarracenia do not protect their meristem in the same way because they are adapted to a temperate environment and go dormant seasonally. For the Sarracenia phylogeny the situation is similar to *Heliamphora* with a long branch and then a recent explosion. If Sarracenia as we know it was in what today is the southeast USA it was effectively reduced to one species going forward 4 million years ago. If it was somewhere else, that is when it arrived.

Ellison *et al.* (2012) were not able to discern much structure within *Sarracenia* but within *Heli-amphora* from their limited data it appears there is no correlation between physical distance between species and genetic distance. For instance they show *Heliamphora neblinae* and *H. pulchella* to be separated temporally by only 600,000 years but they are separated spatially by 600 km with no current *Heliamphora* habitat in between while *H. minor* and *H. heterodoxa*, 55 km apart, are separated genetically by the 9 million years. This implies that the *Heliamphora* species are capable of long distance dispersal, and are not relics that speciated in place as the tepuis eroded. These results are similar to the DNA study of Kok *et al.* (2012) for frogs and lizards in the Guiana Highlands and are consistent with the biogeographic studies of the Guiana Highland flora by Huber (1988) and birds by Mayr and Phelps (1967). Each of these studies discussed in general terms the implications of their work but Rull (2005) proposed an explicit model explaining their results where the biota of

Guiana Highlands have undergone elevation shifts during global climate cycles leading to mixing of species at lower elevations off the tepui tops during colder periods followed by isolation back on the tepui summits during interglacial times.

With each step in the story of the Sarraceniaceae it has gotten more complex and if anything more confusing. The comparative taxonomy got us nowhere and one could even begin wondering at what point the family developed effective pitchers and if it did so more than once. There was certainly plenty of time for independent development. The molecular genetics told us that in spite of deep roots the current set of species in *Heliamphora* and *Sarracenia* are very recent. Because all *Heliamphora* and *Sarracenia* species are interfertile and hybridization and introgression have probably been major factors in the diversification of the genera, a deeper study of their genomes using the techniques of Zellmer *et al.* (2012) with wild collected samples of multiple individuals in each population and large amounts of random sequences is necessary to go the next step in understanding evolution within these genera. The *Sarracenia* study has to be done soon before populations of pure *Sarracenia* has devolved into a massive hybrid swarm.

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