Phytophthora: an ancient, historic, biologically and structurally cohesive and evolutionarily successful generic concept in need of preservation

Clive Brasier1*, Bruno Scanu2, David Cooke3 and Thomas Jung4,5*

Abstract

The considerable economic and social impact of the oomycete genus Phytophthora is well known. In response to evidence that all downy mildews (DMs) reside phylogenetically within Phytophthora, rendering Phytophthora paraphyletic, a proposal has been made to split the genus into multiple new genera. We have reviewed the status of the genus and its relationship to the DMs. Despite a substantial increase in the number of described species and improvements in molecular phylogeny the Phytophthora clade structure has remained stable since first demonstrated in 2000. Currently some 200 species are distributed across twelve major clades in a relatively tight monophyletic cluster. In our assessment of 196 species for twenty morphological and behavioural criteria the clades show good biological cohesion. Saprotrophy, necrotrophy and hemi-biotrophy of woody and non-woody roots, stems and foliage occurs across the clades. Phylogenetically less related clades often show strong phenotypic and behavioural similarities and no one clade or group of clades shows the synapomorphies that might justify a unique generic status. We propose the clades arose from the migration and worldwide radiation ~ 140 Mya (million years ago) of an ancestral Gondwanan Phytophthora population, resulting in geographic isolation and clade divergence through drift on the diverging continents combined with adaptation to local hosts, climatic zones and habitats. The extraordinary flexibility of the genus may account for its global ‘success’. The 20 genera of the obligately biotrophic, angiosperm-foliage specialised DMs evolved from Phytophthora at least twice via convergent evolution, making the DMs as a group polyphyletic and Phytophthora paraphyletic in cladistic terms. The long phylogenetic branches of the DMs indicate this occurred rather rapidly, via paraphyletic evolutionary ‘jumps’. Such paraphyly is common in successful organisms. The proposal to divide Phytophthora appears more a device to address the issue of the convergent evolution of the DMs than the structure of Phytophthora per se. We consider it non-Darwinian, putting the emphasis on the emergent groups (the
Taxonomy's purpose is to foster clear scientific communication and the job of taxonomists is to refine it with that in mind. In doing so, Taxonomists must not only recommend improved communication going forward, but also weigh the costs of altering longstanding, effective communication (Booth 1978).

**INTRODUCTION**

The era of molecular phylogeny has provided strong evidence that the downy mildews (DMs) are as a group polyphyletic, having evolved at least twice from Phytophthora ancestors (Cooke et al. 2000; Runge et al. 2011; Jung et al. 2017a; Bourret et al. 2018; Scanu et al. 2021). This has led to a proposal to distribute the main phylogenetic clades of Phytophthora among several new genera (Runge et al. 2011), further indicated recently by Crous et al. (2021a). To assess the merits of this proposal we review here the environmental, economic and social impact, and the biological and phylogenetic characteristics of the genus, including its relationship to the DMs. We conclude that the case for retaining Phytophthora as a single genus is overwhelming.

**COMMENTARY**

Historical background

Phytophthora is arguably the world's most historic and economically significant genus in plant pathology. A comprehensive timeline of milestones for the genus is given in Table 1. In current classifications Phytophthora is usually assigned to the phylum Oomycota, which in turn are widely accepted as belonging to the heterokont algal-derived, but still somewhat debated, Straminipila within the kingdom Chromista (Dick 2001; Beakes et al. 2012). Together with other oomycetes, Phytophthoras are diploid with gametangial meiosis (Sansome 1961, 1965) and a genetic system akin to that of vascular plants (Brazier 1992; Goodwin 1997). They form indeterminate sporangiophores bearing alga-like sporangia that, in turn, release flagellate zoospores; and alga-like sexual oogonia and antheridia. Like most oomycetes, Phytophthoras exhibit a strong dependence on free water or high humidity for sporangial formation, zoospore spread and infection.

Within the oomycetes, Phytophthora is now assigned to the order Peronosporales, the vast majority of which are plant pathogens (Runge et al. 2011; Thines and Choi 2016; Jung et al. 2017a, 2018a). Amongst others this order includes the genera Halophytophthora and Calycophora, Phytophthora's sister genus Nothophytophthora, and 20 genera of DMs including Bremia, Peronospora, Plasmopara and Sclerospora (cf. Thines and Choi 2016; Jung et al. 2017a; McCarthy and Fitzpatrick 2017; Bourret et al. 2018; Scanu et al. 2021; Maia et al. 2022). Phytophthora and Nothophytophthora are mainly soil and water inhabiting, necrotrophic to hemibiotrophic pathogens forming zoosporic sporangia, whereas the DMs are aerial, obligate biotrophic pathogens with often conidia-like sporangia. The first DM genus, Peronospora, was erected by Corda (1837). The nomenclatural history of Phytophthora began with the potato blight epidemic in western Europe in the 1840s that led to the infamous Irish potato famine (Large 1940; Bourke 1991). The causal agent was initially named Botrytis infestans by Montagne (1845). It was then redesignated Peronospora trifurcata by Unger (1847), Peronospora infestans by Caspary in (1853) (published in Rabenhorst’s Herbarium vivum Mycologicum exsiccati no. 1879), and finally renamed Phytophthora infestans by de Bary (1876), with P. infestans as the ‘type species’ for the new genus Phytophthora (Table 1).

With the expansion of plant pathology as a discipline in the early 1900s the number of described Phytophthora species gradually increased. Rosenbaum (1917), Tucker (1931) and Leonian (1934) produced the first morphologically based keys to meet the growing need for accurate communication. Tucker (1931) accepted twenty species and was notable in emphasising the value of sporangial and gametangial morphology and temperature-growth relations as taxonomic criteria (Brasier 1991). Waterhouse (1963) developed a key based on assigning around 40 species to six morphological groups, introducing a sense of cohesion to a rather loosely structured mass of information (Gallegly 1983). Later, Waterhouse (1970) listed 60 Phytophthora species with a Latin description and/or a designated type, but 19 of these were later discarded in the Phytophthora monograph of Erwin and Ribeiro (1996), who
accepted 58 species; seven of which were later considered invalidly published or lost. The Waterhouse morphological system was developed further in the keys of Newhook et al. (1978) and Stamps et al. (1990). By the late 1980s, however, population-based, karyotype-based and molecular polymorphism-based systematic criteria were being advocated, heralding advancement towards a revised species concept and a more natural evolutionary phylogeny, including the likelihood that the *Phytophthora* genetic system was generating interspecific hybrids (Brasier 1991; Hansen 1991).

Table 1  Timeline of biological milestones in the genus *Phytophthora*

| Year | Milestones in the genus *Phytophthora* | References |
|------|--------------------------------------|------------|
| 1845 | Potato blight epidemic in Europe | Bourke (1991) |
| 1876 | *Phytophthora infestans* designated cause of potato blight | de Bary (1876) |
| 1892 | *Phytophthora* assigned to *Peronosporales* | Fischer (1892) |
| 1922 | ‘Heterothallism’ discovered (*P. faberi*, syn. *P. palmivora*) | Ashby (1922), Gadd (1924) |
| 1925 | First genus-wide physiological studies in *Phytophthora* | Leonian (1925) |
| 1931 | First major taxonomic treatments on *Phytophthora* | Tucker (1931), Leonian (1934) |
| 1935 | Thiamin requirement for growth demonstrated | Ronsdorf (1935), Leonian and Lilly (1938) |
| 1952 | Mode of evolution from lower to higher *Peronosporales* proposed | Gäumann and Wynd (1952) |
| 1960 | A1 and A2 compatibility types in ‘heterothallics’ are bisexual | Galindo and Gallegly (1960) |
| 1963 | Taxonomic key to ~40 known *Phytophthora* species | Waterhouse (1963) |
| 1963 | Oomycetes (*Achlya, Pythium, Phytophthora, Sclerospora*) shown to be diploid with gametangial meiosis | Sansome (1961, 1963, 1965, 1966) |
| 1964 | Exogenous sterols required for sexual reproduction | Elliot et al. (1964), Haskins et al. (1964), Hendrix (1964), Leal et al. (1964) |
| 1972 | ‘Battle for or against diploidy’ convention in Bari, Italy | Brasier (1972), Ko (1978) |
| 1972 | Chemical/hormonal induction of sexual differentiation, including selfing, in A1 x A2 interactions | |
| 1973 | *Phytophthora infestans* shown to be diploid. First report of chromosomal structural hybrids (reciprocal translocation heterozygotes) in A1 x A2 outcrossing *Phytophthoras* | Sansome and Brasier (1973) |
| 1980 | Mitotic segregation of the homozygous from the heterozygous mating type suppressed by reciprocal translocation heterozygosity | Sansome (1980) |
| 1980 | First molecular taxonomy based on protein and DNA polymorphisms | Kaosiri and Zentmyer (1980), Erselius and Shaw (1982), Hansen et al. (1986), Förster et al. (1988, 1990) |
| 1989 | Oomycetes assigned to *Staminigila* | Patterson (1989), Dick (2001) |
| 1990 | Designation of *Halophytophthora* gen. nov. | Ho and Jong (1990) |
| 1996 | *Phytophthora* reaches ~58 described species | Erwin and Ribeiro (1996) |
| 1998 | First interspecific hybrid described | Man in’t Veldt et al. (1998) |
| 1997 | Role of effector molecules in *Phytophthora* host specificity and pathogenesis | Kamoun et al. (1997) |
| 2000 | First molecular phylogeny of the oomycetes. Major *Phytophthora* clades identified. Downy mildews (*Peronospora, Bremia*) shown as evolved from *Phytophthora* | Cooke et al. (2000) |
| 2002 | World-wide surveys reveal many new *Phytophthoras* undetected in natural ecosystems | Jung et al. (2002, 2017b, 2018b, 2020, 2021; d), Brasier et al. (2010), Burgess et al. (2018), Dang et al. (2021) |
| 2004 | First multigene phylogeny of *Phytophthora* | Kroon et al. (2004) |
| 2007 | First multigene phylogeny of the oomycetes. Clade structure sustained | Göker et al. (2007) |
| 2009 | 400–600 *Phytophthora* species predicted | Brasier (2009) |
| 2014 | Divergence of *Phytophthora* Clades predicted at 19.8–39 m years ago | Matari and Blair (2014) |
| 2017 | Designation of *Nothophytophthora* gen. nov. *Phytophthora* origin pre the Gondwana-Laurasia separation (>180 Myr) proposed | Jung et al. (2017a) |
| 2018 | Multiple evolution of downy mildews from *Phytophthoras* demonstrated | Bourret et al. (2018) |
| 2021 | First phylogeny from genome-wide sequencing. Clade structure sustained | Van Poucke et al. (2021) |
| 2021 | *Phytophthora* reaches 200 described species | Scanu et al. (2021), Chen et al. (2022) |
Rapid increase in described Phytophthora species
From around 2000 the number of described species increased rapidly. This was partly due to the unravelling of morphospecies complexes by combinations of classical and molecular methods (e.g. Brasier et al. 2003; Hansen et al. 2019; Jung et al. 2011; Bertier et al. 2013; Ginetti et al. 2014; Safaiefarhani et al. 2015; Weir et al. 2015); and partly to the discovery of many new species and infraspecific lineages during dedicated surveys in forests and natural ecosystems, especially remote regions with low accessibility (e.g. Jung et al. 2003, 2011, 2017b, c, d; Rea et al. 2011; Reeser et al. 2011; Brasier et al. 2012; Scanu et al. 2015; Burgess et al. 2018; Dang et al. 2021).

Within a decade the number of formally described Phytophthoras had surpassed 100, and it was estimated that the number of extant Phytophthora species could be between 200 and 600 (Brasier 2009). Currently, the number of formally described and accepted taxa has reached 200 (Scanu et al. 2021; Chen et al. 2022) and many other new taxa have been designated informally (Brasier et al. 2003; Hüberli et al. 2013; Oh et al. 2013; Jung et al. 2017d, 2018b, 2020). Another 2–400 species may remain to be discovered in the world’s unsurveyed forests and natural ecosystems (Brasier 2009).

Figure 1 summarises the numerical chronology of described Phytophthora species, highlighting the exponential increase in species numbers over the past two decades.

Economic, environmental and social impact of the genus
Having been born out of a disastrous famine in Western Europe the genus Phytophthora was imbued with a degree of notoriety from its inception. Any modern perception of the genus needs to be much broader, in part because a definition of ‘importance’ in solely human terms is an artificial, not a biological, construct: *P. infestans* is no more biologically significant in its natural environment than are most other Phytophthoras in theirs. Nonetheless the genus contains a remarkable number of individually infamous pathogens, including (in addition to *P. infestans*) *P. capsici, P. cinnamomi, P. megakarya, P. nicotianae, P. palmivora, P. plurivora* and *P. ramorum*. Overall, the anthropogenically-related impacts of Phytophthoras are enormous (Tables 2, 3, Additional files 1, 2: Tables S1, S2). Most of these impacts are driven by introductions to environments with highly susceptible hosts, use of crop monocultures, host stress due to ‘off-site’ cultivation and climate change, or a combination of these (e.g. Brasier and Scott 1994; Erwin and Ribeiro 1996; Jung 2009; Jung et al. 2000, 2018a; Rizzo et al. 2002;
Table 2  Examples of the ecological, economic, social and scientific impacts of selected *Phytophthora* species

| *Phytophthora* species | Clade | First described | Environments | Main diseases caused and impacts                                                                                                                                                                                                 | Scopus indexed articles and their citations |
|-----------------------|-------|-----------------|--------------|-----------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------------|--------------------------------------------|
| *P. agathicida*        | 5     | 2015            | Forest, park | Dieback of Kauri, one of the world’s largest and longest-living conifer species, in New Zealand, spreading since 1974. Negative impact on both forest ecosystems and Maori society due to the ecological and cultural significance of Kauri trees | 29/171                                     |
| *P. austrocedri* (syn. *P. austrocedrae*) | 8     | 2005            | Forest, natural ecosystem | Dieback and mortality of native *Austrocedrus* forests in the southern Andes. Dieback and mortality of native Juniper, UK, initially associated with restoration planting of infested nursery stock | 26/135<sup>b</sup>                        |
| *P. cactorum*          | 1     | 1886            | Agriculture, forest, nursery | Root, collar, crown and fruit rots and stem cankers on over 200 species of trees, ornamentals, and fruit crops in 160 genera worldwide | 586/6481                                   |
| *P. capsici*           | 2     | 1922            | Horticulture | Phytophthora blight of *Capsicum* in the Americas and Southeast Asia, and a major limiting factor to vegetable production globally, especially cucurbits, tomatoes, and succulent beans, causing up to 100% losses in individual fields | 1559/18,150                               |
| *P. cinnamomi*         | 7     | 1922            | Forest, heathland, nursery, garden | Dieback of eucalypt forests and woodlands and mass destruction of World Heritage heath flora in Western Australia since 1950s. Heavy mortality of Fagaceae in forests of southeastern US since 1940s and southern Europe since ~ 1990s. Damage to ornamental nursery trade in Europe since 1970s. Listed as one of the 100 worst invasive alien species; pathogenic to ~ 5000 trees, woody ornamentals, and herbaceous plants worldwide | 1331/12,976                                |
| *P. cryptogea*         | 8     | 1919            | Horticulture, nursery, garden | Root and collar rot on a wide range of crops, fruit trees and ornamentals worldwide. Particularly important pathogen in greenhouses | 272/5261                                   |
| *P. fragariae*         | 7     | 1940            | Horticulture | Red core root disease of strawberry since 1920s, causing serious economic losses in strawberry plantations across humid regions of Europe and North America, with severely reduced yields and small poor-quality fruit. In Canada production losses to growers of Can$ 1500 per ha | 178/2091                                   |
### Table 2 (continued)

| Phytophthora species<sup>a</sup> | Clade | First described | Environments | Main diseases caused and impacts | Scopus indexed articles and their citations |
|---------------------------------|-------|-----------------|--------------|----------------------------------|---------------------------------------------|
| *P. infestans*                  | 1     | 1876            | Agriculture, horticulture | Late blight of potato and tomato, notorious for the Irish potato famine 1845–1849 resulting in mass starvation and migration. Currently still a serious threat to global food security worldwide, with US$ 6.7 million annually in yield losses and control costs | 4241/44,346 |
| *P. kernoviae*                  | 10    | 2005            | Forest, heathland, horticulture, park | Aerial bleeding cankers on European beech and leaf and shoot blights of Rhododendron, Magnolia spp., and wild bilberry in the UK and Ireland | 59/976 |
| *P. lateralis*                  | 8     | 1942            | Forest, nursery, park, shelterbelt | Root disease causing heavy mortality of Port Orford cedar (*Chamaecyparis lawsoniana*) in its native range in Oregon and California since 1950s. Serious impact on trade in this valuable commercially harvested timber. Recently spread to ornamental *C. lawsoniana* in western Europe. Social impacts through loss of business in nursery and forestry sectors | 62/1471 |
| *P. megakarya*                  | 4     | 1979            | Agroforestry | Main cause of Black pod disease of cocoa trees in central west Africa since the early 1900s, recently spread to Ghana. Loss of yield often > 30% for the economically important cocoa industry, worth ca US$ 70 billion annually | 106/2301 |
| *P. nicotianae* (syn. *P. parasitica*) | 1     | 1896            | Agriculture, horticulture, nursery, garden | Severe diseases of agricultural and horticultural crops worldwide, including foot rot and gummosis of citrus, black shank of tobacco and collar rot of tomato. Also on ornamentals. Broad host range, infects > 255 genera in 90 plant families | 1235/17,464<sup>b</sup> |
| *P. palmivora*                  | 4     | 1919            | Agroforestry, nursery, garden | Major impact on the production of tropical tree crops including black stripe disease of rubber in Southeast Asia since early 1900s. Also Black pod disease of cocoa in Southeast Asia and the Caribbean, with annual global losses to the cocoa industry of ca 450,000 t valued at > US$ 1 billion. Many ornamental hosts | 567/5892 |
| *P. plurivora*                  | 2     | 2009            | Forest, nursery, park, garden | Root and collar rot and aerial stem cankers on a wide range of woody hosts in Europe and North America; involved in the decline of oak and beech across Europe. Severe impact on the ornamental nursery industry | 66/644 |
These impacts can also be broadly divided into economic impacts, where Phytophthoras are causing losses or damage to cash crops in agricultural, horticultural or forestry systems; environmental impacts, where mainly introduced Phytophthoras are damaging native forest or herbaceous plant communities; and social impacts

| Phytophthora species* | Clade | First described | Environments | Main diseases caused and impacts | Scopus indexed articles and their citations |
|----------------------|-------|----------------|--------------|---------------------------------|------------------------------------------|
| *P. quercina*        | 12    | 1999           | Forest, park | Host-specific fine root pathogen. A main driver of the chronic decline of oak forests across Europe, interacting with climatic extremes | 53/896 |
| *P. ramorum*         | 8     | 2001           | Forest, nursery, garden | Over 200 plant hosts. High impact. Cause of Sudden oak death (native tanoak and other species) in the Western US since ~2000. Through loss of tanoak seed production, a significant impact on local wildlife and native American culture. Cause of Sudden larch death in the UK and Ireland since ~2010 with ~200 km² plantation larch affected and millions felled. Currently a threat to commercial timber production in the US (> US$ 30 billion) and the UK. Also damaging to the ornamental nursery trade in Europe and North America e.g. the rhododendron export trade in Canada (around US$ 5 million) | 627/8560 |
| *P. rubi* (syn. *P. fragariae* var. rubi) | 7     | 2007           | Horticulture | Extremely serious disease of raspberry plantations in Europe, North America, and elsewhere. EPPO A2 list, recommended for phytosanitary treatments | 57/475b |
| *P. sojae* (syn. *P. megasperma* var. sojae) | 7     | 1958           | Horticulture | Devastating root and stem rot of soybean in the US, with an annual cost worldwide of US$ 1–2 billion | 810/13,592b |
| *P. syringae*        | 8     | 1909           | Horticulture, nursery, garden | Root and collar rot, stem cankers, leaf and shoot blights and fruit rot on a medium-wide range of host plants including fruit trees and lilac | 71/2207 |
| *P. ×alni*           | 7     | 2004           | Riparian forest, nursery | Extensive mortality of riparian alder across Europe since 1990s, driven by planting of infested nursery stock. Impacts ecosystem functions and services and riverbank stability. EPPO alert list 1996 to 2001 | 72/772 |
| *P. × cambivora*     | 7     | 1927           | Forest, horticulture, nursery, garden | Root and collar infections (Ink disease) of sweet chestnut and beech in Europe. Root rot of various fruit trees in Europe and the US since 1900s. Significant impact on ornamental nurseries | 142/2259 |

* Associated references are shown in full in Additional file 1: Table S1
* Data include species synonyms

Shearer et al. 2004; Brasier and Webber 2010; Lamour 2013.

Table 2 (continued)
### Table 3  Examples of the ecological, economic and social impacts of disease syndromes or processes involving multiple *Phytophthora* species

| Syndrome or process and location | Environments | No. of *Phytophthora* taxa and hybrids involved<sup>a</sup> | Clades | Impacts |
|---------------------------------|--------------|----------------------------------------------------------|--------|---------|
| Cocoa black pod disease: West Africa, Caribbean, South America, Southeast Asia | Plantation | 5 | 2, 4, 5 | Cocoa pod lesions. Heavy crop losses (cf. *P. megakarya*, Table 2). Impact on small scale local farming communities and on global chocolate industry |
| Oak decline: across Europe | Forest, park | 26 | 1, 2, 3, 5, 6, 7, 8, 10, 12 | Root lesions and sometimes also collar lesions leading to forest declines driven in part by introduced pathogens and interaction with climate change. Impact on forestry and recreation |
| Beech decline: across Europe | Forest, park | 16 | 1, 2, 3, 5, 6, 7, 10, 12 | Root lesions, collar lesions, stem lesions leading to forest declines driven in part by introduced pathogens and interaction with climate change. Impact on forestry and recreation |
| Dieback of Mediterranean maquis vegetation: La Maddalena archipelago, Italy | Natural vegetation | 9 | 6, 7, 8 | Root lesions, collar lesions, stem lesions leading to mortality and decline of natural vegetation in a National Park. Impact on tourism, biodiversity and natural heritage |
| Restoration plantings in native Mediterranean heath vegetation and woodlands: Bay area, California | Planting, specialist nursery | 51 | 1, 2, 4, 6, 7, 8 | Strong evidence for spread to native plant habitats of at least five *Phytophthora* species causing root lesions, collar lesions, dieback and mortality of the vegetation. Impact on biodiversity and natural heritage |
| Dieback of eucalypt forests, Banksia woodlands and heath vegetation: across Western Australia | Forest, natural vegetation | 26 | 65 | Root and collar rot resulting in devastating dieback of whole ecosystems. Many of the *Phytophthora* species involved are considered native; however, the most aggressive species with the widest host ranges are introduced invasives (*P. cinnamomi*, *P. elongata* and *P. multivora*). Impacts on biodiversity, conservation, forestry and natural heritage |
| Woody plant nurseries and outplantings: across Europe | Nursery, outplantings | 65 | 1, 2, 3, 4, 6, 7, 8, 9, 12 | Most of these *Phytophthora* taxa are not native to Europe but are now established in the wider environment causing diseases of trees and shrubs in forests and natural ecosystems (e.g. *P. austrocedri*, *P. cactorum*, *P. cinnamomi*, *P. kernoviae*, *P. multivora*, *P. plurivora*, *P. ramorum*, *P. ×alni*, *P. ×cambivora*). Direct impact on nurseries; indirect impact on forestry, private garden owners, recreation and natural heritage |

---

<sup>a</sup> Lists of the individual taxa involved in each syndrome (including described species and currently informally designated species) and the full citations of the associated references are shown in Additional file 2: Table S2.
where significant damage is done to human communities with outcomes ranging from starvation, death and mass migration to loss of cultural heritage (Tables 2, 3, Additional files 1, 2: Tables S1, S2). In some cases, the impact factors are multiple. For example, the introduced *P. cinnamomi* causes damage to native forests and to important Mediterranean heath ecosystems and is also a serious problem in commercial nurseries and in horticultural and forest plantations (Brasier et al. 1993; Erwin and Ribeiro 1996; Shearer et al. 2004; Jung et al. 2016, 2018a, 2020). In Australia *P. cinnamomi* is considered a key threatening process to the Australian estate under the Environment Protection and Biodiversity Conservation Act 1999. The introduction of *P. ramorum* has had a considerable impact on the ornamental nursery trade in North America and Europe, caused heavy losses of native tanoaks (*Neolithocarpus densiflorus*) in the USA and commercial larch (*Larix kaempferi*) plantations in the UK, and collateral damage to many adjacent tree and shrub species and native ericaceous heaths (e.g. Rizzo et al. 2002; Brasier and Webber 2010; Jung et al. 2016, 2018a). The loss of oak and tanoak acorns has affected native American culture (e.g. Ortiz 2008) and food sources for wildlife. Many disease syndromes or processes involve multiple *Phytophthora* species (Tables 3, Additional file 2: Table S2). Since the 1990s, the number

Fig. 2 Number of important *Phytophthora* declines and diebacks of forests and natural ecosystems over time. Adapted from Jung et al. (2018a).

[1 = ink disease of *Castanea sativa* in Europe (observation of first typical symptoms in 1838), 2 = ink disease of *Castanea dentata* in the USA (observation of first typical symptoms in 1824), 3 = decline of *Fagus sylvatica* in the UK; 4 = littleleaf disease of pines in the USA, 5 = decline and mortality of *Chamaecyparis lawsoniana* in the Pacific Northwest, 6 = jarrah dieback in Western Australia (WA; observation of first typical symptoms in 1920s), 7 = ink disease of *C. crenata* and chestnut hybrids in Korea, 8 = eucalypt dieback in Victoria (observation of first typical symptoms in 1935), 9 = kauri dieback in New Zealand, 10 = Dieback of *Nothofagus* forests in Papua New Guinea, 11 = Mediterranean oak decline, 12 = Alnus mortality in Europe, 13 = temperate European oak decline, 14 = decline of *F. sylvatica* in mainland Europe, 15 = Sudden Oak Death in California and Oregon, 16 = littleleaf disease of *Pinus occidentalis* in the Dominican Republic, 17 = mortality of *Austrocedrus chilensis* in Argentina (observation of first typical symptoms in 1948), 18 = leaf and shoot blight of eucalypt plantations in New Zealand, 19 = oak decline in the Eastern USA, 20 = root and collar rot of eucalypt plantations in South Africa, 21 = needle cast and defoliation of *Pinus radiata* in Chile, 22 = dieback of *Eucalyptus gomphocephala* in WA, 23 = dieback of riparian *Eucalyptus rudis* in WA, 24 = Sudden Larch Death in the UK, 25 = dieback of *Araucaria excelsa* in Brazil, 26 = Ash decline in Denmark and Poland, 27 = dieback of *Nothofagus* spp. in the UK, 28 = mortality of *Juniperus communis* in the UK, 29 = red needle cast of *P. radiata* in New Zealand, 30 = leaf and twig blight of *Ilex aquifolium* in Corsica and Sardinia, 31 = dieback of Mediterranean maquis vegetation, 32 = Dieback of Fagaceae-Lauraceae monsoon forests in Northern Taiwan, 33 = Dieback of subtropical Fagaceae forests in Southern Taiwan, 34 = poplar dieback in Serbia, 35 = dieback of Valdivian rainforests in Chile, 36 = gummosis of *Acacia mearnsii* plantations in Brazil, 37 = collar rot of *P. radiata* plantations in New Zealand, 38 = dieback of laurisilva cloud forests in Northern Vietnam, 39 = black butt of *Acacia mangium* plantations in Vietnam, 40 = decline of *Cinnamomum cassia* plantations in Vietnam, 41 = cankers and dieback of Western hemlock and Douglas fir in the UK.]
of previously unknown *Phytophthora* declines of forests and natural ecosystems globally has increased exponentially, from 11 to currently 41 (Fig. 2).

The number of publications and citations for selected *Phytophthoras* is also indicated in Table 2. Again, this should not be automatically interpreted as indicating the 'relative importance' of a species. While they do to an extent reflect a species' scientific profile, such statistics can also be biased by the length of time a problem has been recognised; the economic value of particular cash crops—especially food crops; exacerbation of problems by subsequent disease management or biosecurity breaches; and the often more generous research grants available in developed countries. Furthermore, publications and citations related to environmental impacts, even major impacts such as loss of Kauri pines (*Aga thesis robusta*) to *P. agathidicida* in New Zealand, or loss of entire species-rich heath vegetation in southwest Australia to *P. cinnamomi* (Table 2, Additional file 3: Table S3), tend to be substantially fewer, usually as a consequence of limited research funding.

The monophyly of *Phytophthora* and the sustained structural stability of its phylogenetic clades

The first molecular phylogenetic analysis of *Phytophthora* and other oomycetes was published by Cooke et al. (2000). It was based on ITS profiles and included 50 described *Phytophthora* species. Unlike other oomycete genera such as *Pythium* or *Halophytophthora*, *Phytophthora* was revealed as a tight monophyletic cluster of eight major clades (Clades 1–8), plus two putatively more distantly related clades (Clades 9 and 10). Unsurprisingly, the clades were also shown to transcend previous morphological groupings. Strikingly, *Peronospora sparsa* clustered within *Phytophthora* Clade 4, indicating a relatively recent evolution of *Peronospora* and other DMs from *Phytophthora*. Cooke et al. (2000) suggested *Peronospora* and *Bremia* were obligate, conidial *Phytophthoras*, in support of an earlier proposal by Gäumann (1952).

Since then, at least 12 other molecular phylogenetic studies of *Phytophthora* have been undertaken, ranging from increasingly complex multigene analyses (Martin and Tooley 2003; Kroon et al. 2004, 2012; Blair et al. 2008; Robideau et al. 2011; Martin et al. 2014; Rahman et al. 2015; Jung et al. 2017c; Yang et al. 2017; Bourret et al. 2018; Scanu et al. 2021; Chen et al. 2022) to a genome-wide sequence-based phylogeny (Van Pouncke et al. 2021). These studies have been carried out against the background of the rapidly increasing number of described *Phytophthora* species outlined above.

Despite these fresh analyses and the addition of many new species our perception of the infrageneric structure of *Phytophthora* has changed little since Cooke et al. (2000). The overall clade structure has remained stable and generally accepted. The number of major clades, i.e. those with four or more species (therefore, excluding monospecific Clades 11, 13 and 14, currently represented by *P. lili*; the undescribed *P.* taxon mugwort; and *P. cyperi*, which is probably a DM; Ho et al. 2004; Bourret et al. 2018) has increased from ten to eleven (Jung et al. 2017c; Chen et al. 2022). The phylogenetic positions of some species have been clarified and multiple new subclades have been added.

Above all, the major *Phytophthora* clades are still confirmed to be a relatively tight, bush-like, fundamentally monophyletic, evolutionary cluster. Indeed Clades 9 and 10, considered by Cooke et al. (2000) to be more distant, are now more closely aligned with the other major clades (Jung et al. 2017c; Yang et al. 2017; Scanu et al. 2021; Van Pouncke et al. 2021; Chen et al. 2022). This is in contrast to the oomycete genus *Pythium* which, beginning with Cooke et al. (2000), has been shown to be evolutionarily divergent and polyphyletic, and in consequence was split into several monophyletic genera (de Cock et al. 2015; Uzuhashi et al. 2010).

Moreover, lineages encompassing *Peronospora* and the other 19 DM genera have now been shown to have evolved from *Phytophthora* at least twice (Bourret et al. 2018; Scanu et al. 2021; Fig. 3), confirming that *Phytophthora* is 'paraphyletic' in relation to its DM descendants (Cooke et al. 2000). Downy mildews with pyriform haustoria (DMPHs; e.g. *Bremia, Plasmopara*) and the obligate biotrophic *Phytophthora cyperi* form a monophyletic cluster in sister position to *Phytophthora* Clade 1 (Fig. 3). In contrast, those with coloured conidia (DMCCs; *Peronospora* and *Pseudoperonospora*), the graminicolous DMs (GDMs; e.g. *Sclerotinia*) and the brassicicolous DMs (BDMs; e.g. *Hyaloperonospora*) form a monophyletic cluster which diverged from a common ancestor with *Phytophthora* Clades 1–5, 12, and the DMPHs (Bourret et al. 2018; Scanu et al. 2021; Fig. 3). The DMs as a group therefore appear to be fundamentally polyphyletic.

*Phytophthora* is a biologically sound and cohesive genus

The continued acceptance by the scientific community of *Phytophthora* as an assemblage of clades has probably also reflected a perception that this structure exhibits strong biological cohesion. Thus, the 11 major phylogenetic clades share a wide range of characters, both morphological and behavioural, that collectively characterise the genus (Figs. 4, 5; Table 4, Additional files 3, 4, 5: Tables S3, S4, S5). Also these characters often show as much variation between species within a clade as they do between clades (Table 4, Additional files 3, 4, 5: Tables S3, S4, S5).
Fig. 3  Phylogenetic tree of the *Phytophthora* clades and representative downy mildews. Redrawn from Scanu et al. (2021). A fifty percent majority rule consensus phylogram derived from maximum likelihood analysis of a concatenated four-locus (ITS, Btub, cox1, nadh1) dataset of representative species from phylogenetic Clades 1–12 of *Phytophthora* and the four downy mildew groups DMPH, DMCC, GDM, and BDM. Maximum likelihood bootstrap values and Bayesian posterior probabilities are indicated but not shown below 60% and 0.80, respectively. *Nothophytophthora amphigynosa* was used as outgroup taxon (not shown). Scale bar = 0.01 expected changes per site per branch.
For example, caducous (deciduous) sporangia are found in nine major clades and persistent sporangia in all 11 clades (Fig. 4; Additional file 3: Table S3). Chlamydo- spores are produced by 59 species in ten clades; and 137 species across all eleven clades lack the ability to produce them (Fig. 4; Additional file 3: Table S3). Eight of the clades contain both self-fertile (homothallic) species and species with an A1/A2 outcrossing (heterothallic) breeding system. Both amphigynous and paragynous antheridia are found in eight clades (Fig. 4; Additional file 3: Table S3). Sterile species occur in five clades.

Conspicuous morphological similarities are shared between species in phylogenetically divergent clades (Table 4). For example, both *P. infestans* (Clade 1) and *P. ramorum* (Clade 8) have caducous, semi-papillate sporangia, and both are A1/A2 outcrossing with amphigynous antheridia. Both *P. pseudosyringae* (Clade 3) and *P. foliiorum* (Clade 8) produce semipapillate, partly caducous sporangia, and both are self-fertile with mostly paragynous antheridia. *Phytophthora multivesiculata* (Clade 2) and *P. europaea* (Clade 7) are both self-fertile with paragynous antheridia and both form non-papillate persistent sporangia; while *P. clandestina* (Clade 1) and *P. kernoviae* (Clade 10) are both self-fertile with amphigynous antheridia and both produce papillate caducous sporangia (Table 4).

Furthermore, phylogenetically divergent clades often share strong similarities in ‘lifestyle’. An aerial dispersal lifestyle occurs across eight and a soilborne lifestyle across all 11 major clades. Apparently very flexibly-adapted species exhibiting both an aerial and a soilborne lifestyle are found in seven clades (Fig. 5; Additional file 4: Table S4). The ability to infect and seriously damage roots, bark (phloem) and even xylem (Brown and Brasier 2007; Parke et al. 2007) tissues of woody hosts as well as herbaceous tissues is something of a *Phytophthora* speciality among the oomycetes, and largely distinguishes the genus from the obligately biotrophic DMs. It is found in 71.9% of the species and across all the clades (Fig. 5; Additional file 4: Table S4). Species with wide, medium and narrow host ranges and species exhibiting host specificity are found in all or nearly all clades (Fig. 5; Additional file 4: Table S4).

In addition, currently at least 85 *Phytophthora* species (43%), representing all 11 major clades, have been shown to disperse in an aquatic environment (Fig. 5; Additional file 4: Table S4) and to live saprotrophically, free from the host. This property also distinguishes *Phytophthora* from the DMs. Forty-one species in five clades, including 25 of the 27 sterile species, have a primarily aquatic lifestyle as
litter decomposers and opportunistic pathogens (Fig. 5; Additional file 4: Table S4).

A wide adaptation to climatic conditions is another feature shared across the clades: the majority contain species with either low, medium or high cardinal temperatures for growth (Fig. 4; Additional file 5: Table S5). Indeed, phylogenetically divergent Clades 1 and 8 are remarkably similar both in terms of the proportion of species adapted to low, medium or high optimum temperatures and in their maximum temperature tolerances (Additional file 5: Table S5). Furthermore, within each of the 21 different ‘lifestyle and behavioural categories’ listed in Additional files 4 and 5: Tables S4 and S5 the number of clades with taxa that exhibit the attribute is consistently high: average 9.1 across the eleven major clades; range 5–11.

Cooke et al. (2000) also reviewed the morphological and behavioural properties of their 50 Phytophthora taxa and proposed that Clades 1–5 comprised predominantly aerially dispersed species with papillate caducous sporangia and Clades 6–8 predominantly soil dispersed species with persistent non-papillate sporangia, consistent with an earlier proposal for two evolutionary trends in the genus (Brasier 1983). In the present analysis and that of Yang et al. (2017) this last proposal is no longer fully supported. Of the 75 species in Clades 1–5, for example, 32% are papillate caducous and aerial, another 30.7% are papillate persistent and soil inhabiting and the remaining 36.3% represent a mixture of attributes (Additional file 6: Table S6). For Clades 6–8 however the proposal does have support. Of 89 species, 80.9% are non-papillate persistent and soil inhabiting, compared to 11.2% semi-papillate persistent and soil inhabiting and 4.5% semi-papillate and partly caducous (Additional file 6: Table S6). Clades 1–5, therefore, appear more flexible in terms of their present day ‘lifestyle’ variability than Clades 6–8.

Across the clades unusual developmental features are exhibited by a small number of species (Table 5, Additional file 7: Table S7). Collectively, these are another indication of the behavioural adaptability of the genus. Sporangiophore constrictions are found in P. pinifolia (Clade 6) and P. constricta (Clade 9), presumably to facilitate aerial dispersal in otherwise non-papillate, soil and waterborne species. These appear to be an example of convergent evolution (Rea et al. 2011), as do the ultra-long sporangial pedicels produced by P. capsici (Clade 2) and P. hibernalis (Clade 8) (Kunimoto
et al. 1976; Erwin & Ribeiro 1996). These pedicels may promote sporangial clustering (cf. Granke et al. 2009) and adherence to surfaces. Also unusual are the stromata formed by P. cinnamomi (Clade 7) and by P. ramorum and P. lateralis (Clade 8) (Table 5; Moralejo et al. 2006; Brasier et al. 2010; Jung et al. 2013), which may be adaptations for nutrient storage and eruption through tough leaf or periderm surfaces.

Evolution and adaptability of the *Phytophthora* clades

Considering the many ecological niches and environments it has occupied, phenotypically the genus *Phytophthora* appears to have changed remarkably little. This raises the question of the evolutionary processes that have resulted in its clades being phylogenetically divergent, yet still exhibiting strong biological and behavioural conformity coupled with high adaptability. We suggest that the clades developed as a result of the migration and worldwide radiation of an ancestral Gondwanan or pre-Gondwanan *Phytophthora* population on the emerging continents, beginning around 140 Mya and that the resulting geographic isolation led to a degree of clade divergence through genetic drift, and also local adaptation to the different hosts and parts of hosts, habitats and climatic zones on the diverging continents.

We suggest the sustained biological similarity across the clades (Figs. 4, 5; Table 4; Additional files 3, 4, 5: Tables S3, S4, S5) is due to the *Phytophthora* lifestyle and genome being highly versatile in terms of (1) switching between different spore morphologies or dispersal modes (Fig. 4; Additional file 3: Table S3); (2) a range of trophic options from saprotrophy on degraded vegetation through necrotrophy and transient biotrophy on diverse tissues of pteridophytes, gymnosperms and angiosperms (Fig. 5; Additional file 4: Table S4); (3) different breeding strategies (Fig. 4; Additional file 3: Table S3), including an inbreeding system (‘homothallism’) with probably no barrier to outbreeding (cf. Whisson et al. 1994) and an outcrossing system (A1/A2 compatibility or ‘heterothallism’) that also enables selfing (Sansome 1980; Brasier 1992; Judelson 2009); (4) an ability to rapidly modify host–pathogen recognition processes via effector molecules (Kamoun et al. 1997; Kamoun 2006) and other pathogenicity factors associated with the more rapidly evolving component of the ‘two speed’ genome (Raffaele and Kamoun 2012; Zhang et al. 2019; Dale et al. 2019); (5) rapid evolution of asexual clones via mitotic recombination and transposon induced mutagenesis (e.g. Kasuga et al. 2016; Dale et al 2019); and (6) rapid evolution via interspecific hybridisation (Brasier et al. 1999, 2004; Man in’t Veld et al. 2012; Bertier et al. 2013; Burgess 2015; Aguayo et al. 2016; Jung et al. 2017d; Van Poucke et al. 2021). These attributes may have facilitated adaptation to new hosts and new biogeographic environments without...
significant biological change; often resulting in convergent evolution between otherwise geographically isolated clades (Figs. 4, 5; Table 4).

In summary, we consider that *Phytophthora* is a highly flexible and highly successful biological ‘model’ that has survived well over aeons of time, leading to molecularly detectable divergence between its clades without a marked disjunction in their general morphology or behaviour. This great adaptability is probably a major factor in their being high risk pathogens when introduced into new environments.

**Emergence of the Downy Mildews**
A major development among the *Phytophthora* clades has been the emergence of the DMs. On one occasion this was from a common ancestor with Clades 1–5 and Clade 12, on the other from a common ancestor with Clade 1 (Bourret et al. 2018; Fig. 3). This has resulted in the emergence of organisms with a broadly similar range of properties, apparently via convergent evolution. Probably the most definitive characteristics of the DMs are obligate biotrophy of angiosperms, usually accompanied by intracellular haustoria; and sporangiophores of determinate growth, facilitating synchronised dispersal. Major host shifts including adaptive radiation on particular plant families such as the *Poaceae* and *Brassicaceae,* combined with high levels of host specificity on largely herbaceous plant parts, are believed to have played a major role in their emergence (Göker et al. 2007; Thines and Choi 2016; Bourret et al. 2018). Also characteristic is aerial dissemination; and, in 10 of the 20 genera, directly germinating conidiosporangia (Gäumann 1964; Hall 1996; Göker et al. 2007; Thines and Choi 2016; Fletcher et al. 2019).

Given the adaptiveness and global spread of Phytophthoras it is perhaps surprising that this process has succeeded only twice. However, these developments involved abandoning definitive *Phytophthora* properties. This includes: necrotrophic ability (Thines and Choi 2016; Fletcher et al. 2018), important in *Phytophthora* parasitism of woody tissues and resulting in DMs being more benign pathogens or even endophytes; nutrient transporters linked to saprotrophic ability, enabling Phytophthoras nutrient gain in competition with other microorganisms; loss of the ability to utilise inorganic nitrogen and sulphur (Yin et al. 2017); loss of indefinite sporangiophore development, an adaptation best suited to continuously wet or aquatic conditions; and for many DMs, loss of the mechanism of zoosporogenesis (Fletcher et al. 2018) and therefore zoospore mediated infection. Indeed, detailed comparisons of *Phytophthora* and DM genomes support synteny and a common origin but also

### Table 5 Unusual morphological or developmental features among *Phytophthora* species

| Species         | Clade | Features*                                                                 | Possible adaptations                                                                 |
|-----------------|-------|---------------------------------------------------------------------------|-------------------------------------------------------------------------------------|
| *P. infestans*  | 1     | Sporangiophore apophyses                                                  | Mechanism of indeterminate sporangiophore growth                                    |
| *P. capsici*    | 2     | Long pedicels 30–100 µm Umbellate sympodia                                | Splash dispersal, sporangial clustering and adherence to host surfaces               |
| *P. litchii*    | 4     | Downy white mycelium                                                       | Resistance to desiccation on suberised fruit pericarp surface                       |
| *P. heterospora*| 4     | Determine sporangiophores and synchronous sporangial formation            | Rapid synchronized sporulation on exposed fruit pericarp surface                     |
| *P. pinifolia*  | 6     | Narrowing of sporangiophores near sporangial bases                         | Adaptation to both moist and drier habitats or seasonal or diurnal climate          |
| *P. cinnamomi*  | 7     | Tough mycelium Stromata                                                    | Facultative caducity, enabling aerial infection and dispersal in a Clade of soil- and water-borne species |
| *P. lateralis*  | 8     | Stromata and sporangiomata                                                 | Competitive growth through soil and litter Nutrient storage for seasonal hibernation and subsequent sporulation Long term survival |
| *P. hibernalis* | 8     | Long pedicels 20–80 µm                                                     | Splash dispersal, sporangial clustering and adherence to host surfaces               |
| *P. ramorum*    | 8     | Stromata and sporangiomata                                                 | Pressure eruption through tough leaf cuticle or fruit periderm followed by sporulation |
| *P. constricta* | 9     | Sporangiophore constrictions near the sporangial bases                     | Facultative caducity and aerial dispersal in a Clade of soil- and waterborne species |
| *P. insolita*   | 9     | Production of oospores without antheridia (presumed gametangial apomixis) | Inbreeding mechanism. Survival in periodically dry waterways without cost of less adapted recombinant offspring |

*Associated references are shown in Additional file 7: Table S7*
demonstrate that DMs have lost conserved domains encoding some of these properties (Fletcher et al. 2018, 2021). The narrower host and nutrient specificity, reduction in the effector repertoire and consistently reduced pathogenicity gene complements of the DMs (Fletcher et al. 2018) could render DM species more prone to extinction in a changing environment (Thines and Choi 2016). Equally, by completing their life cycles in the more sheltered and homogeneous milieu of living plant tissues DMs are probably less exposed to competition from other microorganisms; and due to their host specificity at less ‘risk’ of interspecific hybridisation, especially compared with the ecologically more flexible Phytophthoras (Brasier 2001; Van Poucke et al. 2021).

The much longer average branch lengths in the DMs, largely distinguish them from the more tightly clustered ‘bush-like’ Phytophthora clades, and probably reflect a relatively rapid evolution towards enhanced host specialisation and biotrophy (Bourret et al. 2018; and Fig. 3): an evolutionary jump, perhaps driven by strong directional selection associated with ensuing host–pathogen ‘arms races’. In consequence, many early stages in the evolution of the DMs are probably lost to extinction. Indeed, despite the thousand or so extant Phytophthora and DM taxa, examples of prominent Phytophthora-like characters among the DMs and vice versa are patchy and often somewhat equivocal. Thines (2009) discusses what may be ‘intermediate taxa’ or ‘bridging taxa’. But such terms are, however, subjective and should probably be treated with caution because of: (1) the unknown progenitor taxa; (2) intervening extinctions and reticulations; (3) the possibility of convergent evolution; and (4) the lack of information on the genetic control of many characters, such as haustorial form or sporangiophore development.

Amongst Phytophthoras, the recently described P. podocarpi (previously P. taxon totara) on Podocarpus shoots in New Zealand shares a common ancestor with the DMCCs and their relatives (Bourret et al. 2018) but otherwise has Phytophthora characteristics (Dobie et al. 2022). Phytophthora litchii (syn. Peronosphythora litchii) in Clade 4 resembles DMs in producing determinate sympodial sporangiophores, a ‘downy white mycelium’ on lychee (Litchi chinensis) fruits, and smaller gene families (Chen 1961; Ho et al. 1984; Ye et al. 2016; Sun et al. 2017). Otherwise, P. litchii causes necrotic lesions, is able to grow on artificial media, and displays a typical Phytophthora-type zoospore release and gametangial morphology. The aerial and soil inhabiting P. heterospora (also Clade 4), which causes bark and root necroses on various woody host plants, produces both zoospore-releasing sporangia and directly germinating pseudoconidia (Scanu et al. 2021).

The possession of such unusual characters by P. litchii and P. heterospora does not, however, confirm them as proto-DMs. The ‘DM-like’ characteristics of P. litchii (Tables 5, Additional file 7: Table S7) could be evidence of a common ancestor shared with the DMs, but they could also reflect convergent evolution (Ye et al. 2016) or ancestral hybridisation. The determinate sporangiophores and the ‘downy’ mycelium could be adaptations to reduce or avoid desiccation on the exposed surface of the leathery lychee pericarp. The downy mycelium might be more refractive due to a protective cell surface hydrophobin or mucin (cf. Meijer et al. 2006). The pseudoconidia of P. heterospora, at face value a DM-like feature, may be an adaptation allowing extra reproductive versatility in alternating moister and drier diurnal or seasonal conditions (Scanu et al. 2021). As already discussed, there are comparable unique or unusual developmental features in other Phytophthora species (Table 5, Additional file 7: Table S7).

**Phytophthora-like characters among the DMs**

Ten of the 20 DM genera, including *Pseudoperonospora, Plasmodpara* and *Sclerophthora* produce sporangia which release zoospores (Bourret et al. 2018). *Peronospora cyperi* (syn. *Kawakamia cyperi*), was later renamed *Phytophthora cyperi*, probably because of the reportedly *Phytophthora*-like caducous sporangia and paragynous antheridia (Erwin and Ribeiro 1996). Whether *P. cyperi* produces zoospores is uncertain, but it should probably be accepted under the name *K. cyperi* based on it being a biotroph (Thines et al. 2015) and its phylogenetic status (Bourret et al. 2018; and Fig. 3).

Three monotypic graminicolous DM genera, *Graminivora, Poakatesthia* and *Viennotia* show features not found in other DMs. In particular, the indeterminate sporangiofores and *P. infestans*-like sporangial apophyses in *V. oplismeni*, and the occurrence of intracellular hyphal growth in *Poakatesthia pennisetis*, which casts doubt on whether it is an obligate biotroph (Thines 2009). These features were suggested by Beakes and Thines (2017) to be evolutionary hangovers from *Phytophthora*, but some could also reflect convergent evolution.

Perhaps the best example of a DM with Phytophthora-like characters is the graminicolous genus *Sclerophthora*. This is widely presumed to be biotrophic (Kenneth 1981; Erwin and Ribeiro 1996; Thines and Choi 2016) and in molecular phylogenies exhibits the characteristic long branch length, or evolutionary jump, of the DMs (Thines et al. 2008; Bourret et al. 2018). *Sclerophthora macrospora* has Phytophthora-like sporangiophore and sporangial morphology, and a wide but highly specialised graminicolous, and therefore non-Phytophthora-like, host range (Kenneth 1981; Erwin and Ribeiro 1996; Telle and Thines...
Tokura (1975) reported culturing *S. macrospora* on artificial media but was unable to obtain zoospores or to infect rice seedlings with these cultures. Tokura’s observations have yet to be repeated. Any attempt to do so will hopefully include diagnosis of any resulting axenic growth with molecular markers.

Among a panoply of around 800 species across 20 DM genera it is likely that some ancestral *Phytophthora*-like characteristics will have survived as long as they did not confer a marked selective disadvantage. On the evidence of Thines (2009), retention of *Phytophthora*-like characters in the DMs is associated with particular host groups. Specifically, in *Sclerophthora* and *Viennotia* in respect of the *Poaceae*, and in ‘*P. cyperi*’ the *Cyperaceae*. Perhaps these associations involved host jumps, and possibly horizontal gene transfer (cf. Brasier 1995; Bourret et al. 2018; Fletcher et al. 2021), resulting in such closed host-pathogen systems that drivers towards further adaptation were less intense.

**Proposals to split *Phytophthora* into separate genera are biologically and phylogenetically inappropriate**

**Evolutionary process versus taxonomic cladism**

None of the authors of the many molecular phylogenies of *Phytophthora* published since Cooke et al. (2000) have suggested there is a case either for a merging of the clades, or for their nomenclatural designation as subgenera or sections. This apparent acceptance of the clade structure has probably also reflected a perception of their biological cohesion and of the enormous significance of the genus for scientific communication and global biosecurity (discussed later).

Nonetheless, to resolve the paraphyly of *Phytophthora*, reflected in the evidence that the DMs have evolved from the genus (Cooke et al. 2000; Jung et al. 2017a; Bourret et al. 2018; Scanu et al. 2021), and applying the terminology of cladism (not to be confused with cladistics), Runge et al. (2011) proposed either (1) placing all DMs and *Phytophthora* species in a single genus under the oldest generic name *Peronospora*, which would require renaming all *Phytophthora* species and those in 19 DM genera, resulting in a highly heterogeneous group; or (2) the description of at least six new genera within *Phytophthora* in order to conserve the DM genera. We consider the first of these suggestions intrinsically flawed as it would assimilate the ancient ancestral genus *Phytophthora* with its broad suite of morphological and behavioural characters into its highly specialised descendant, *Peronospora*, an evolutionary absurdity. Runge et al. (2011) suggested that their second alternative, splitting *Phytophthora* into around six new genera, would be most appropriate, but only on the highly questionable grounds that it would require fewer name changes. In terms of evolutionary process, their third option, reclassifying all DMs under the parental group *Phytophthora* would probably be the most logical. However, this would be in conflict with the International Code of Nomenclature for algae, fungi and plants (ICNafp) (Turland et al. 2018), which gives the first described genus *Peronospora* nomenclatural priority (though this could be overcome by conservation of *Phytophthora* over *Peronospora*); and, most importantly, it would ignore the biological realities. Voglmayr (2008) considered that none of the above alternatives would receive broad acceptance, “representing a dilemma for classification”.

Crous et al. (2021a) have also indicated that “it can be expected that *Phytophthora* will resolve into several genera in future studies”. Whether or not the paraphyly of *Phytophthora* is considered a genuine problem, that it was necessary for Runge et al. (2011) to propose two different ‘solutions’ is of concern. In our view this highlights the artificiality and subjectivity of the taxonomic process: such an approach takes little account of the often considerable biological and phylogenetic distances between *Phytophthora* and the DMs. This raises the question whether paraphyly, the emergence of one distinct life form from another without the progenitor becoming extinct, needs to be resolved by a different taxonomy at all, but should simply be accepted as a common feature of evolution in many ancient and successful genera. Around 20% of animal and 20–50% of plant species are paraphyletic in these terms (Crisp and Chandler 1996; Ross 2014). This makes it a common trait of evolution to be accepted at all taxonomic levels. Indeed, the somewhat negative use of the term ‘paraphyly’ in cladistics has been characterised as “Disparaging phylogenetic jargon for a cladogram’s representation of a progenitor in a macro-evolutionary series” (Zander 2013).

Further, in terms of the definitions of Ashlock (1971) and Aubert (2015), based on the original definition of a phylon as the totality of organisms ‘related by blood and descended from a common typical ancestor’ (Haeckel 1877), *Phytophthora* is clearly monophyletic because all known *Phytophthora* species share the same common ancestor; yet *Phytophthora* is also paraphyletic because it does not contain all descendants of the common ancestor, i.e. it does not include the DMs. Collectively, however, *Phytophthora* and the DMs are holophyletic, since they contain all the descendants of their shared common ancestor. In contrast to this essentially Darwinian definition of monophyly, cladism does not discriminate between monophyly and holophyly, but focusses the definition of monophyly on the descendants, not on the ancestor as in the original Haeckelian sense (Hennig 1966; Ashlock 1971; Hörandl 2006, 2007; Aubert 2015). As a consequence, in cladism the terms monophyly and
paraphyly are sometimes applied to phylogenetic trees as if they describe fixed or immutable entities. In reality they are useful generalisations that attempt to define a complex or continuum but are often unreliable due to past reticulations, significant gaps due to extinctions and, in the case of Phytophthora, the numerous undiscovered taxa (Brasier 2009). This renders the question of applying taxonomic weight, or names, to phylogenetic dichotomies contentious.

We consider that the key question for determining an appropriate taxonomy is not the existence of a node, but the evolutionary processes that gave rise to the divergence of the lineages and the extent of the biological changes involved (cf Bray 2009). This approach takes into account the main geographic, genetic or biological drivers of the dichotomy. Often these can only be retrospectively inferred, rather than critically established.

In terms of many analysts (e.g. Ashlock 1971; Crisp and Chandler 1996; Brummit 1998; Mayr and Bock 2002; Brummit and Sosef 2003; Hörandl 2006, 2007; Zander 2013; Aubert 2015), this is an evolutionary approach as opposed to a purely cladistic approach to establishing a meaningful and practical, if still approximate, taxonomy; and one in which classification is not allowed to trump evolution. Zander (2013) has pithily summarised this viewpoint as follows: 

**Phylogenetics imposes a classification on the results of cladistic analysis without a process-based explanation of those results. The sister-group structure is taken to be a classification itself. Evolution is not clustering, classification is. Evolution is not nesting, classification is. Phylogenetics leaps from the clustering and nesting of cladistic analysis straight to classification without explanation of the analysis in terms of serial transformations of one taxon into another, which is the nut of (Darwinian) macroevolutionary theory.**

On the above basis and taking into account our suggestions regarding the post-Gondwanan expansion of the Phytophthora clades, we consider Phytophthora to be a fundamentally monophyletic cluster that has at least twice given rise to the evolution of descendents with a distinct set of biological traits linked to obligate biotrophy, the DMs, via paraphyletic ‘jumps’. Since such jumps are common in nature, we see no current biological justification or systematic need for subjecting Phytophthora, a successful, ancient and biologically coherent mother genus, to segregation into separate genera. Further, proposals to do so appear mainly to be a device to address the convergent evolution of the DMs rather than a problem related to the structure of Phytophthora per se. This, despite the fact that it has also been acknowledged by Crous et al. (2021b) that, as a unit, a ‘genus’ should be defined not only by phylogeny but also by common morphological, ecological and chemical properties (= synapomorphies).

Further, because of their substantial differences in lifestyle, we consider it appropriate to accept a much broader generic concept for Phytophthora than for the DMs. In the more morphologically limited and behaviourally specialized DMs, genera have tended to be discriminated by conidiosporangial pigmentation and conidiophore and haustorial morphology, host specificity and, more recently, phylogenetic separation (e.g. Göker et al. 2003, 2007; Voglmayer 2008; Thines et al. 2015). For example, the BDM genus *Hyaloperonospora* is distinguished from the DMCC genus *Peronospora* largely by globose haustoria, non-pigmentation of the conidial walls and brassicaceous versus broad host specialisation. The other BDM genus, *Perofacia*, is distinguished from *Hyaloperonospora* largely by uniformly ellipsoidal conidia and the pseudodichotomous and appressed conidiophores (Constantinescu and Fatehi 2002). These generic differences are even more limited than the differences between some phylogenetically very closely related *Phytophthora* species. Thus *P. ramorum* and *P. hibernalis* in *Phytophthora* Clade 8c differ in sporangial pedicel morphology, the presence of chlamydospores, breeding system, antheridial type, oospore size, optimum temperature for growth and their main host families.

Overall, we consider that the biological and evolutionary case for retaining *Phytophthora* as a single genus is overwhelming. We have already shown that the 11 major *Phytophthora* clades share a characteristic diversity and plasticity across an extensive suite of morphological features, breeding systems and lifestyles. While there are trends, none of the clades are distinguished by a unique special character (synapomorphy) or combination of characters (cf. Bennett et al. 2017). In addition to monophyly, we consider the latter should be an indispensable requirement for recognition of a separate genus. Consequently, we propose for *Phytophthora*, rather than a monophyly-centred cladistic concept, a more ‘Darwinian’ generic concept based on similarity (synapomorphies) and common descent (monophyly in the original Haeckelian sense; Mayr and Bock 2002; Hörandl 2006). This allows similarities within the older parental group (*Phytophthora*) which exists in parallel to the descendant group (DMs) while excluding similarities resulting solely from convergent evolution.

**Scientific communication and biosecurity importance of Phytophthora as a cohesive genus**

Through being both well biologically defined and widely accepted the generic name *Phytophthora* is currently an engine of understanding and communication for a large body of scientists operating across disciplines ranging
from mycology and plant pathology to biosecurity and social history. This wider ‘Phytophthora community’ has, over time, not only generated dedicated books (e.g. Erwin et al. 1983; Lucas et al. 1991; Erwin and Ribeiro 1996; Lamour 2013) but an enormous scientific literature base. Currently there are ca 14,000 articles in Scopus (compared with ca 4000 for all 20 DM genera combined), and that literature continues to grow rapidly. This unifying scientific communication value would be seriously damaged by an inappropriate and unnecessary attempt to break up the genus.

The impact on biosecurity needs to be seen in the context of the extensive damage caused by Phytophthoras to cash crops and forests (Tables 1, 2; Tables S1, S2); the new epidemics and pandemics resulting from introductions of exotic Phytophthoras via international trade in plants and international travel (Fig. 2); and the many new Phytophthora species being discovered in underexplored ecosystems with the potential to cause further pandemics, especially in host plants they have not previously encountered. Because of these threats, coherent unambiguous communication about the genus is extremely important for developing sound, evidence-based biosecurity and plant health protocols at both the international and local scale. The current understanding among regulators and scientists about what is meant by Phytophthora when developing plant health regulation is a valuable asset in crop and habitat protection. An unnecessarily designation of multiple new genera could seriously damage this understanding, resulting in confusion and, at worse, weakened biosecurity, adding another threat to an already fragile global environment. These problems would probably be further exacerbated by the often long time lags between taxonomic changes and names being incorporated into plant health legislation or into extension programmes.

Regarding the scope for engendering confusion, many natural and managed ecosystems are inhabited by multiple Phytophthora species. For example, 27 and 39 Phytophthora species respectively, from seven clades in each case, have recently been found in the forests and natural ecosystems of Taiwan and Vietnam (Brasier et al. 2010; Jung et al. 2017b, d, 2020); while horticultural nurseries across Europe are infested by at least 65 Phytophthora taxa from nine clades (Table 3; Additional file 2: Table S2; Moralesjo et al. 2009; Jung et al. 2016). Moreover, the same disease syndromes are often caused by multiple Phytophthoras. At least 26 Phytophthora species from nine clades are associated with the current pan-European declines of oak forests (Table 3; Additional file 2: Table S2); and at least 51 species and hybrids are associated with damage to native plant communities around San Francisco (Table 3; Additional file 2: Table S2). So common are Phytophthoras on trees that commercial lateral flow devices are available to diagnose the genus (e.g. Tomlinson et al. 2010). Any designation of new genera that unnecessarily dissected the common biological properties of Phytophthoras would negatively impact communication and management in these and many similar situations and cause confusion among practitioners such as farmers, horticulturalists, forest managers and nursery owners reliant on scientific extension programmes for guidance.

The potential impact on verbal discourse deserves its own consideration. At a recent meeting of pathologists, forest health surveyors and plant health regulators in Britain addressing a previously unrecorded Phytophthora attacking Western Hemlock (Tsuga heterophylla) (Pérez-Sierra et al. 2022), discussion ranged across ten forest Phytophthora species from six different clades. To have referred to these by multiple generic names would have rendered the discussion unnecessarily complex and confusing to taxonomists and non-taxonomists alike. Similarly, upwards of 70 Phytophthora species across all 11 major clades are routinely discussed at the biennial International Forest Phytophthora Symposia. These meetings are an important channel of communication at a time of major Phytophthora threats to forests and natural ecosystems. Any unnecessary break-up of the genus would seriously undermine the value and purpose of this research community.

Numerous Government and NGO websites worldwide are dedicated to Phytophthora threats (e.g. Anonymous 1–4). Furthermore, terms such as ‘Phytophthora dieback’, ‘Phytophthora decline’ ‘Phytophthora root rot’, ‘Phytophthora collar rot’, ‘Phytophthora leaf blight’ and ‘Phytophthora bleeding canker’ are used widely in books, scientific papers and at meetings to discriminate the main disease syndromes common across the Phytophthora clades. The differences of meaning between the terms are well understood by academic plant pathologists, field surveyors and plant health regulators alike. In Britain and Australia terms such as ‘Phytophthora root rot’ and ‘Phytophthora dieback’ are even in common usage in horticultural magazines and the popular media. Broadly, these terms are an important component of the language centred around Phytophthora behaviour, disease management and biosecurity.

The risks to effective scientific communication inherent in over-zealous application of formal taxonomic practices in the context of molecular phylogenetics can be seen in the recent debate around Fusarium, another historic pathogen genus of high biosecurity importance. Regrettably, there now appears to be a damaging split in the international Fusarium community over what does, and does not, constitute the genus (Crous et al. 2021b;
CONCLUSIONS AND RECOMMENDATIONS

Arguably *Phytophthora* remains one of, if not the best-known and most important genus of plant pathogens. Despite the rapid increase in the number of described *Phytophthora* species and improvements in molecular phylogeny the genus has remained structurally coherent and biologically well understood. In our view, no one *Phytophthora* major clade or combination of clades exhibits a sufficiently distinct set of biological characteristics to warrant a unique generic status. Paraphyletic jumps, such as the emergence of the DMs from *Phytophthora* ancestors, should be considered a normal feature of evolution in ancient and successful genera such as this. Enthusiasm to ‘dice and slice’ along the lines of cladistic nuances should not trump evolutionary or biological coherence or overlook the fact that the primary purpose of names is to facilitate communication.

We are aware that under the orthodoxies, idiosyncrasies and sometimes vague constructs of the ICNafp (cf. Hawksworth 2020) a taxonomic restructuring of *Phytophthora* could be published relatively unchallenged by any author or group regardless of their familiarity with the genus as long as they follow certain somewhat subjective rules. ‘*Phytophthora*’ is surely now bigger than all of us. In which case its status needs to be policed by a wide consensus of the scientific community, perhaps through a recommendation of a working group of the International Commission on the Taxonomy of Fungi (ICFT). We contend therefore that any proposal for a major restructuring of the circumscription of the genus should be presented to and considered by an international working group of *Phytophthora* researchers, perhaps under ICTF auspices. There are already strong precedents for the international community coming together on aspects of *Phytophthora* research. For example, an international meeting debated the case ‘for or against diploidy in *Phytophthora*’ at the University of Bari, Italy in May 1972 (Brasier 2008); and there have been large International Symposia on *Phytophthora* at the University of California, Riverside in 1982 (Erwin et al. 1983) and at Trinity College, Dublin in 1989 (Lucas et al. 1991).

The issue might also be usefully addressed by a special session during the next International Mycological Congress in Maastricht in 2024.

While a case might be made for assigning sub-generic or section names to the various *Phytophthora* clades, as is the practice in some other large genera such as *Agaricus*, *Aspergillus*, *Cladonia*, *Hebeloma* and *Penicillium*, we doubt this would add much to our communication or understanding, and could be even more confusing to end users.

Considering all the above issues, and especially the lack of unequivocal evidence that defining *Phytophthora* clades as discrete genera would result in more biologically meaningful entities, we recommend that the current broad generic concept be retained. This would preserve the cultural history of the genus. It would also maintain the currently enormously effective value of the name *Phytophthora* in scientific communication, including for the many applied biologists and regulators dealing with *Phytophthoras* on a daily basis.

Supplementary Information

The online version contains supplementary material available at https://doi.org/10.1186/s43008-022-00097-z.

Acknowledgements

We thank Marilia Horta Jung (Mendel University in Brno) for redrawing the phylogenetic tree in Figure 3 from Scanu et al. (2021).

Author contributions

CMB, TJ, BS and DELC conceptualized this study, reviewed the *Phytophthora* and DM literature and interpreted the data. TJ, CMB and BS collated the data and analysed phenotypic traits for the tables and figures. CMB, TJ and BS wrote the first draft of the manuscript. All authors have read and agreed to the published version of the manuscript.

Funding

This work was supported by the Project Phytophthora Research Centre Reg. No. CZ.02.1.01/0.0/0.0/15_003/0000453 cofinanced by the Czech Ministry for Education, Youth and Sports and the European Regional Development Fund; by the ‘fondo di Ateneo per la ricerca 2019’, an internal funding provided by the University of Sassari; by the Scottish Government; and by Brasier Consultancy.
Availability of data and materials
Not applicable.

Declarations

Ethics approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Author details
1 Forest Research, Alice Holt Lodge, Farnham, Surrey GU10 4LH, UK. 2 Department of Agricultural Sciences, University of Sassari, Viale Italia 39A, 07100 Sassari, Italy. 3 James Hutton Institute, Invergowrie, Dundee DD2 5DA, UK. 4 Department of Forest Protection and Wildlife Management, Phytophthora Research Centre, Mendel University in Brno, 613 00 Brno, Czech Republic. 5 Phytophthora Research and Consultancy, 83131 Nussdorf, Germany.

Received: 7 March 2022   Accepted: 25 May 2022
Published online: 27 June 2022

References
Aguayo J, Halkett F, Husson C, Nagy ZÁ, Szigethy A, Bakonyi J, Frey P, Marçais B (2016) Genetic diversity and origins of the homoploid-type hybrid Phytophthora xalni. Appl Environ Microbiol 82:7142–7153
Anonymous 1 Government of Western Australia. 2 University of Berkely California. Presidio management of Agricultural Sciences, University of Sassari, Viale Italia 39A, 07100 Sassari, Italy. 3 The James Hutton Institute, Invergowrie, Dundee DD2 5DA, UK. 4 Department of Forest Protection and Wildlife Management, Phytophthora Research Centre, Mendel University in Brno, 613 00 Brno, Czech Republic. 5 Phytophthora Research and Consultancy, 83131 Nussdorf, Germany.

Bourke A (1991) Potato blight in Europe in 1845. The scientific controversy. In: Lucas JA, Shattock RC, Shaw DS, Cooke LR (eds) Phytophthora. Cambridge Univ. Press, Cambridge, pp 12–24
Bourret T, Choudhury RA, Mehl HK, Blomquist CL, McRoberts N, Rizzo DM (2018) Multiple origins of downy mildews and mito-nuclear discordance within the paraphyletic genus Phytophthora. PLoS ONE 13:e0192502. https://doi.org/10.1371/journal.pone.0192502
Brasier CM (1972) Observations on the sexual mechanism in Phytophthora palmivora and related species. Trans Brit Mycol Soc 58:237–251
Brasier CM (1983) Problems and prospects in Phytophthora research. In: Erwin DC, Tsao PH, Bartnicki-Garcia S (eds) Phytophthora, its biology, ecology and pathology. American Phytopathological Society, St Paul, Minnesota, pp 351–364
Brasier CM (1991) Current questions in Phytophthora systemsatics: The role of the population approach. In: Lucas JA, Shattock RC, Shaw DS, Cooke LR (eds) Phytophthora. Cambridge Univ. Press, Cambridge, pp 104–128
Brasier CM (1992) Evolutionary biology of Phytophthora part I: genetic system, sexuality and the generation of variation. Annu Rev Phytopathol 30:153–171. https://doi.org/10.1146/annurev.phytopath.30.090919.001101
Brasier CM (1995) Episodic selection as a force in fungal microevolution with special reference to clonal speciation and hybrid introgression. Can J Bot 73:1213–1221. https://doi.org/10.1139/b95-381
Brasier CM (2001) Rapid evolution of introduced plant pathogens via interspecific hybridization. Bioscience 51:123–133. https://doi.org/10.1641/0006-3568(2001)051[123:REOPPI]2.0.CO;2
Brasier CM (2008) The biosecurity threat to the UK and global environment from international trade in plants. Plant Pathol 57:792–808. https://doi.org/10.1111/j.1365-3599.2008.01886.x
Brasier CM (2009) Phytophthora biodiversity: How many Phytophthora species are there? In: Goheen EM, Frankel SJ (eds) Phytophthoras in Forests and Natural Ecosystems: Fourth Meeting of the International Union of Forest Research Organizations (IUFRO) Working Party S07.02.09, General Technical Report PSW-GTR-221, USDA Forest Service, Pacific Southwest Research Station, Albany, California, pp 101–115
Brasier CM, Cooke DEL, Duncan JM (1999) Origins of a new Phytophthora pathogen through interspecific hybridisation. P Natl Acad Sci USA 96:5878–5883. https://doi.org/10.1073/pnas.96.10.5878
Brasier CM, Cooke DEL, Duncan JM, Hansen EM (2003) Multiple new pheno-typic taxa from trees and riparian ecosystems in Phytophthora gonapodyides-P. megasperma ITS clade 6, which tend to be high-temperature tolerant and either inbreeding or sterile. Mycol Res 107(3):277–290. https://doi.org/10.1017/S0953756203007380
Brasier CM, Franceschini S, Vetraino AM, Hansen EM, Green S, Robin C, Webber JF, Vannini A (2012) Four phenotypically and phylogenetically distinct lineages in Phytophthora lateralis. Fungal Biol 116(12):1232–1249. https://doi.org/10.1016/j.fungalbiol.2012.10.002
Brasier CM, Kirk SA, Delcan J, Cooke DEL, Jung T, Man int’t Veld WA (2010) Phytophthora alni sp. nov. and its variants: designation of emerging lineages in Phytophthora lateralis. J Phytopathol 158:595–603. https://doi.org/10.1111/j.1365-3059.2009.01405.x
Brasier CM, Robredo F, Ferraz JFP (1993) Evidence for Phytophthora cinnamomi from international trade in plants. Phytopathology 83:1237–1239. https://doi.org/10.1094/PHYTO.1993.83.10.1237
Brasier CM, Vettraino AM, Chang TT, Vannini A (2010) Four phenotypically and phylogenetically distinct lineages in Phytophthora lateralis. Fungal Biol 116(12):1232–1249. https://doi.org/10.1016/j.fungalbiol.2012.10.002
Brasier CM (1991) Sudden larch death. Nature 466:824–825. https://doi.org/10.1038/366824a
Brasier CM, Vettraino AM, Chang TT, Vannini A (2010) Phytophthora lateralis discovered in an old growth Chamaecyparis forest in Taiwan. Plant Pathol 59:595–603. https://doi.org/10.1111/j.1365-3059.2010.02278.x
Brasier CM, Webber J (2010) Sudden larch death. Nature 466:824–825. https://doi.org/10.1038/466824a
Brooks AV, Brasier CM (2007) Colonization of tree xylem by Phytophthora ramorum, P. kernoviae and other Phytophthora species. Plant Pathol 56:227–241. https://doi.org/10.1111/j.1365-3059.2006.01511.x
Brummitt RK, Sosef MSM (1998) Paraphyletic taxa are inherent in Linnaean classification: a reply to Freudenstein. Taxon 47:411–412. https://doi.org/10.2307/2267771
Brummitt RK (2003) Further dogged defence of paraphyletic taxa. Taxon 52:803–804. https://doi.org/10.2307/3647353
**Phytophthora ramorum**. Mol Plant Microbe in 19:1348–1358. https://doi.org/10.1094/MPMI-19-1348

Man in’t Veld WA, Veenbaas-Rijks WJ, Ilieva E, de Cock AWAM, Bonants PJM, Pieters R (1998) Natural hybridization of Phytophthora nicotianae and P. cactorum demonstrated by isozyme analysis and random amplified polymorphic DNA. Phytopathology 88:922–929. https://doi.org/10.1094/phyto.1998.88.9.922

Man in’t Veld WA, Rosendahl KCHM, Hong C (2012) *Phytophthora x serendipita* sp. nov. and *P. x pelgrandis*, two destructive pathogens generated by natural hybridization. Mycologia 104:1390–1396. https://doi.org/10.3852/11-272

Martin FN, Blair JE, Coffey MD (2014) A combined mitochondrial and nuclear multilocus phylogeny of the genus *Phytophthora*. Fungal Genet Biol 66:19–32. https://doi.org/10.1016/J.FGB.2014.02.006

Martin FN,Tookey PW (2003) Phylogenetic relationships among *Phytophthora* species inferred from sequence analysis of multizonally encoded cytochrome oxidase I and II genes. Mycologia 95:269–284. doi.org/10.2307/3758706

Matai NH, Blair JE (2014) A multilocus timescape for oomycete evolution estimated under three distinct molecular clock models. BMC Evol Biol 14:101

Mayr E, Bock WJ (2002) Classifications and other ordering systems. J Zool 357–379

McCarthy CGP, Fitzpatrick DA (2017) Phylogenomic reconstruction of the oomycete phylogeny derived from 37 genomes. Nature 541:165–170. https://doi.org/10.1038/nature21111

Montagne JFC (1845) Note sur la maladie qui ravige les pommes de terre et caracteres du *Botrytis infectans*. (Note on the disease that ravages potatoes and characters of *Botrytis infectans*). Bull Soc Sci Philom Paris 13:312–313

Moralez E, Pérez-Sierra A, Alvarez LA, Belbañí L, Lefort F, Descals E (2009) Multiple alien *Phytophthora* taxa discovered on diseased ornamental plants in Spain. Plant Pathol 58:100–110. https://doi.org/10.1111/j.1365-3059.2008.01930.x

Moralez E, Puig M, García JA, Descals E (2006) Stromata, sporangiomata and chlamydosori of *Phytophthora ramorum* on inoculated Mediterranean woody plants. Mycol Res 110:1323–1332. https://doi.org/10.1017/j.mycres.2006.09.004

Newhoock FJ, Waterhouse GM, Stamps DJ (1978) Tabular key to the species of *Phytophthora* de Bary. Commonwealth Mycological Institute, Kew, UK. Mycol Pap. 143, p 20

Oh E, Gray-henmouth M, Wingfield BD, Wingfield MJ, Burgess TI (2013) Surveys of soil and water reveal a goldmine of *Phytophthora* diversity in South African natural ecosystems. IMA Fungus 4:123–131. https://doi.org/10.5598/imafungus.2013.04.01.12

Ortiz BR (2008) Contemporary Californian Indians, oaks, and Sudden Oak Death (*Phytophthora ramorum*). In: Merenlender A, McCreary D, Pur- death: *Phytophthora ramorum*) in California. Plant Dis 86:205–214. https://doi.org/10.1094/POIS.2008.86.3.205

Rea AJ, Burgess TJ, Hardy GESTJ, Strukely MJC, Jung T (2011) Two novel and potentially endemic species of *Phytophthora* associated with epizootic dieback of kwongan vegetation in the south-west of Western Australia. Plant Pathol 60:1055–1068. https://doi.org/10.1111/j.1365-3059.2011.02463.x

Reeser PW, Sutton W, Hansen EM, Remigi P, Adams GC (2011) *Phytophthora* species in forest streams in Oregon and Alaska. Mycologia 103:22–35. https://doi.org/10.3852/10-013

Rizzo DM, Garbelotto M, Davidson JM, Slaughter GW, Koike ST (2002) *Phytophthora ramorum* as the cause of extensive mortality of Quercus spp. and *Lithocarpus densiflorus* in California. Plant Dis 86:205–214. https://doi.org/10.1094/POIS.2002.86.3.205

Robideau GP, De Cock AWAM, Coffey MD, Voolmajay R, Brouwer H, Bala K, Chitty DIW, Desaulniers N, Eggertson QA, Gachon CMM, Hu C-H, Kupper FC, Rintoul TL, Sarhan E, Verstappen ECP, Zhang Y, Bonants PJM, Ristaino JB, Lévesque CA (2011) DNA barcoding of oomycetes with cytochrome c oxidase subunit I and internal transcribed spacer. Mol Ecol Resour 11:1002–1011. https://doi.org/10.1111/j.1471-8286.2011.02041.x

Rondot RL (1935) Vergleichende Untersuchungen über die Wirkung verschiedener Wuchsstoffe auf das Wachstum einiger Pilze. Arch Mikrobiol 63:309–325

Ross HA (2014) The incidence of species-level paraplygy in animals: a reassessment. Mol Phylogenet Evol 76:10–17. https://doi.org/10.1016/j.mpev.2014.02.021

Rosenbaum J (1917) Studies of the genus *Phytophthora*. J Agric Res 8:233–276

Runge F, Telle S, Ploch S, Savory E, Day B, Sharma R, Thines M (2011) The inclusion of downy mildews in a multi-locus-dataset and its reanalysis reveals a high degree of paraphyly in *Phytophthora*. IMA Fungus 2:163–171. https://doi.org/10.5598/imafungus.2011.02.02.07

Safaiefarhani B, Mostowfizadeh-Ghalamfarsa R, Hardy GESTJ, Burgess TE (2015) Reevaluation of the *Phytophthora cryptogaea* species complex and a description of a new species *Phytophthora pseudocryptogaea* sp. nov. Mycol Prog 14:108. https://doi.org/10.1007/s11557-015-1129-9

Samson RA, Hubika V, Varga J, Hourbaken J, Hong S-B, Klaassen CHW, Perrone G, Seiffert KA, Magista D, Visagie CM, Kocsube S, Szegeti G, Yuguchi T, Peterson SW, Frisvad JC (2017) Response to Pitt & Taylor 2016: conserva- tion of Aspergillus with *A. niger* as the conserved type is unnecessary and potentially disruptive. Taxon 66:1439–1446. https://doi.org/10.12705/6606.10

Sanso E (1961) Meiosis in the oogonium and antheridium of *Pythium debaryanum* Hesse. Natur 191:827–828

Sanso E (1963) Meiosis in *Pythium debaryanum* Hesse and its significance in the life history of the biflagellates. Trans Brit Mycol Soc 46:63–72

Sanso E (1965) Meiosis in diploid and polyploid sex organs of *Phytophthora* and *Achlya*. Cytologia 30:103–117

Sanso E (1966) Meiosis in the sex organs of the Oomyctes. In: Darlington CD, Lewis KR (eds) Chromosomes today, vol 1. Oliver and Boyd, Edin- burgh, pp 77–83

Sanso E, Brasier CM (1973) Diploidy and chromosomal structure hybridity in *Pythium* infestans. Nature 241:344–345

Sanso E (1980) Reciprocal translocation heterozygosity in heterothal- lic species of *Phytophthora* and its significance. Trans Brit Mycol Soc 74:175–185. https://doi.org/10.2246/PHY/74-7-185

Scaru B, Jung T, Maisoal H, Linaldeedu BT, Horta Jung M, Brândano A, Mostowfizadeh-Ghalamfarsa R, Janoušek J, Riolo R, Cacciola SO (2021) *Phytophthora heterospora* sp. nov., a new pseudocandin-producing sister species of *P. palmivora*. J Fungi 7:870. https://doi.org/10.3390/jof7100807

Scaru B, Linaldeedu BT, Deidda A, Jung T (2015) Diversity of *Phytophthora* species from declineing Mediterranean maquis vegetation, including two new species, *Phytophthora cinnamomi* and *P. omanamenta* sp. nov. PLoS ONE 10:e0143234. https://doi.org/10.1371/journal.pone.0143234

Shearer BL, Crane CE, Cochrane A (2004) Quantification of the susceptibility of the native flora of the South-West Botanical Province, Western Australia, to *Phytophthora cinnamomi*. Aust J Bot 52:435–443. https://doi.org/10.1071/BO03131
