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Phylogeny, classification and biogeography of *Philothea* sect. *Erionema* (Rutaceae) based on nrDNA sequences

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ABSTRACT

Philothea sect. *Erionema* includes 14 species from eastern Australia and one from south-western Australia. We conducted a phylogenetic analysis of the section, including samples of all species, using sequences of the ITS and ETS regions of nuclear ribosomal DNA. Results were broadly congruent with a previous analysis based on morphological and flavonoid data. The analysis is consistent with the monophyly of the section and supports the monophyly of six species represented by multiple samples. *Philothea verrucosa* (A. Rich.) Paul G. Wilson was resolved as paraphyletic with respect to *P. freyciana* Rozefelds but with poor support. *Philothea glasshousiensis*, *P. myoporoides* and *P. myoporoides* subsp. *myoporoides* were clearly polyphyletic, including separate geographic clades and the classification of each of these taxa requires revision. In particular, disjunct northern populations of *P. glasshousiensis* probably represent a distinct species, the five subspecies of *P. myoporoides* could be treated as separate species and at least two other distinct groups that are currently included under the circumscription of subsp. *myoporoides* could be treated as species. The phylogeny revealed deeply divergent, geographically overlapping clades in eastern Australia and substantial distances (up to 900 km) between sister taxa. We infer that biogeography of the group has been shaped largely by vicariant differentiation of taxa.

Keywords: Australia, biogeography, nrDNA, *Philothea freyciana*, *Philothea glasshousiensis*, *Philothea myoporoides*, phylogeny, Rutaceae, taxonomy.

Introduction

Philothea sect. *Erionema* (F.Muell.) Paul G. Wilson is one of four sections recognised in the Australian genus *Philothea* Rudge (Wilson 2013). The section currently includes 14 species (21 taxa, including subspecies) from eastern Australia and 1 species, *P. brucei*¹ (with 3 subspecies) from south-western Australia (Fig. 1). This is distinguished from other sections of the genus primarily on the basis of the that have two (or rarely several) glands at the base of the white anther apiculum and by the seeds that are ellipsoid, laterally flattened, have a linear hilum on the adaxial face and have a basal chalazal region (Wilson 1998, 2013). Members of the section are insect pollinated (Armstrong 1991) shrubs or small trees that occur in open eucalypt forests or woodlands or in heathlands, sometimes associated with rock outcrops (Wilson 2013).

Classification of sect. *Erionema* has seen several changes in the last two decades. The section was transferred from *Eriostemon* Sm. (*sensu* Wilson 1970) to *Philothea* (Wilson 1998) based on Armstrong's (1991) morphological phylogenetic analyses of Australian Rutaceae. Circumscription of the section was amended by Wilson (1998) to include the Western Australian (WA) species, *P. brucei* (previously included in a monotypic section, *Eriostemon* sect. *Osmanthos* Paul G. Wilson), on the basis of shared anther and seed morphology. New taxa have also been assigned to sect. *Erionema* including: a new species, *P. freyciana* that was segregated from *P. verrucosa* by Rozefelds (2001a);

¹Author names for species and subspecies of *Philothea* are given in Table 1.

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a new subspecies of *P. buxifolia* described by Wilson (1998); and a number of new taxa or combinations that were published in a complex of species related to *P. myoporoides* (Bayly 1998; Rozefelds 2001b; Forster 2005). This last group, in particular, has been taxonomically problematic, with up to 10 largely allopatric subspecies being recognised under a widespread and broadly circumscribed *P. myoporoides* (Rozefelds 2001b). The five Queensland (Qld) subspecies of this group were elevated to species rank by Forster (2005) but that treatment did not consider the status of the remaining subspecies, resulting in markedly different species concepts for taxa on either side of the New South Wales (NSW)–Qld state border.

Relationships in sect. *Erionema* have been assessed using datasets based on morphology and leaf flavonoids (Bayly 2001). Those analyses supported monophyly of the section, including *P. brucei* but many relationships were not well resolved. In particular, most members of the *P. myoporoides* group (including currently accepted subspecies and recent segregate species) formed a large polytomy that also included *P. verrucosa*, *P. scabra*, *P. hispidula* and *P. buxifolia*. Relationships inferred among that group were based largely on homoplasious morphological characters.

A small number of *Philotheca* species have been included in molecular phylogenetic studies assessing higher-level relationships of Australian Rutaceae (Chase *et al.* 1999; Scott *et al.* 2000; Gropo *et al.* 2008; Salvo *et al.* 2010; Bayly *et al.* 2013; Morton and Telmer 2014; Appelhans *et al.* 2021) or as outgroups in analyses of other genera (Othman *et al.* 2010; French *et al.* 2016). At most, these studies have included one species of sect. *Erionema*. As such, there are currently no molecular data for assessing the circumscription of this section or the relationships of taxa within this.

The current study uses newly generated sequences of the internal transcribed spacers (ITS) and external transcribed spacer (ETS) of nuclear ribosomal DNA to assess relationships among members of *Philotheca* sect. *Erionema*. The aims of this study were: (1) to test the monophyly of the section, in particular the inclusion of *P. brucei*; (2) to test the classification of species and subspecies, especially those associated with *P. myoporoides*; and (3) to gain insight into the biogeographic history of this group in eastern and south-western Australia.

Materials and methods

Taxon sampling and DNA sequencing

Plant material was obtained from field collections, cultivated plants of known provenance, and in a few cases from existing herbarium specimens (Table 1).

In total, 49 ingroup samples were used, representing all 15 species of sect. *Erionema* (19 of the 24 currently accepted taxa including segregate subspecies; the 5 subspecies not

sampled were *Philotheca brucei* subsp. *brevifolia* and subsp. *cinerea*, *P. buxifolia* subsp. *falcata* and subsp. *obovata*, and *P. scabra* subsp. *latifolia*). Initial results indicated that *P. myoporoides* subsp. *myoporoides*, the most geographically widespread subspecies in the section, was genetically heterogeneous. This taxon was therefore sampled more intensively using material from 19 different locations. Representatives of the three other sections of *Philotheca* were used as outgroups: *P. coccinea*, *P. glabra* and *P. thryptomenoides* from sect. *Philotheca*; *P. fitzgeraldii* from sect. *Corynonema* (Paul G. Wilson) Paul G. Wilson; *P. spicata* from sect. *Cyanochlamys* (Bartl. ex F. Muell.) Paul G. Wilson.

DNA isolation, amplification and sequencing of ITS and ETS followed the methods described by Bayly *et al.* (2015).

Sequence editing and alignment

Contiguous sequences for each region were assembled and edited using Sequencher (ver. 4.8, Gene Codes Corporation, Ann Arbor, MI, USA) or Geneious (ver. 9, see <https://www.geneious.com/>; Kearse *et al.* 2012). Sequencing chromatograms were carefully scrutinised for double signal peaks that could indicate variation among the many copies of ITS and ETS regions that are present in the genome of each plant. Standard IUPAC nucleotide ambiguity codes were used for any positions in the sequence showing evidence of such polymorphisms. Most sequences included some polymorphic sites (fewer than ten across the entire span of the ITS + ETS regions) but two samples had very high numbers, namely *Philotheca myoporoides* subsp. *brevipedunculata* and *P. buxifolia*, with 29 and 27 polymorphic sites respectively across the combined rDNA sequences. These two sequences were retained in the dataset so that these taxa could be represented in analyses, although we recognise that the large number of polymorphic nucleotide sites in these sequences could create uncertainty about their phylogenetic placement. In both of these samples, the 5.8S rDNA region was highly conserved when compared with the flanking ITS-1 and ITS-2 regions (showing no differences in *P. buxifolia* and only one ambiguous base in *P. myoporoides* subsp. *brevipedunculata*), making the recovered sequences unlikely to include rDNA pseudogenes (see Bailey *et al.* 2003).

Sequences were aligned in Geneious with some manual adjustment. Insertion–deletion events (INDELs) were coded using the ‘simple INDEL coding’ approach of Simmons and Ochoterena (2000), with a single character appended to the end of the data matrix, representing each INDEL, whether single- or multi-base. The final alignment, including INDEL characters (identified as distinct ITS and ETS INDEL CHARSETs in the nexus file), is deposited in TreeBase (see <http://treebase.org/>; study accession number 26368).

Phylogenetic analyses

Combined ITS and ETS sequences were analysed using maximum parsimony (MP) as implemented in PAUP* 4.0 beta

Table 1. Voucher specimens and GenBank accession numbers for sequences analysed in the study.

Taxon	Location	Coll no.	Voucher	ITS	ETS
<i>Philothea</i> sect. <i>Erionema</i>					
<i>P. brucei</i> (F.Muell.) Paul G.Wilson subsp. <i>brucei</i>	WA, Great Western Hwy 88 km SW of Paynes Find	MJB1918	MEL 2383578A	–	MK109942
<i>P. brucei</i> subsp. <i>brucei</i>	WA, 37 km N of Cleary on Mouroubra Rd	MJB1938	MEL 2383572A	MK109993	MK109943
<i>P. buxifolia</i> (Sm.) Paul G.Wilson subsp. <i>buxifolia</i>	NSW, Royal NP, 1.3 km along Sir Bertram Stevens Dr from the intersection with Bundeena Ave	ELB13a	MELU D105835	MK109967	MK109916
<i>P. conduplicata</i> (Paul G.Wilson) P.I.Forst.	Qld, Girraween NP, 2.3 km E of park headquarters	MTM377	BRI AQ0755781	MK109989	MK109939
<i>P. conduplicata</i>	ACT, cult. ANBG (loc. 112, propagation ID 617438)	MJB2009	MELU D105851	MK109991	MK109940
<i>P. epilosa</i> (Paul G.Wilson) P. I.Forst.	NSW, Boonoo Boonoo NP	MJB269	MELU D105982	–	MK109945
<i>P. epilosa</i>	Qld, Girraween NP, upper Bald Rock Creek	PIF34471	BRI AQ0745288	MK109995	MK109946
<i>P. freyciana</i> Rozefelds	Tas., Freycinet NP, Mt Mayson West (cult. Royal Tasmanian Botanical Gardens)	NP412	MELU D108699	MK109992	MK109941
<i>P. glasshousiensis</i> (Domin) P.I.Forst.	Qld, Mt Cooroora, Pomona	MTM608	BRI AQ0846021	MK109968	MK109917
<i>P. glasshousiensis</i>	Qld, Kroombit Tops NP, Helipad on side track off Razorback Ridge	MTM643	BRI AQ0846033	MK109969	MK109918
<i>P. glasshousiensis</i>	Qld, Kroombit Tops NP, clifftops near radio towers, ~2 km NW of barracks	MTM293	BRI AQ0813599	–	MK109949
<i>P. glasshousiensis</i>	Qld, Glass House Mountains NP, Mt Coonowrin (Crookneck)	PIF34310	BRI AQ0744216	MK109998	MK109950
<i>P. hispidula</i> (Sieber ex Spreng.) Paul G.Wilson	NSW, Blue Mountains NP, adjacent to the walking track near the car park for Jellybean Pool	PHW3326	MEL 2396738A	MK109970	MK109919
<i>P. hispidula</i>	NSW, Thirlmere Lakes NP, Lake Werri Berri	ELB12a	MELU D105834	–	MK109951
<i>P. myoporoides</i> (DC.) Bayly subsp. <i>myoporoides</i>	NSW, Cox River, South Bowenfels		CBG 8742	–	MK109914
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	Vic., Mt Donna Buang Rd	MJB2025E	MELU D105829	MK109971	MK109920
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	Vic., Alpine NP, north-east of Lake Tali Karng on Gillios Tk	MJB2026C	MELU D105824	MK109972	MK109921
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	Vic., Briagolong SF	MJB2030D	MELU D105818	MK109973	MK109922
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Merricumbene Forest	L.G.Adams s.n.	CANB 381821	MK109974	MK109923
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Morton NP, The Castle	Craven10056	CANB 632861	MK109975	MK109924
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, 6.5 km E of Reids Flat, on road to Bigga	S.Donaldson2371	CANB 619523	MK109976	MK109925
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Kosciuszko NP, 7.5 km from turnoff to Olsen's Lookout, on Geehi Rd	ELB1d	MELU D105783	MK109977	MK109926

(Continued on next page)

Table 1. (Continued)

Taxon	Location	Coll no.	Voucher	ITS	ETS
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Tinderry Nature Reserve	ELB2c	MELU D105787	MK109978	MK109927
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Kanangra–Boyd NP	ELB3a	MELU D105791	MK109979	MK109928
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Bents Basin State Conservation Area	ELB5d	MELU D105800	MK109980	MK109929
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW Warragamba Park, Nortons Basin	ELB6d	MELU D105805	MK109981	MK109930
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Bargo-Picton Rd (Remembrance Dr), under bridge over Bargo River	ELB7d	MELU D105810	MK109982	MK109931
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Nullica SF, track to Nethercote Falls	ELB8e	MELU D105816	MK109983	MK109932
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Mumbulla Mountain	N.Fisher129	CANB 544781	MK109984	MK109934
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	Vic., Toolangi SF, Quarry Rd	MJB1973	MELU D105838	MK109985	MK109935
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Blue Mountains NP, Jellybean Pool	PHW3325	MEL 2396739A	MK109988	MK109938
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	Vic., Lake Mountain	MJB1866	MELU D105839	–	MK109953
<i>P. myoporoides</i> subsp. <i>myoporoides</i>	NSW, Blue Mountains NP, Glenbrook causeway	ELB4	MELU D105796	MK110000	MK109954
<i>P. myoporoides</i> subsp. <i>acuta</i> (Blakely) Bayly	NSW, Mt Bunganbil	MJB174	MELU D105846	MK109999	MK109952
<i>P. myoporoides</i> subsp. <i>brevipedunculata</i> Bayly	ACT, cult. ANBG (loc. 46, propagation ID 631817)	MJB2008	MELU D105852	MK109966	MK109915
<i>P. myoporoides</i> subsp. <i>euroensis</i> Bayly	Vic., Garden Range, Euroa	MJB s.n.	MELU D105847	–	MK109933
<i>P. myoporoides</i> subsp. <i>petraea</i> Rozefelds	Vic., Mt Stewart, Gippsland		MEL 0004133A	MK109987	MK109937
<i>P. obovalis</i> (A.Cunn.) Paul G.Wilson	NSW, S of Bell on Lithmore-Windsor Rd	MJB205	MELU D105842	MK110001	MK109955
<i>P. obovatifolia</i> (Bayly) P.I.Forst.	NSW, Mt Werrikimbe		MEL 2278581A	MK109986	MK109936
<i>P. queenslandica</i> (C.T.White) P.I.Forst.	Qld, Tinnanbar via Tin Can Bay–Maryborough Rd	MTM273	BRI AQ0746015	MK109990	–
<i>P. queenslandica</i>	Qld, Great Sandy NP, adjacent to Cooloola Coast Cemetery	PIF34188	BRI AQ 743513	MK110002	MK109956
<i>P. scabra</i> (Paxton) Paul G.Wilson subsp. <i>scabra</i>	NSW, Georges River NP, Picnic Point	ELB11	MELU D105836	MK110003	MK109957
<i>P. trachyphylla</i> (F.Muell.) Paul G.Wilson	Vic., Boggy Creek, near bridge on Princes Hwy, Nowa Nowa	MJB1900	MELU D105850	MK110005	MK109959
<i>P. trachyphylla</i>	NSW, Deua NP, road from Araluen to Moruya, 5 km past Kennys Creek bridge	ELB10	MELU D105837	–	MK109960
<i>P. verrucosa</i> (A.Rich.) Paul G.Wilson	Vic., cult. Heidelberg, ex. Mt Difficult, Grampians	MJB2475	MELU D112272	KU377587	MK109961
<i>P. verrucosa</i>	Vic., Mt Arapiles	WN7a	MELU D108908	MK110007	MK109963
<i>P. verrucosa</i>	Vic., Brisbane Ranges, DeMotts Rd ~300 m W of Clarkes Rd	MJB2199	MELU D121711	MK110008	MK109964

(Continued on next page)

Table 1. (Continued)

Taxon	Location	Coll no.	Voucher	ITS	ETS
<i>P. verrucosa</i>	Tas., East Risdon Nature Reserve	WN17a	MELU D108906	MK110009	MK109965
<i>P. virgata</i> (Hook.f.) Paul G.Wilson	Vic., Cooracambra NP, Mt Kaye walking track	MJB266	MELU D105843	MK110006	MK109962
<i>P. sect. Corynonema</i>					
<i>P. fitzgeraldii</i> (C.R.P.Andrews) Paul G.Wilson	WA, near intersection of Lake King–Norseman Rd and Coolgardie–Esperance Hwy	MJB1942	MEL 2383574A	MK109996	MK109947
<i>P. sect. Cyanochlamys</i>					
<i>P. spicata</i> (A.Rich.) Paul G.Wilson	WA, Alexander Morrison NP, Tootbardie Rd, 1.3 km south from intersection with Coorow–Green Head Rd	MJB1907	MEL 2383588A	KU861302	KU861261
<i>P. sect. Philotheca</i>					
<i>P. coccinea</i> (C.A.Gardner) Paul G.Wilson	WA, Boorabbin NP at Boorabbin (Koorarawalyee) Rest Area	MJB1929	MEL 2383614A	MK109994	MK109944
<i>P. glabra</i> (Paul G.Wilson) Paul G.Wilson	WA, Great Northern Hwy, 1 km S of White Wells turnoff	MJB1917	MEL 2383617A	MK109997	MK109948
<i>P. thryptomenoides</i> (S.Moore) Paul G.Wilson	WA, Great Northern Hwy, 93.8 km SW of Paynes Find	MJB1921	MEL 2383582A	MK110004	MK109958

Herbarium abbreviations follow Index Herbariorum (<http://sweetgum.nybg.org/science/ih/>). Collector abbreviations: ELB, Erin L. Batty; MJB, Michael J. Bayly; MTM, Michael T. Mathieson; PHW, Peter H. Weston; PIF, Paul I. Forster. Locality abbreviations: ACT, Australian Capital Territory; ANBG, Australian National Botanic Gardens; cult., cultivated; Dr, Drive; Hwy, Highway; Mt, Mount; NP, National Park; NSW, New South Wales; Qld, Queensland; Rd, Road; RBGDT, Royal Botanic Gardens & Domain Trust (Sydney); SF, State Forest; Tas., Tasmania; Tk, Track; Vic., Victoria; WA, Western Australia. GenBank accession numbers in bold indicate sequences that are newly published as part of this study.

(ver. 10, D. L. Swofford, see <https://paup.phylosolutions.com/>) and by Bayesian inference (BI) using MrBayes (ver. 3.1.2, see <https://nbisweden.github.io/MrBayes/>; Ronquist and Huelsenbeck 2003), with and without the presence of INDEL characters. MP analyses were performed using heuristic tree searches, a CLOSEST addition sequence, TBR branch swapping, MAXTREES set at 30 000, all characters equally weighted and gaps treated as missing data. MP Bootstrap analyses used 1000 'full heuristic' replicates with MAXTREES set at 3000 per replicate. Models for BI analyses were selected using the Akaike Information Criterion as implemented in MrModeltest (ver. 2.3, J. Nylander, see <https://github.com/nylander/MrModeltest2>). The chosen models were GTR + Γ + I for ITS and GTR + Γ for ETS; INDELs were analysed under the 'restriction' model. BI analyses used the default settings of MrBayes and each included two runs of four chains, each run for 2 000 000 generations. Trees were sampled every 1000 generations and a majority rule consensus was computed (with trees from the first 500 000 generations discarded as burn-in). To determine that the runs had converged on a stationary distribution and that the burn-in period was adequate, the distribution of likelihood values in Tracer (ver. 1.5, A. Rambaut, see <https://beast.community/tracer.html>) and the standard deviation of split frequencies (that was < 0.01 at the end of the runs) were evaluated.

Results

The combined nrDNA dataset included 1279 characters, of which 365 were variable and 237 were parsimony informative (119 parsimony informative characters from ITS, including 4 INDEL characters and 119 from ETS, including 6 INDEL characters). BI and MP analyses of the combined ITS and ETS dataset produced congruent results, as did analyses with and without INDEL characters included, therefore only the BI tree based on the analysis including INDEL characters is presented here (Fig. 2), showing both Bayesian posterior probabilities (PP) and parsimony bootstrap support (BS) values. The MP analysis (not shown) produced 30 000 equally parsimonious trees with a length 650 steps and a consistency index of 0.66.

The analysis (Fig. 2) is consistent with the monophyly of sect. *Erionema* (PP 0.99, BS 84%) as currently circumscribed, including the WA species *P. brucei*. The basal nodes in the BI tree are not well supported but both *P. brucei* and a clade of three species (*P. obovalis*, *P. virgata* and *P. trachyphylla*) clearly sit outside a large, well-supported clade (PP 1, BS 92%) that includes all other taxa. That large clade is referred to here as the 'pedunculate clade' because the members are characterised by inflorescences with pedicels (one to many) borne on a peduncle; in contrast, pedicels in *P. brucei*, *P. obovalis*, *P. virgata* and *P. trachyphylla* arise directly from the subtending leaf axil and lack a distinct peduncle. Pedunculate, cymose, axillary inflorescences are inferred to

be apomorphic within sect. *Erionema* and within the genus based on the morphological analysis of Bayly (2001).

Within the pedunculate clade, a well-supported basal dichotomy separates a clade of *P. queenslandica* sister to two samples of *P. glasshousiensis* from all other samples.

Ten taxa in the analysis were represented by multiple accessions that allowed their monophyly to be tested. Among these, six were supported as monophyletic, namely *P. trachyphylla* (PP 1, BS 87%), *P. brucei* (PP 1, BS 100%), *P. queenslandica* (PP 1, BS 100%), *P. conduplicata* (PP 1, BS 100%), *P. hispidula* (PP 1, BS 92%), and *P. pilosa* (PP 1, BS 100%). *Philothea verrucosa* was resolved as paraphyletic with respect to *P. freyciana* but support for that relationship was lacking with PP of 0.79 and BS < 50%. Three taxa were notably resolved as polyphyletic, namely *P. myoporoides* (with the currently recognised subspecies spread across multiple clades), *P. myoporoides* subsp. *myoporoides* and *P. glasshousiensis*.

Samples of *P. myoporoides* subsp. *myoporoides* fell into two distinct parts of the nrDNA tree, separated by well-supported nodes. These two clusters of samples have a geographic basis. Neither cluster was resolved as monophyletic, with samples being placed at polytomous nodes that also included other taxa. The two clusters are identified here (Fig. 2) as the 'southern' group that includes samples from the montane highlands of eastern Victoria (Vic.) and southern NSW, and the 'northern' group that includes samples from near Sydney to the South Coast of NSW, generally at lower altitudes.

The samples of *P. glasshousiensis* were also resolved in two distinct geographic clades. These are identified here as the 'southern' samples (Fig. 2) from the Glasshouse Mountains and Mount Cooroora, and the 'northern' samples from Kroombit Tops National Park.

Discussion

Comparison of nrDNA phylogeny with previous analyses based on morphological and flavonoid data

The nrDNA phylogeny for sect. *Erionema* is largely congruent with that of Bayly (2001) that was based on morphological and leaf flavonoid characters. The degree of congruence between the two studies using different datasets is highly unlikely to result from chance and adds confidence to interpretations of relationships. Here we briefly summarise some of the key points of congruence and incongruence between results of the two studies, with the latter largely relating to nodes that are poorly supported in one study or the other.

Both the current molecular and the earlier morphological and flavonoid studies resolved the pedunculate clade as a group, outside of which *Philothea brucei* from WA, and *P. obovalis*, *P. trachyphylla* and *P. virgata* from south-east Australia, were resolved as early diverging lineages.

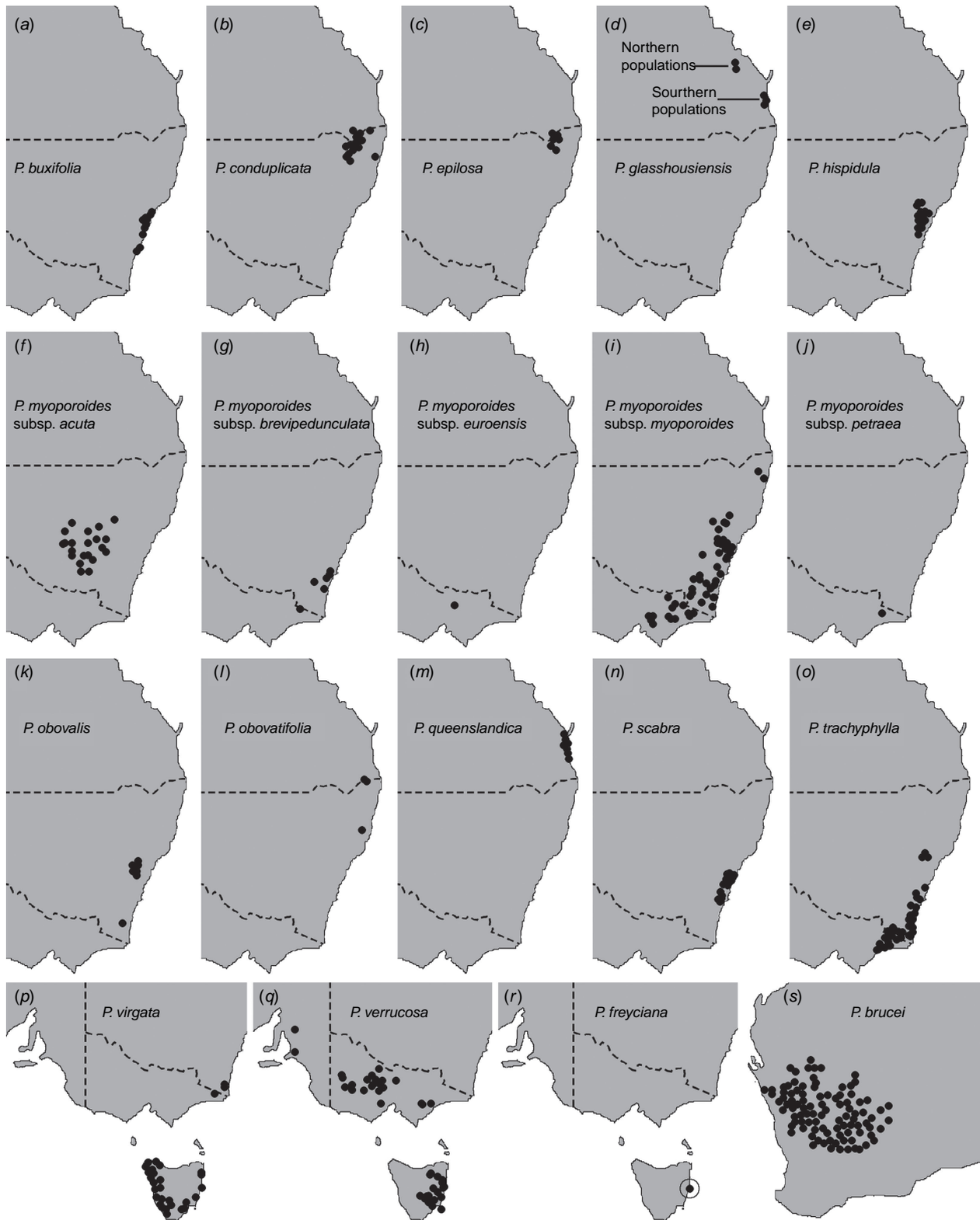


Fig. 1. Distributions of taxa of *Philotheca* sect. *Erionema*. Maps are based on those of [Wilson \(2013\)](#) and [Rozeffelds \(2001a, 2001b\)](#) plus our examination of herbarium material. Note that infraspecific taxa for *P. buxifolia*, *P. brucei* and *P. scabra* are not mapped separately.

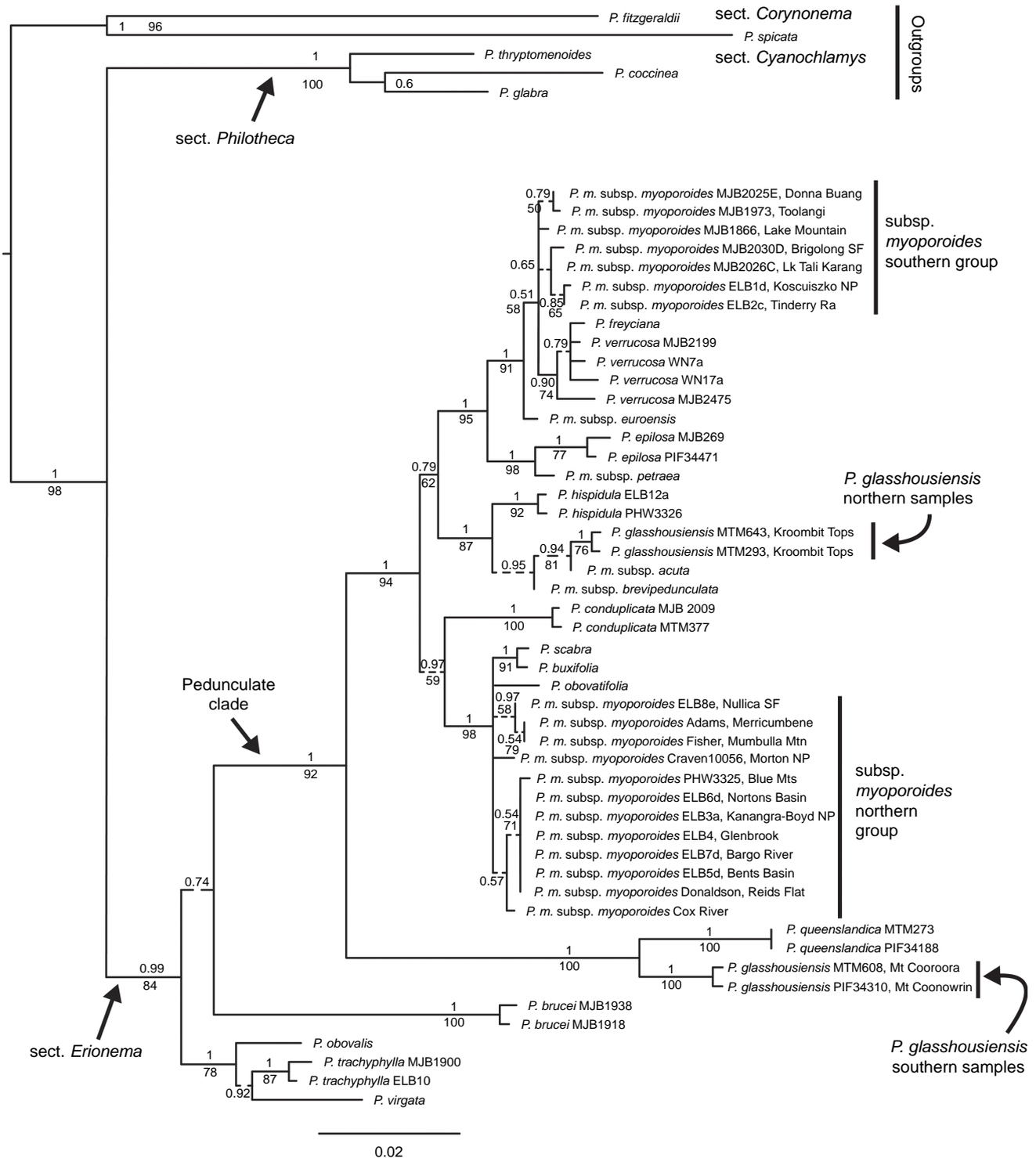


Fig. 2. Majority rule consensus tree based on BI analysis of combined ITS and ETS dataset, showing mean branch lengths. Bayesian posterior probabilities are shown (either first, or above branches) along with MP bootstrap values (either second or below branches; only shown where bootstrap support is >50%). Dashed branches were not present on the MP strict consensus tree.

Relationships among those branches were slightly different in the two studies. In the morphological and flavonoid study (Bayly 2001), *P. obovalis* and *P. virgata* formed a clade that was sister to the rest of the section, one node below

P. trachyphylla in the tree; Bayly (2001) incorrectly indicated that *P. obovalis* and *P. virgata* were united by having 4-merous flowers (a state found consistently only in *P. virgata*) but the two do share the trait, unique in the genus, of having a

reduced number of carpels (<5) per flower (Wilson 1970, 2013). In the nrDNA tree, *P. obovalis* and *P. virgata* were placed with *P. trachyphylla* in a well-supported clade (PP 1, BS 78%) in which *P. trachyphylla* was resolved, but not well supported (PP 0.92, BS <50%), as sister to *P. virgata*.

Within the pedunculate clade, the nrDNA tree provided better resolution than Bayly's (2001) tree based on morphological and flavonoid data. Also, by having multiple accessions for many taxa in the present study, the nrDNA tree has highlighted relationships that could not be investigated in the morphological and flavonoid analysis, in which each taxon was represented by a single aggregate unit. Within the pedunculate clade, key points of congruence between the two studies are: (1) the placement of *P. queenslandica* + southern populations of *P. glasshousiensis* together as a clade sister to all other taxa; (2) the nesting of *P. verrucosa*, *P. buxifolia*, *P. scabra* and *P. hispidula* within *P. myoporoides*. A difference between the studies is that the morphological and flavonoid tree resolved *P. buxifolia*, *P. scabra* and *P. hispidula* as a clade, whereas the nrDNA tree showed *P. hispidula* as well separated from a *P. buxifolia* + *P. scabra* clade, with strong support. In this case, the relationships in the morphological and flavonoid tree were based entirely on homoplasious characters (see fig. 5.15 in Bayly 2001).

Implications for taxonomy

Polyphyly of *Philothea myoporoides*

A significant result from the phylogenetic study presented here is the strong evidence that *Philothea myoporoides* is not monophyletic either in the current circumscription (Forster 2005; Wilson 2013) or the previously broader circumscriptions of Wilson (1970), Bayly (1998) or Rozefelds (2001b). The current circumscription of *P. myoporoides* should not be maintained and our results add weight to the argument for elevating the remaining subspecific taxa to the rank of species, as Forster (2005) did with the subspecies from south-eastern Qld. This would entail raising the segregate subspecies (subsp. *acuta*, subsp. *brevipedunculata*, subsp. *euroensis* and subsp. *petraea*) to species rank. Such an elevation in the rank of these taxa is supported by their generally allopatric distributions (Fig. 1) and generally distinct morphology from each other (albeit largely in leaf shape and size; Bayly 1998; Rozefelds 2001b), and from other species in the genus. Although the monophyly of each of these taxa was not explicitly tested in the current study, each being represented by a single accession, they are all resolved in distinct positions in the phylogeny. For subsp. *acuta*, an earlier name at species rank, *Eriostemon affinis* Sprague, is available (Bayly 1998; Wilson 2013) but there are no existing names at species rank for the other segregate subspecies and therefore new combinations or new names are required.

An alternative response to the polyphyly of *P. myoporoides* would be to retain a broad circumscription of the species and to subsume into this, as synonyms or subspecies, all other members of the pedunculate clade (as identified on Fig. 2).

That would require uniting 18 currently accepted taxa under one species, including long-accepted and morphologically distinct species such as *P. verrucosa*, *P. hispidula*, *P. scabra* and *P. buxifolia*. We consider this option as untenable, given the degree of morphological, ecological and genetic differentiation within the pedunculate clade, and such a broad species concept is inconsistent with other species in the family and would seem unparalleled in current classifications of any other group in the Australian vascular flora.

Polyphyly of *Philothea myoporoides* subsp. *myoporoides*

Philothea myoporoides subsp. *myoporoides* is clearly polyphyletic, with samples falling in two well-separated parts of the nrDNA tree (Fig. 2). These two genetic groups are geographically and ecologically separated and align, at least partly, with morphological variants previously discussed by Wilson (1970, 2013) and Bayly (1998). In Fig. 3 we have

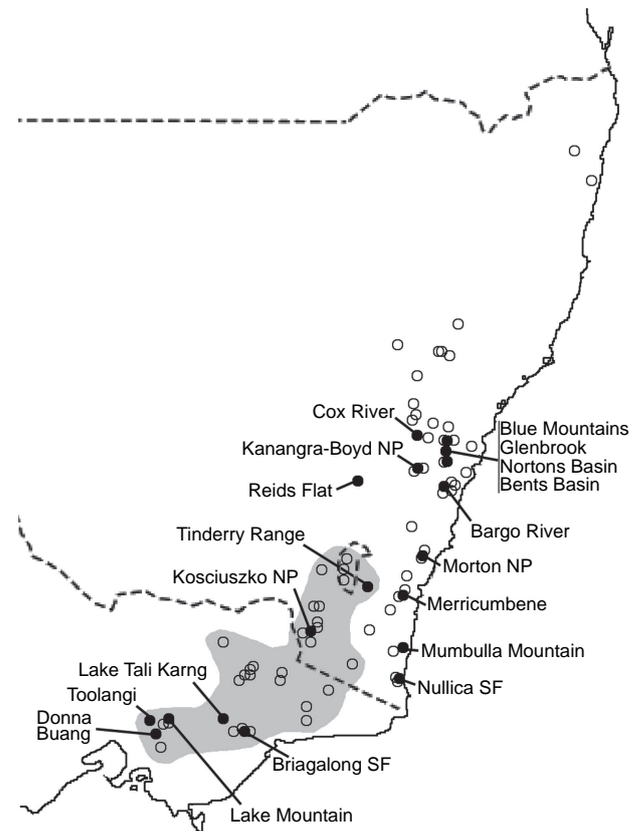


Fig. 3. Distribution of *Philothea myoporoides* subsp. *myoporoides*, including the distribution of samples used in this study (filled circles, with locality names given) and the nrDNA groups these belong to. Grey shading shows the distribution of specimens that are morphologically assigned to the 'southern' nrDNA group; samples from other areas are assigned here to the 'northern' group. Map is based on records in the Australasian Virtual Herbarium and our examination of specimens from MEL, NSW and CANB (primarily in terms of leaf shape, leaf size and habitat). NP, National Park; SF, State Forest.

mapped the distribution of the two genetic groups and have also indicated, based on preliminary examination of herbarium material, what we infer to be the geographical ranges of these groups.

The ‘southern’ genetic group includes the samples from the highlands of Vic., southern NSW and the Australian Capital Territory (ACT). These typically grow in montane or subalpine forests, commonly in Vic. in forests dominated by Mountain Ash (*Eucalyptus regnans* F.Muell.) or Alpine Ash (*Eucalyptus delegatensis* R.T. Baker), often around granite outcrops but also on other substrates. This group equates to the ‘mountain form’ of subsp. *myoporoides* discussed by Wilson (1970, 2013) and Bayly (1998). The earliest species name relating to this group is *Eriostemon lancifolius* F.Muell. Plants of this group often have leaves that are broader, relative to their length, than those of the ‘northern’ genetic group but are variable in leaf shape and size (e.g. Fig. 4).

The ‘northern’ nrDNA group in subsp. *myoporoides* is restricted to NSW and is disjunct from populations of the ‘southern group’ (Fig. 3). It occurs in a range of habitats, mostly at lower altitudes than populations of the ‘southern group’ and is morphologically variable. The samples in our dataset from the Blue Mountains, Glenbrook, Kanangra–Boyd National Park, Bargo River, Bents Basin and Nortons Basin (all from the Central Coast and Central Tablelands of NSW, along water courses, mostly at low altitudes), morphologically match the type of *P. myoporoides* and have relatively long, linear leaves (Fig. 4). Other samples in the ‘northern’ nrDNA

group have different leaf tips and the leaf shapes are relatively short and broad. These include the sample from Cox River that morphologically resembles the type of *Eriostemon cuspidatus* A.Cunn. from the same locality. Other samples with shorter or broader leaves include that from Reids Flat near Bigga in the Central Tablelands and samples from rocky areas in hills and escarpments of the South Coast or Southern Tablelands regions (Merricumbene Forest in Deua National Park, Morton National Park and Mumbulla Mountain). The last group of specimens occurs in localities close to those of *P. myoporoides* subsp. *brevipedunculata* (in Deua National Park and nearby areas) in similar upland habitats; although approaching that taxon in leaf dimensions they have larger leaves, longer peduncles and distinct nrDNA from the single sample of subsp. *brevipedunculata* included here. Nonetheless, the morphological, ecological and genetic distinctiveness of subsp. *brevipedunculata* in this area could be worthy of further detailed investigation based on more intensive sampling.

The two distinct genetic groups in subsp. *myoporoides* should be recognised as at least two different species. However, further work is required to clearly circumscribe them. Although we assign herbarium samples to morphological groups (as in the maps in Fig. 3), there is substantial morphological variation within these groups, especially the northern one, and despite the morphological extremes being fairly distinct, there is a lack of clear separation in leaf attributes (or reproductive features) when all specimens are considered.

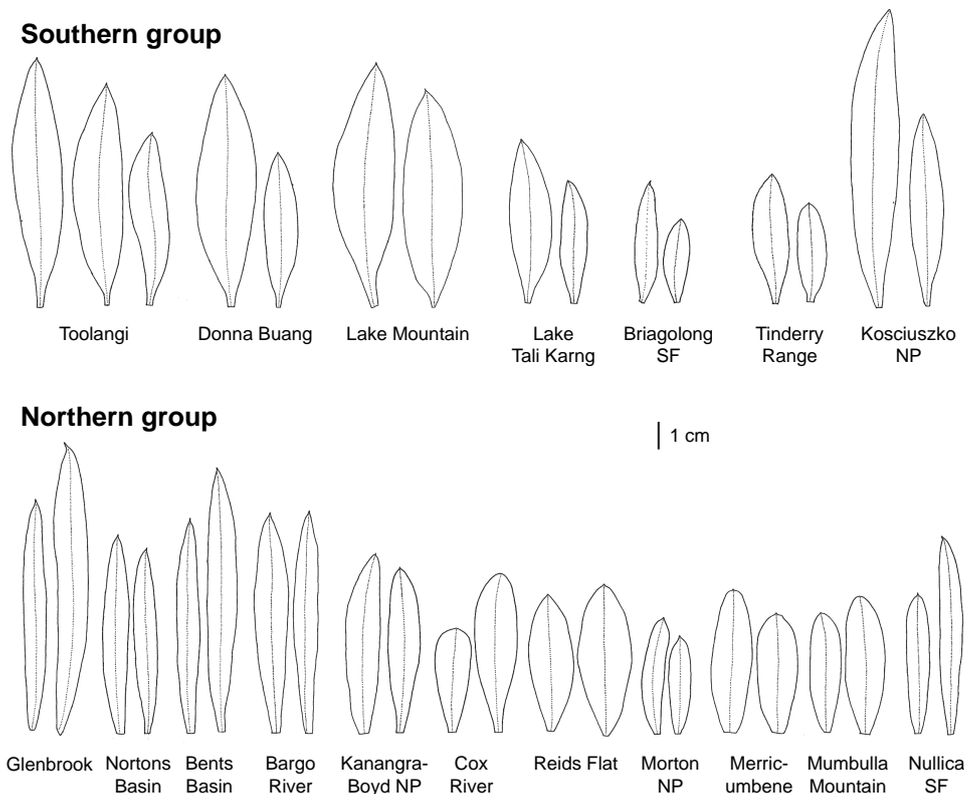


Fig. 4. Examples of leaf variation within the two nrDNA groups in *Philotheca myoporoides* subsp. *myoporoides*. Drawn from representative leaves of samples used in this study. NP, National Park; SF, State Forest.

Also, in the analysis here, neither the 'southern' nor 'northern' nrDNA groups in subsp. *myoporoides* was resolved as monophyletic. This could be due to a lack of signal in the ITS and ETS markers or, amongst other hypotheses, it could reflect the presence of multiple taxa in these nrDNA groups. Other species that group closely with the genetic groups of subsp. *myoporoides* in the nrDNA tree (e.g. *P. scabra*, *P. buxifolia* and *P. obovatifolia* with the 'northern' nrDNA group, and *P. verrucosa* with the 'southern group') are morphologically distinct, to the extent that they could not reasonably be treated as conspecific. Further resolution of both genetic relationships and morphological variation in the nrDNA groups of subsp. *myoporoides* is needed to inform the delimitation of taxa.

Polyphyly of *Philothea glasshousiensis*

The polyphyly of *P. glasshousiensis* provides strong evidence that northern and southern populations should be recognised as two distinct species. The southern populations, from the Glasshouse Mountains and Mount Cooroora, represent *P. glasshousiensis sensu stricto*, the type being from the Glasshouse Mountains (Mount Coonowrin). These southern populations are most closely related to but clearly distinct from *P. queenslandica*, being separated by long branches on the phylogenetic tree (Fig. 2). Although these two species both occur in south-eastern Qld, there is a clear ecological separation between them, with *P. glasshousiensis* growing in rocky areas towards the summits of the mountains and *P. queenslandica* restricted to lowland heaths in wallum vegetation that is periodically inundated.

Herbarium specimens from the northern populations, at Cania Gorge (not sequenced here) and Kroombit Tops, were included in the circumscription of *P. glasshousiensis* (or *P. myoporoides* subsp. *leichhardtii* (Benth.) Paul G. Wilson) by Bayly (1998) and Forster (2005) on the basis of morphological resemblance and their occurrence in similar cliff line habitat. Our recent, preliminary comparisons indicate that, although the northern populations resemble *P. glasshousiensis sensu stricto* in most qualitative features (many of which are highly conserved across sect. *Erionema*), they usually have larger leaves than those of *P. glasshousiensis sensu stricto* but there is overlap in leaf sizes. More detailed study is needed to clarify the extent to which the two genetic groups can be distinguished morphologically.

The status of *Philothea freyciana* warrants further investigation

Philothea freyciana was described by Rozefelds (2001a) for populations from Freycinet Peninsula, Tasmania (Tas.), that were previously included in the more widespread species *P. verrucosa* (e.g. Wilson 1970; as *Eriostemon verrucosus* A.Rich.). Rozefelds (2001a) distinguished *P. freyciana* from *P. verrucosa* on the basis of habit, leaf size and anther apex shape. Because of its limited distribution and small population sizes, *P. freyciana* is listed as Endangered under

both Australia's *Environment Protection and Biodiversity Conservation Act 1999* and Tasmania's *Threatened Species Protection Act 1995*.

Our analysis placed the single sample of *P. freyciana* in a clade with weak–moderate support (PP 0.90, BS 74%) with samples of *P. verrucosa* from Vic. and Tas. There is low sequence variation among samples in this clade and the BI consensus tree suggests that *P. freyciana* could be nested within *P. verrucosa*, although with little support (PP 74, BS <50%). Given this result, and that morphological differences between the two species are slight (Duretto 2009), their relationships and genetic distinctiveness are worthy of further investigation. This seems especially worthwhile given the conservation listings of *P. freyciana* and the potential conservation funding that might be spent to preserve it. Such a study, using additional samples and genetic markers, is currently underway (W. Neal in prep.).

Implementation of taxonomic changes

Although our results highlight the need for several taxonomic changes in sect. *Erionema*, these changes are not formally implemented here. One reason for this is that further morphological study is needed to assist with the delimitation of some taxa that are identified here on genetic grounds; this particularly applies to the two genetic groups within *P. myoporoides* subsp. *myoporoides* and to recognising the northern populations of *P. glasshousiensis* as a distinct species. The other reason is that the generic circumscription of *Philothea* is uncertain and we would prefer not to create new names or combinations for taxa until that is resolved. Our unpublished data, and previous studies (Bayly *et al.* 2013), strongly suggest that *Philothea* is not monophyletic but relationships of the four sections of *Philothea* to each other and to related genera are yet to be clarified. It is possible that sect. *Erionema* might be recognised as a genus distinct from *Philothea*. Some of us (MJB and collaborators) are working to resolve relationships of this group and to propose a revised generic classification. Our intention is that new names or combinations for members of sect. *Erionema* (e.g. raising the segregate subspecies of *P. myoporoides* to species rank) would be published as part of that work, rather than creating additional, potentially briefly used names in the interim.

Implications for biogeography

The relationships resolved here in sect. *Erionema* indicate some interesting biogeographic patterns. These include: (1) the disjunction between western and eastern Australia (between *Philothea brucei* and all other taxa); (2) the presence of potential deep geographical overlaps of two eastern clades (i.e. the pedunculate clade and the *P. obovalis* + *P. trachyphylla* + *P. virgata* clade); (3) early and substantial divergence of a south-eastern Qld lineage (*P. glasshousiensis* + *P. queenslandica*) within the pedunculate clade; (4) two distinct connections between the mainland and Tas. (i.e.

P. verrucosa and *P. virgata*, both in separate clades in the phylogeny); and (5) striking geographic disjunction between the sister taxa *P. myoporoides* subsp. *petraea* and *P. epilosa* (~900 km; Fig. 1c, j), and between the northern populations of *P. glasshousiensis* s.l. and *P. myoporoides* subsp. *acuta* (~850 km; Fig. 1d, f).

The seeds from members of sect. *Erionema* usually fall within a few metres of the parent and, unlike those from other sections of *Philotheca*, have no distinctive features to promote dispersal by animals (Armstrong 1991; Bayly 2001). It thus seems likely that long-distance dispersal has not been of major importance in the history of the group and that biogeographic patterns have mostly been shaped by vegetation shifts associated with past climatic and geological changes and a history of differentiation of allopatric taxa through vicariance.

The ages of divergences within sect. *Erionema* have not been estimated via molecular dating but some inferences can be made from previous work. We have not attempted molecular divergence dating here because fossils most suitable for calibration sit outside the family Rutaceae (Pfeil and Crisp 2008; Bayly et al. 2013) in taxa to which alignment of ITS and ETS markers is problematic and therefore likely to lead to spurious results. Nonetheless, the previous dating study of Bayly et al. (2013), based on chloroplast markers that display a more conservative rate of change (*rcbL* and *atpB*), did include one member of sect. *Erionema* (*P. buxifolia*) and representatives of the other three subgenera of *Philotheca* (including two samples of subg. *Philotheca*). The genus was not resolved as monophyletic in that study and relationships within the group were poorly supported, which, apart from other uncertainties regarding calibrations and mutation rate estimation, clouds understanding of divergence times. The branch connecting *P. buxifolia* to other taxa in the tree, i.e. a possible stem age for sect. *Erionema*, was 18–34 Ma (mean 28 Ma). The crown age for the section could be much younger than this, but this estimate at least allows the possibility that sect. *Erionema* dates to the Paleogene. A Paleogene age would be consistent with a vicariant separation of western and eastern Australia, potentially in the mid Miocene, as inferred for a range of other plant groups of the temperate mesic zone (Crisp and Cook 2007). A history on that timescale could also help to account for the presence of highly disjunct, potentially relictual lineages in south-eastern Australia, and the presence of multiple, deeply diverged but geographically overlapping clades.

The suggested relationship between *P. myoporoides* subsp. *petraea* and *P. epilosa* is particularly remarkable, given not just the distance between them, but also the presence of other members of the section in intervening areas. Despite the substantial distance, a relationship between the two taxa is morphologically plausible as they show a strong resemblance in leaf shape and size, as also noted by I. C. Clarke in annotations on one of the only two herbarium specimens of subsp. *petraea* (MEL 2030756A). However, if this resemblance is

taken as evidence of relationship, a high level of conservation in leaf shape is implied despite substantial isolation in distance and presumably time, and across differing climates.

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Data availability. The data supporting this study are publicly available. DNA sequences are available in GenBank (see <https://www.ncbi.nlm.nih.gov/genbank/>) with accession numbers as indicated in Table 1. The final DNA alignment, including INDEL characters, is available in TreeBase (see <http://treebase.org/>; study accession number 26368).

Conflicts of interest. Dr Murphy and Dr Bayly are both editors for *Australian Systematic Botany*. Despite this relationship, these authors did not at any stage have editor-level access to this manuscript while in peer review, as is the standard practice when handling manuscripts submitted by an editor to this journal. *Australian Systematic Botany* encourages editors to publish in the journal and has protocols that keep editors completely separate from the decision-making processes for their manuscripts. The authors have no further conflicts of interest to declare.

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