

Standard Paper

Coenogonium nimisii – a new isidiate epiphytic lichen similar to Porina rosei

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Abstract

Our floristic work in British ancient forests resulted in a description of a frequently reported but misidentified species, *Coenogonium nimisii*. Its thallus is very similar to *Porina rosei*, but the apothecia and pycnidia correspond with *C. luteum*. Sterile collections are not easy to distinguish but the new species differs from *P. rosei* in several microscopic characters of the isidia. *Coenogonium nimisii* is so far known from bark and epiphytic bryophytes, rarely mossy rocks, in ancient humid forests of Great Britain and Ireland. The genus *Coenogonium* is poorly represented by molecular data in the GenBank database. Our preliminary results revealed distinct genetic lineages within two traditionally circumscribed species, *C. luteum* and *C. pineti*, which may represent cryptic species.

Keywords: ancient forests; cryptic species; Great Britain; phylogeny; taxonomy; *Trentepohliaceae*

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Introduction

Coenogonium (*Ostropales*, *Coenogoniaceae*) is a large genus of crustose or filamentous lichenized fungi, currently including c. 90 species (Rivas Plata *et al.* 2006; Lücking *et al.* 2017). It is distributed almost worldwide with its centre of diversity in the tropics (Kantvilas *et al.* 2018). In Europe, three species are so far known: *C. luteum* (Dicks.) Kalb & Lücking, *C. pineti* (Ach.) Lücking & Lumbsch and *C. tavaresianum* (Vězda) Lücking *et al.* In addition, five tropical species have been reported from Macaronesia: *Coenogonium frederici* (Kalb) Kalb & Lücking (Aptroot & Rodrigues 2005; as *Dimerella frederici*), *Coenogonium implexum* Nyl. (Rodrigues & Aptroot 2005), *C. interplexum* (Tavares 1952), *C. luteolum* (Kalb) Kalb & Lücking (Kalb & Hafellner 1992; as *D. luteola*) and *C. subluteum* (Rehm) Kalb & Lücking (Follmann 1990; as *D. epiphylla*).

During our fieldwork in British ancient forests, we collected isidiate thalli of ‘*Porina rosei*’ with apothecia or pycnidia of ‘*Coenogonium luteum*’. These strange specimens were sequenced and we revealed an undescribed *Coenogonium* species. Subsequently, this taxon was mentioned in the *Revisions of British and Irish Lichens* (Vol. 3) under the provisional name *C. confusum* Malíček & Sanderson *ined.* (Cannon *et al.* 2021). Finally, we decided to describe it here formally as *C. nimisii*.

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Material and Methods

Sampling, morphology and chemistry

Collected specimens are deposited in PRA and the personal herbaria of J. Malíček and N. Sanderson. Microscopic descriptions are based on hand-cut sections mounted in water or 10% KOH for observations of the isidia surface. Lichen secondary metabolites were identified using thin-layer chromatography (TLC) in solvents A, B’ and C (Orange *et al.* 2010). The images were captured using 1) an Olympus SZX 12 stereomicroscope with an Olympus DP 70 (resolution 12.5 Mpx) cooled colour digital camera with the software QuickPHOTO MICRO 3.0 (Promicra), using an extended depth of field module Deep Focus, and 2) an Olympus BX 43 microscope with a Promicra 3–5CP (resolution 5 Mpx) colour digital camera with the same software.

DNA extraction, PCR amplification and sequencing

The Invisorb Spin Plant Mini Kit (Invitex) and Chelex protocol (Ferencová *et al.* 2017) were used for DNA extractions. The fungal ITS rDNA (henceforth ITS) and mitochondrial SSU (mtSSU) were amplified with the following primers: ITS1F (Gardes & Bruns 1993) and ITS4 (White *et al.* 1990), mrSSU1, mrSSU2R and mrSSU3R (Zoller *et al.* 1999). The mycobiont from three samples of *Porina rosei* was amplified with ITS-A (Blattner 1999; Škaloud & Rindi 2013) and ITS4. PCR reactions of ITS and mtSSU were prepared for a 20 µl final volume, containing 14 µl double-distilled water, 4 µl MyTaq polymerase reaction buffer, 0.2 µl MyTaq DNA polymerase, 0.4 µl of each of the 25 mM primers, and 1 µl of the sample. Amplifications of both loci consisted of an initial 1 min denaturation at 95 °C, followed by 35



cycles of 30 s at 95 °C, 30 s at 56 °C, 30 s at 72 °C, and a final extension of 7 min at 72 °C. For the combination ITS-A and ITS4, a cycle was programmed with an initial 1 min denaturation at 94 °C, followed by 30 cycles of 25 s at 94 °C, 30 s at 54 °C, 45 s at 72 °C, and a final extension of 10 min at 72 °C. The PCR products were visualized on a 0.8% agarose gel and cleaned with ExoSAP-IT™ PCR Product Cleanup Reagent (ThermoFisher Scientific), according to the manufacturer's protocols. The algal ITS and *rbcl* sequences were amplified as described by Černajová *et al.* (2022). In total, 19 new ITS, 19 mtSSU and 9 *rbcl* sequences were generated (Table 1). Sequences of the *Porina rosei* mycobiont, photobiont and *Coccomyxa* ITS were not included in the phylogenetic analysis.

Sequence alignment and phylogenetic analysis

The newly produced sequences were edited in BioEdit v. 7.2.5 (Hall 1999). The final analyses of the mycobiont included the 28 newly generated sequences and nrITS and mtSSU sequences of the genus *Coenogonium* available in the GenBank database. *Gyalecta jenensis* and *G. russula* were selected as outgroups because they form a sister clade to *Coenogonium* (see Miadlikowska *et al.* 2014) and both ITS and mtSSU sequences were available in GenBank. The ITS and mtSSU regions were aligned separately using MAFFT v. 7 (Katoh & Standley 2013) with the L-INS-i method (Katoh *et al.* 2005). The ITS alignment contained 531 positions and 20 sequences; the mtSSU alignment had 719 positions and 25 sequences. Both regions were analyzed as single-locus datasets (see Supplementary Material Figures S1 & S2, available online). For the final tree, we created a concatenated dataset of 27 sequences, containing 462 ITS positions and 719 mtSSU positions. Ambiguous positions (i.e. > 50% of missing data) were excluded from the final analysis. Support values on nodes were checked against single-gene trees and no conflict among well-supported branches (maximum likelihood bootstrap percentages > 0.7) was detected.

The photobiont analysis was based on seven newly generated *rbcl* sequences and 88 sequences retrieved from GenBank (16 reported as *Coenogonium* photobionts, 10 as *Porina* photobionts and 62 other reference sequences, both lichen photobionts and free-living algae). The *rbcl* alignment contained 87 unique sequences and 752 positions.

Phylogenetic trees were inferred with Bayesian inference (BI) using MrBayes v. 3.2.7a (Ronquist & Huelsenbeck 2003; Ronquist *et al.* 2012), maximum likelihood (ML) analysis using GARLI v. 2.0 (Zwickl 2006), and maximum parsimony (MP) analysis using PAUP v. 4.0b10 (Swofford 2003). BI and ML analyses were carried out on a partitioned dataset to differentiate among ITS1, 5.8S and ITS2 rDNA, and mtSSU regions for mycobionts. For the photobiont *rbcl* tree, the dataset was partitioned according to particular codon positions. Substitution models TN93 + G (ITS1), JC + G (5.8S), K2 + I (ITS2), T92 + G (mtSSU), and JC + G + I, T92 + G and K2 + G + I (*rbcl*) were selected using the Bayesian information criterion (BIC) as implemented in jModelTest2 (Guindon & Gascuel 2003; Darriba *et al.* 2012). Two parallel MCMC runs, with four chains, were carried out for 10 million generations. Trees and parameters were sampled every 100 generations. Finally, the burn-in values were determined using the 'sump' command. The ML analysis was carried out using default settings, five search replicates, and the automatic termination set at 5 million generations. The MP analysis was performed using heuristic searches with 1000 random sequence

addition replicates and random addition of sequences (the number was limited to 10⁴ per replicate). ML and MP bootstrap support values were obtained from 100 and 1000 bootstrap replicates, respectively. Only one search replicate was applied for ML bootstrapping. The alignment used in this study is publicly available on Zenodo as doi: 10.5281/zenodo.7627869.

Taxonomy

Coenogonium nimisii Malíček & Sanderson sp. nov.

Mycobank No.: MB 847489

Similar to *Coenogonium fruticulosum* L. Ludw., so far known from New Zealand, but different in several anatomical characters, such as the higher hymenium (80–110 µm), and larger photobiont cells (10–18 µm) and ascospores (9–11 µm). The isidia of *C. nimisii* are 25–70 µm thick, glossy in fresh material, with one or rarely more layers of thin periclinal hyphae. Apothecia and pycnidia rarely present, ±identical with *C. luteum*.

Type: Great Britain, England, V.C. 11, by Penderley Lodge, Stubbs Wood, New Forest, veteran *Quercus robur* in pasture woodland, 2020, *N. Sanderson* 2744 (PRA 21373—holotype). GenBank Accession nos: OQ366546 and OQ366530.

(Fig. 1)

Thallus crustose, thin, up to 40 µm thick, pale (green-)grey to whitish, without prothallus, often with yellow/ochre tinge, with abundant isidia, forming dense mounds away from the apothecia, more discrete near apothecia. *Isidia* coralloid, moniliform (i.e. with frequent constrictions), green and glossy in fresh material, pale grey-green, grading to ochre-orange where well lit, up to 0.5 mm high, 25–70 µm diam., filled with very abundant photobiont cells with 3–5 cells per isidium width in optical view, cortex poorly defined and composed of one or rarely more layers of colourless periclinal hyphae which are more visible in fresh material or KOH. *Photobiont* trentepohlioid, ±globose cells (7–)10–18 µm diam.

Apothecia rare, (0.5–)1–1.5 mm diam., sessile on the thallus (rarely formed on isidia), distinctly constricted at the base. *Disc* yellow-orange to dark orange, with creamy to yellow, sometimes flexuose margins. *Exciple* colourless, outermost part yellow-brown in dark-pigmented apothecia, composed of branching, conglutinated, 5–8 µm thick hyphae. *Hypothecium* colourless. *Hymenium* 80–110 µm tall, colourless, I+ brownish orange to locally pale blue, K/I+ blue; *epihymenium* pale yellow-brown in dark-pigmented apothecia; *paraphyses* 1.5–2.0 µm thick, upper cells sometimes moniliform, the apical cell up to 4.5 µm diam. *Asci* narrowly cylindrical, c. 30 × 4 µm, without an amyloid ring around the pore, 8-spored, uniseriate. *Ascospores* colourless, fusiform-ellipsoid, 1-septate, straight or rarely one cell slightly curved, (7–)9–11(–14) × 2.5–3(–3.5) µm (Supplementary Material Figure S3, available online).

Pycnidia rare, ±globose, 0.3–0.5 mm diam., sessile to shallowly immersed in the thallus and substratum, beige, pycnidia wall colourless in section, 15–30 µm thick, composed of branched hyphae of mostly parallel orientation. *Conidia* aseptate, colourless, ellipsoid, 3.5–5 × 2 µm.

Chemistry. No lichen compounds detected by TLC. Spot reactions: C–, K–, KC–, Pd–, UV– or UV+ whitish grey or ochre (parts with the orange/ochre pigment).

Table 1. GenBank Accession numbers and voucher information for sequenced specimens of *Coenogonium*. Sequences in bold are newly produced.

Taxon	Source - Specimen	Mycobiont ITS	Mycobiont mtSSU	Algal ITS	Algal <i>rbcL</i>
<i>Coenogonium isidiatum</i>	Russia, Sakhalin, Shikotan Island, <i>S. Chesnokov</i> 172 (LE-L-14438)	MH179135	MH179140	–	–
<i>C. isidiatum</i>	Russia, Sakhalin, Iturup Island, <i>L. Konoreva</i> 606 (LE-L-14436)	MH179137	MH179142	–	–
<i>C. isidiatum</i>	Russia, Sakhalin, Iturup Island, <i>L. Konoreva</i> 618 (LE-L-14437)	MH179136	MH179141	–	–
<i>C. isidiatum</i>	Russia, Sakhalin, Sakhalin Island, <i>L. Konoreva</i> 198 (LE-L-14441)	MH179138	–	–	–
<i>C. isidiatum</i>	Russia, Kamchatka, <i>I. Stepanchikova</i> Nik-1790 (LE-L-14442)	–	MH179143	–	–
<i>C. luteolum</i>	Portugal, Madeira, São Jorge, <i>J. Malíček</i> 14105 (hb. <i>J. Malíček</i>)	OQ366541	OQ366534	–	–
<i>C. luteum</i> s. lat.	Austria, Niederösterreich, Rothwald, <i>J. Vondrák</i> 26140 et al. (PRA)	OQ366542	OQ366535	–	–
<i>C. luteum</i> s. lat.	Great Britain, Scotland, Lochgilphead, <i>J. Malíček</i> 9141 et al. (hb. <i>J. Malíček</i>)	–	OQ366536	–	–
<i>C. luteum</i> s. lat.	Russia, Caucasus, <i>J. Malíček</i> 10624 et al. (hb. <i>J. Malíček</i>)	–	OQ366537	–	–
<i>C. luteum</i> s. lat.	USA, California, Santa Rosa Island, <i>B. Ryan</i> 31430 (ASU)	HQ650710	AY584699	–	–
<i>C. luteum</i> s. str.	Great Britain, Scotland, Lochgilphead, <i>J. Malíček</i> 12550 et al. (hb. <i>J. Malíček</i>)	–	OQ366526	–	–
<i>C. luteum</i> s. str.	Great Britain, Scotland, Oban, <i>J. Malíček</i> 12390 et al. (hb. <i>J. Malíček</i>)	–	OQ366527	–	–
<i>C. luteum</i> s. str.	Portugal, Madeira, Ponta do Sol, <i>J. Malíček</i> 14095 (hb. <i>J. Malíček</i>)	OQ366548	OQ366528	–	–
<i>C. nimisii</i>	Great Britain, England, New Forest, <i>J. Vondrák</i> 19844 (PRA)	OQ366545	OQ366532	OQ371695	OQ400977
<i>C. nimisii</i>	Great Britain, England, New Forest, <i>J. Vondrák</i> 19912 (PRA)	OQ366544	OQ366529	–	OQ400978
<i>C. nimisii</i>	Great Britain, England, New Forest, <i>M. Powell</i> 4685 (hb. <i>J. Malíček</i>)	OQ366543	OQ366531	OQ371694	OQ400974
<i>C. nimisii</i>	Great Britain, England, New Forest, <i>N. Sanderson</i> 2744 (PRA-00021373, holotype)	OQ366546	OQ366530	–	OQ400975
<i>C. nimisii</i>	Great Britain, Scotland, Oban, <i>J. Malíček</i> 12373 et al. (hb. <i>J. Malíček</i>)	OQ366547	OQ366533	OQ371693	OQ400973
<i>C. pineti</i>	Canada, Ontario, Waterloo Region, <i>R. T. McMullin</i> (BIOUG24047-G12)	KT695346	–	–	–
<i>C. pineti</i>	Czech Republic, Southern Bohemia, Šumava Mts, <i>J. Vondrák</i> 23798 & <i>S. Svoboda</i> (PRA)	OQ366550	OQ366538	–	–
<i>C. pineti</i>	Czech Republic, Central Bohemia, Český kras, <i>Z. Palice</i> 32241 (PRA)	OQ366551	OQ366539	–	–
<i>C. pineti</i>	Czech Republic, Southern Bohemia, Šumava Mts, <i>Z. Palice</i> 32447 (PRA)	OQ366552	OQ366540	–	–
<i>C. pineti</i>	Germany, Nordrhein-Westfalen, Solingen, <i>Z. Palice</i> 15531 (PRA)	–	AY300884	–	–
<i>C. pineti</i>	USA, Alaska, Glacier Bay, <i>T. Spribille</i> 39097 (GZU)	–	KR017337	–	–
<i>C. tavaresianum</i>	Great Britain, England, New Forest, <i>J. Vondrák</i> 19776 et al. (PRA)	OQ366549	OQ407486	–	–
<i>Gyalecta jenensis</i>	Canada, British Columbia, Selkirk Mts, <i>T. Spribille</i> s. n. (GZU)	KR017099	KR017330	–	–
<i>G. russula</i>	Sweden, Dalarna, <i>J. Hermansson</i> 14140 (UPS)	HM244759	HM244735	–	–
<i>Porina rosei</i>	Czech Republic, Šumava Mts, <i>J. Malíček</i> 12126 & <i>Z. Palice</i> (hb. <i>J. Malíček</i>)	–	OQ407485	OQ371692	OQ400972
<i>P. rosei</i>	Russia, Caucasus, <i>J. Vondrák</i> 15450 (PRA)	OQ407480	OQ407483	–	OQ400976
<i>P. rosei</i>	Russia, Caucasus, <i>J. Vondrák</i> 22903 (PRA)	OQ407482	OQ407484	–	OQ400979
<i>P. rosei</i>	Russia, Caucasus, <i>J. Vondrák</i> 22938 (PRA)	OQ407481	–	–	OQ400980

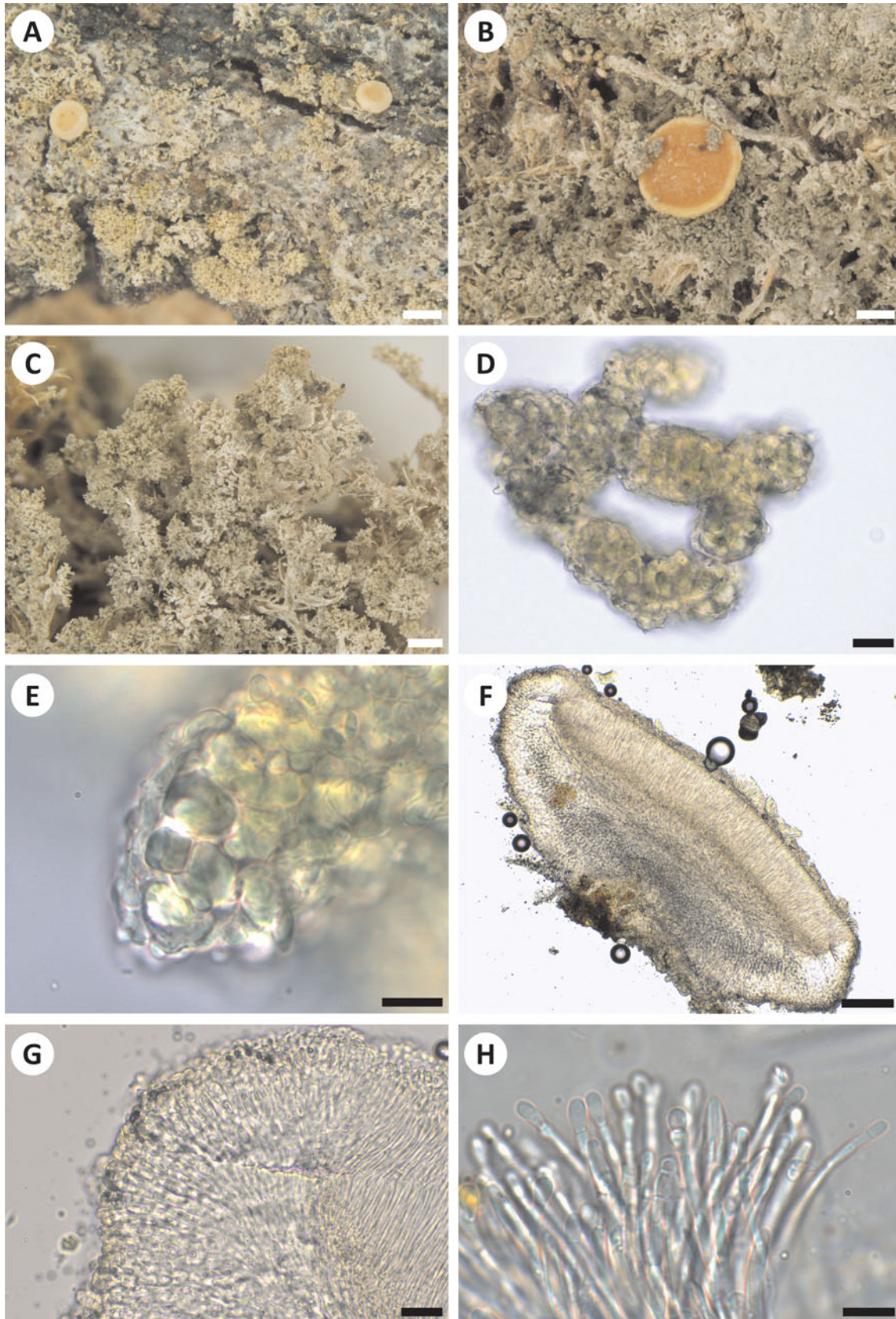


Figure 1. Habitus of *Coenogonium nimisii*. A, holotype (PRA). B, apothecium (Sanderson 436). C, isidia (Sanderson 436). D, isidia in water (Malíček 12373). E, isidia in KOH (Malíček 12373). F, apothecial section in water (holotype, PRA). G, exciple in water (holotype, PRA). H, paraphyses in water (Sanderson 436). Scales: A–C = 0.5 mm; D & G = 20 μ m; E & H = 10 μ m; F = 100 μ m. In colour online.

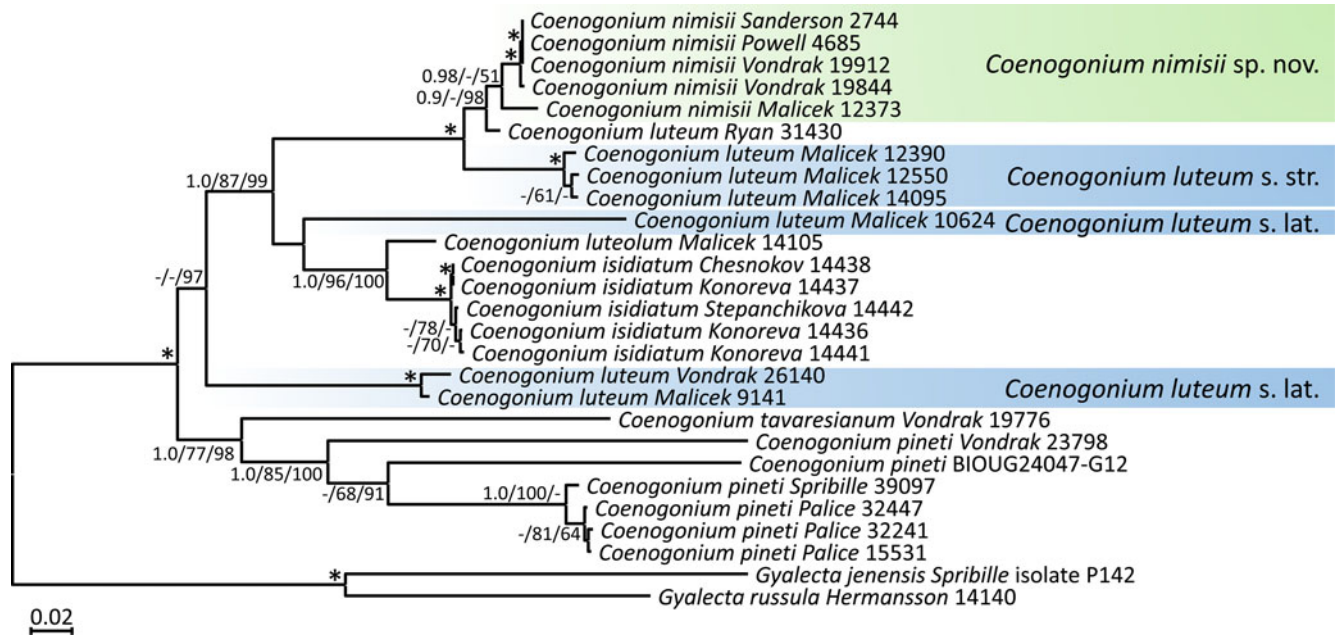


Figure 2. Phylogenetic hypothesis of *Coenogonium* resulting from the Bayesian analysis of combined ITS rDNA and mtSSU sequences. Values at nodes indicate the statistical supports of Bayesian posterior probability (left), maximum-likelihood bootstrap (middle) and maximum parsimony bootstrap (right). Fully supported branches (1.0/100/100) are marked with an asterisk. Scale bar shows the estimated number of substitutions per site. In colour online.

Phylogeny. The new species forms a well-supported clade in the phylogenetic tree (Fig. 2). The sequences of *C. nimisii* are very uniform with only one change in the mtSSU region. However, unexpectedly large variability was observed in the Scottish specimen (Malíček 12373), which differs in 31 ITS positions and four mtSSU positions. The new species is very closely related to the North American specimen of *C. luteum* (Ryan 31430), which could theoretically represent a non-isidiate form of *C. nimisii*. Nevertheless, the original collection has been examined only as a picture.

Etymology. The specific epithet honours Pier Luigi Nimis, the famous Italian lichenologist, who celebrates his 70th birthday this year.

Habitat and distribution. On a wide range of tree species with base-rich bark, especially on *Quercus robur*, frequently overgrowing epiphytic mosses and liverworts; rarely also on mossy rocks. *Agonimia octospora* Coppins & P. James, *Lepraria finkii* (B. de Lesd.) R.C. Harris and *Thelotrema lepadinum* (Ach.) Ach. are examples of associated species. *Coenogonium nimisii* is a typical species of ancient oceanic woodlands, so far known only from Great Britain and Ireland. It is widespread but local, known from SW England, Wales, SW Scottish Highlands and Ireland.

Notes. In Great Britain and Ireland *C. nimisii* is very rarely fertile and has been long mistaken for the very similar looking but much rarer *Porina rosei* (Fig. 3A). True *P. rosei* is also often sterile and isidia are very variable in colour, size and shape in both species. However, the two species differ mainly in microscopic characters of their isidia. The poorly defined cortex in *C. nimisii* is formed by periclinal hyphae, in contrast to a thick layer of irregular rounded cells in *P. rosei*, as demonstrated by Sérusiaux (1991) and Orange *et al.* (2020). In poorly developed isidia, we recommend observing this character carefully in KOH.

Additionally, isidia in *P. rosei* are more slender (20–50 µm thick), dull in fresh material, thin isidia contain only one or two layers of photobiont cells (Fig. 3B), they are squarrose when richly branched and contain an unknown pigment which is K+ pale brown directly after application and pale purple after several hours (Fig. 3B & C).

Coenogonium nimisii and *Porina rosei* also differ in their trentepohlioid photobionts despite their morphological similarity. *Porina rosei* photobionts in our phylogenetic hypothesis, based on the *rbcl* gene (Supplementary Material Figure S4, available online) were recovered in clade 1 *sensu* Nelsen *et al.* (2011), together with all previously published *Porina* photobionts regardless of their geographical origin (including the tropics). The photobionts of *Porina rosei* originating from the Caucasus and Czech Republic were identical or closely related to those of *Porina leptalea* (Durieu & Mont.) A.L. Sm. from France in *rbcl* sequences (Borgato *et al.* 2022). *Phycopeltis*, which has been repeatedly mentioned as the photobiont of *Porina* spp. (Grube *et al.* 2017; McCarthy & Kantvilas 2017), could be placed within clade 1 (Zhu *et al.* 2015). However, the genera *Trentepohlia*, *Printzina* and *Phycopeltis* are polyphyletic (Škaloud *et al.* 2018); their taxonomic revision is beyond the scope of this study and we therefore maintain the clade nomenclature introduced by Nelsen *et al.* (2011). *Coenogonium nimisii* photobionts were recovered in clade 2 together with the majority of previously published *Coenogonium* photobionts (Supplementary Material Figure S4). According to the BLAST search, the algal ITS sequences retrieved from both lichen species are congruent with *rbcl* sequences in terms of the closest relatives. From one sample of both lichen species, we obtained a sequence of *Coccomyxa* sp. and also observed its colonies (Fig. 3D) several times. The observations indicate the colonies are probably free-living but in a close association with the lichens.

Apothecia of *C. luteum* s. lat. are almost identical to those of *C. nimisii*. We observed only a slightly lower hymenium (up to

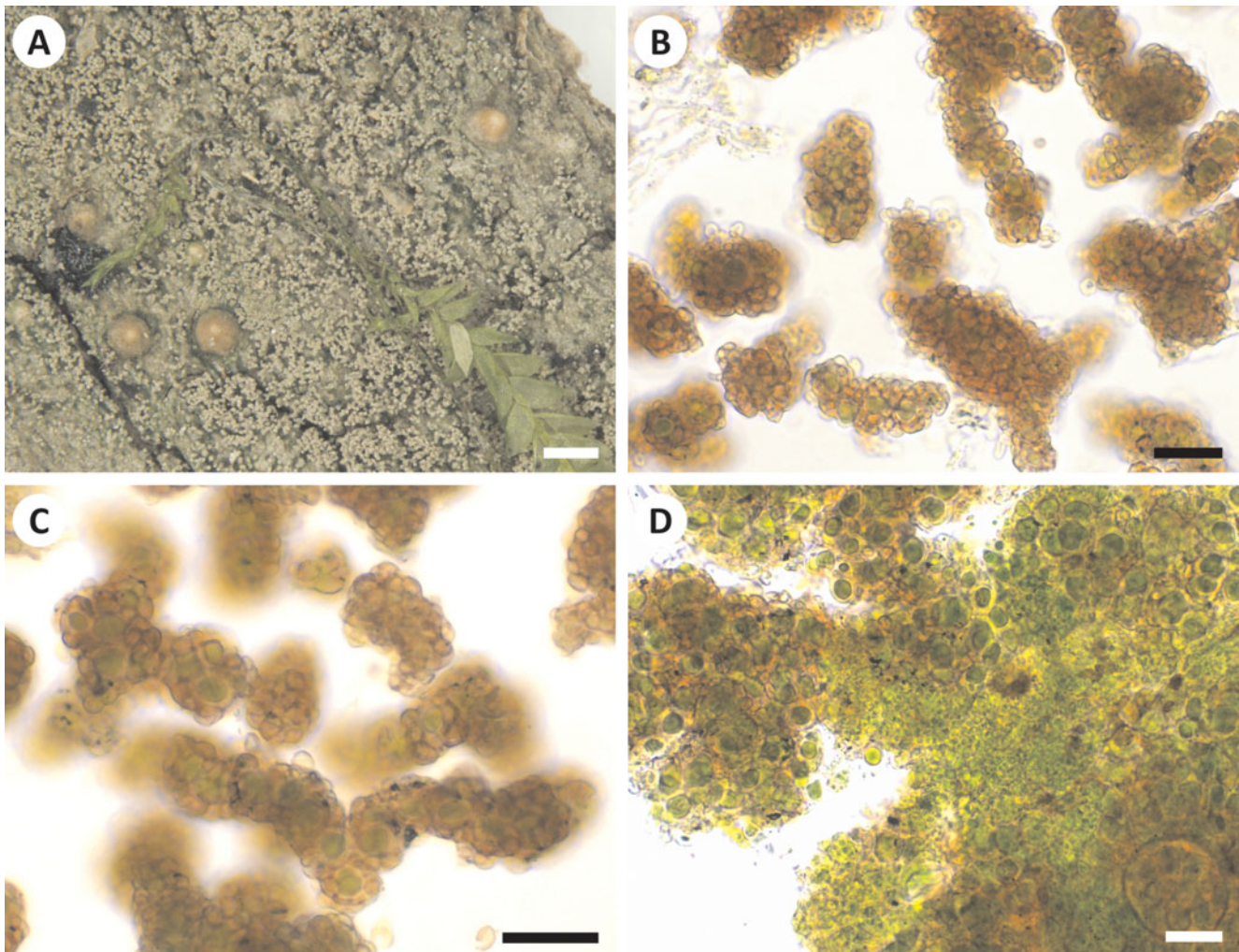


Figure 3. Habitus of *Porina rosei*. A, isidiate thallus with perithecia (PRA-JV22903). B & C, purple pigment in isidia c. 20 hours after KOH application (PRA-ZP29299). D, trentepohlioid photobiont (large cells) and a colony of *Coccomyxa* (small cells) from isidia in KOH (PRA-JV22903). Scales: A = 0.5 mm; B & C = 50 μ m; D = 20 μ m. In colour online.

90 μ m) in *C. luteum* s. lat. A comparison of the 10 isidiate taxa of the genus known so far was recently made (Davydov *et al.* 2021). Until now, only *C. isidiatum* (G. Thor & Vězda) Lücking *et al.* has been reported from the temperate to boreal zone of the Northern Hemisphere. It differs in the bluish grey, locally shiny thallus with concolorous isidia and bigger ascospores of 9–14 \times 3–4 μ m (Thor & Vězda 1984; Davydov *et al.* 2021). *Coenogonium nimisii* is very similar to the New Zealand species *C. fruticulosum* L. Ludw., which differs in several minor anatomical characters (see the diagnosis) and having no observed pycnidia and a wider ecology, including also rotting tussock bases in subalpine grasslands and bryophytes or detritus in montane shrublands (Ludwig 2014).

Enterographa brezhonega Sparrius & Aptroot is occasionally parasitic on *C. nimisii* and rarely on *C. luteum*, and could easily be mistaken as myxomycete fruiting bodies or blobs of *Lepraria*; however, if examined closely the convoluted white lirellae are highly distinctive.

Additional specimens examined. **Great Britain:** England: V.C. 5, South Somerset, Ley Combe, Hawkcombe, SS885.456, *Quercus* high forest derived from coppice on steep slope, on

base-rich streak on downhill side of *Quercus* stub, 2001, N. Sanderson 436 (hb. Sanderson; fertile!); V.C. 11, South Hampshire, Southampton, New Forest National Park, c. 4 km west of Lyndhurst, 50.86893°N, 1.63327°W, 60 m, on mossy bark of *Fagus sylvatica*, 2018, N. Sanderson, A. Acton, M. Powell & J. Vondrák 19844 (PRA); *ibid.*, c. 3 km east of Lyndhurst, 50.86941°N, 1.52586°W, 40 m, on bark of *Quercus*, 2018, N. Sanderson, A. Acton, M. Powell & J. Vondrák 19912 (PRA); Matley Wood, on flushed side of old *Quercus* trunk, 2018, M. Powell 4685 (hb. Powell, hb. Malíček). **Scotland:** V.C. 98, Argyll Main, Grampian Mts, Oban, Invercreran, Glen Creran, old-growth oak-dominated forest with wet places, ESE-facing slope, 56°35'48"N, 5°11'23"W, 40–60 m, on bark of *Quercus robur*, 2018, J. Malíček 12373, A. Acton, Z. Palice, M. Powell & J. Vondrák (hb. Malíček); Ardfern, Eilean Mhic Chrion, old-growth hazel-dominated wood on a ridge and slopes below, 56°10'25"N, 5°32'23"W, alt. 50–60 m, on mossy *Fraxinus*, infected by *Enterographa brezhonega*, 2018, A. Acton, B. Coppins, J. Malíček, Z. Palice 25138, M. Powell & J. Vondrák (PRA).—**Ireland:** V.C. H2, North Kerry, Killarney, Meeting of the Waters, on *Quercus*, infected by *Enterographa brezhonega*, 1996, B. Coppins 25862 (E).

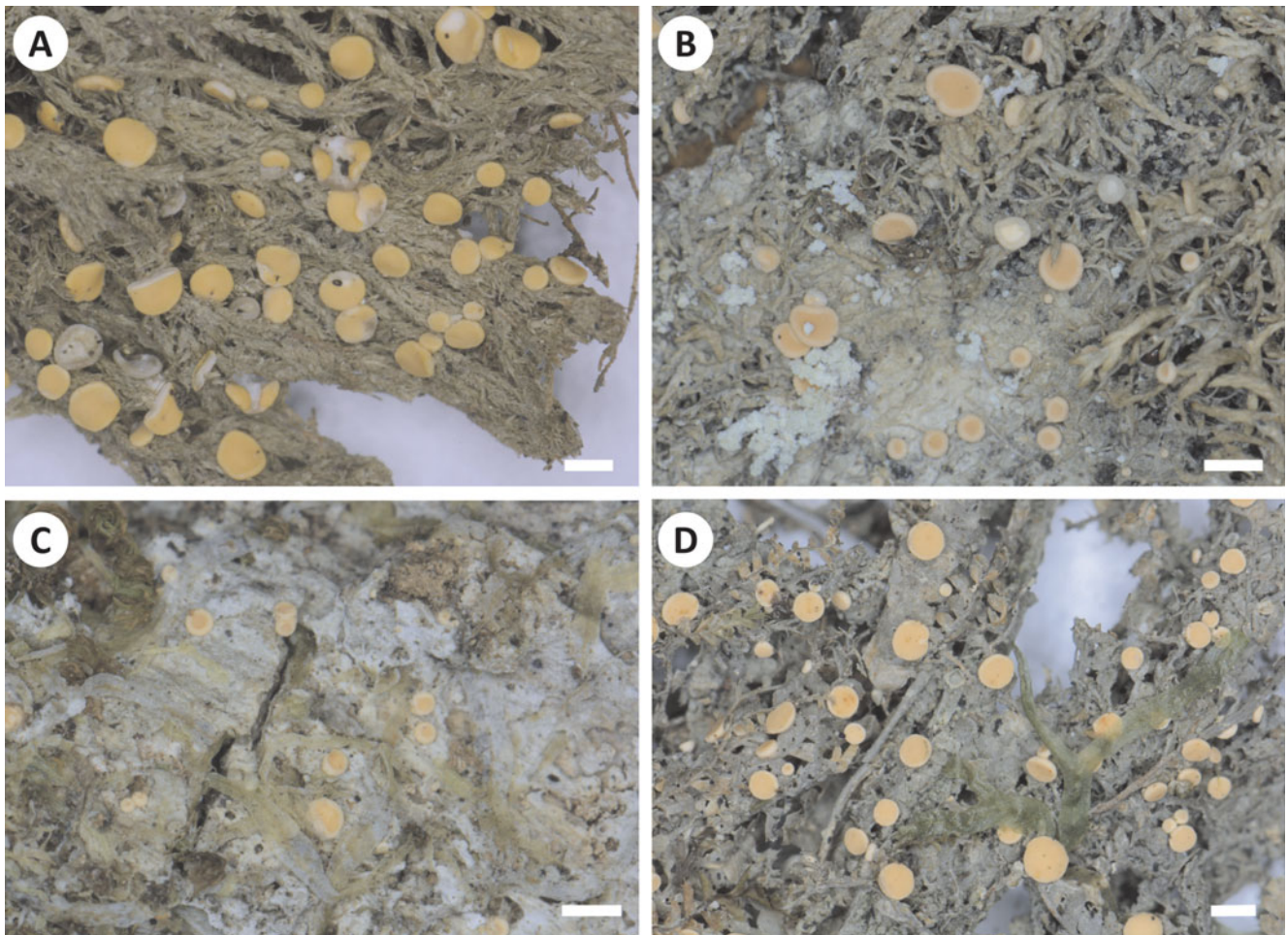


Figure 4. Variability of *Coenogonium luteum* s. lat. (A–C) and *C. luteolum* (D). A, specimen JM 12550. B, JM 10624. C, JM 9141. D, JM 14105. Scales = 1 mm. In colour online.

Specimens of Porina rosei examined. **Czech Republic:** South Bohemian Region: Šumava, Lenora, Zátoňská hora Nature Reserve, old-growth scree forest on S-facing slope of Mt Zátoňská hora, 48°56'27"N, 13°49'52"E, 900 m, on overhanging siliceous rock, 2018, J. Malíček 12126 & Z. Palice (hb. Malíček); Šumava, Prachatice, the valley of Blanice, rock outcrop with a fragment of relic pine forest above the right bank of the rivulet, 48°57'47.2"N, 13°55'21.8"E, 760 m, on shaded overhanging granite rock, 2020, Z. Palice 29299 (PRA).—**France:** Vercors, Gorges de la Vernaison, en amont de Echevis, amont du lieu-dit Grangeage, 400–450 m, taillis avec buis en bord de rivière, 1986, E. Sérusiaux (PRA-V-03285, isotype).—**Great Britain:** England. V.C. 11, South Hampshire, New Forest, Busketts Wood, Great Stubby Hat, 50.896588°N, 001.564699°W, 35 m, fertile on bark of senescent *Fagus sylvatica* in old-growth *Fagus-Ilex* pasture woodland, 2017, N. Sanderson (hb. Sanderson); *ibid.*, Coomy Hat, 50.897609°N, 001.558788°W, 30 m, on bark of ancient *Fagus sylvatica* in old-growth *Fagus-Ilex* pasture woodland, 2020, N. Sanderson (hb. Sanderson).—**Russia:** Adygea: Maykop, Guzeripl, protected area Kavkazskiy zapovednik, 43.98889°N, 40.12507°E, 700 m, on bark of *Carpinus orientalis*, 2016, J. Vondrák 15450 (PRA). **Krasnodar Region:** Adler, Khosta, protected area Tiso-samshitovaya roshcha, 43.53116°N, 39.87683°E, 120 m, on twig of *Buxus sempervirens*, 2019, J. Vondrák 22903

(PRA); *ibid.*, on bark of *Quercus*, 2019, Z. Palice 33401, J. Vondrák 22938 (PRA).

Coenogonium luteum (Dicks.) Kalb & Lücking

(Fig. 4)

For a comparison of phylogenetic relationships, we included seven specimens of *C. luteum* in the analysis. However, we found this to be a polyphyletic species, represented by at least three lineages in this study (Fig. 2). The lectotype (E00455320, coll. J. Dickson 1785) of *C. luteum* comes from the United Kingdom. Sequences cover two distinct clades from Britain, but the specimens differ in the ascospore length: 7.0–9.0(–10.5) versus 9.5–12 µm. We consider the clade represented by three collections (marked s. str.), which contain shorter ascospores, to be identical with the type specimen with ascospores of (6.7–)8–9(–11) µm. The clade represented by the specimen JM10624 corresponds to *C. luteolum*, which is characterized by yellowish apothecia. Kalb & Hafellner (1992) also described differences in the length of ascospores (7–9 µm in *C. luteolum* vs 8–12 µm in *C. luteum*); however, these differences were not observed by us and we regard them as part of intraspecific variability. The remaining two lineages (marked s. lat.) probably represent undescribed or similar tropical


species (see Rivas Plata *et al.* 2006). The identity of the North American *C. luteum* (Ryan 31430), which is strikingly close genetically to *C. nimisii*, is debatable and the possibility that it could be *C. nimisii* cannot be ruled out. The sequenced *C. nimisii* samples come from a geographically limited area (i.e. Great Britain), which may account for the low genetic variability.

Conclusions

In contrast to the high number of described species in the genus *Coenogonium*, the representation of molecular data in the GenBank database is very poor (< 10 species). In this study, we produced 16 mitochondrial SSU and 12 nuclear ITS sequences from five morphologically delimited taxa of *Coenogonium*. Additionally, we produced three mycobiont sequences of mtSSU and three ITS for *Porina rosei*, as well as four ITS and nine *rbcL* sequences of algal partners isolated from both species. The sequences of *C. tavaresianum*, *C. luteolum* and *Porina rosei* are published here for the first time. *Coenogonium tavaresianum* is genetically close to *C. pineti*. *Porina rosei* is published here as new for the Czech Republic.

The newly described species *C. nimisii* forms a well-defined and supported branch within the phylogenetic tree (Fig. 2), but this is in contrast to two other well-known, widely distributed and traditionally defined taxa, *C. luteum* and *C. pineti*. The first mentioned taxon represents at least three distinct species according to molecular data. Similarly, *C. pineti* is genetically variable and consists of at least two distinct but apparently cryptic species. The species complexes of the traditionally circumscribed taxa *C. luteum* and *C. pineti* are deliberately not addressed in detail within this paper as they merit a more comprehensive study.

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References

- Aptroot A and Rodrigues AFF (2005) New lichen records for the Azores, with the report of some tropical species new to Europe. *Cryptogamie, Mycologie* **26**, 273–280.
- Blattner FR (1999) Direct amplification of the entire ITS region from poorly preserved plant material using recombinant PCR. *Biotechniques* **27**, 1180–1186.
- Borgato L, Ertz D, Van Rossum F and Verbeke A (2022) The diversity of lichenized trentepohlioid algal (*Ulvophyceae*) communities is driven by fungal taxonomy and ecological factors. *Journal of Phycology* **58**, 582–602.
- Cannon P, Malíček J, Sanderson N, Benfield B, Coppins B and Simkin J (2021) *Ostropales: Coenogoniaceae*, including the genus *Coenogonium*. *Revisions of British and Irish Lichens* **3**, 1–4.
- Černajová I, Schiefelbein U and Škaloud P (2022) Lichens from the littoral zone host diverse ulvophycean photobionts. *Journal of Phycology* **58**, 267–280.
- Darriba D, Taboada GL, Doallo R and Posada D (2012) jModelTest 2: more models, new heuristics and parallel computing. *Nature Methods* **9**, 772.
- Davydov EA, Yakovchenko LS, Konoreva L, Chesnokov S, Ezhkin A, Galanina IA and Paukov AG (2021) New records of lichens from the Russian Far East. II. Species from forest habitats. *Opuscula Philolichenum* **20**, 54–70.
- Ferencová Z, Rico VJ and Hawksworth DL (2017) Extraction of DNA from lichen-forming and lichenicolous fungi: a low-cost fast protocol using Chelex. *Lichenologist* **49**, 521–525.
- Follmann G (1990) Zur Kenntnis der Flechtenflora und Flechtenvegetation von Madeira und den umliegenden Inseln. I. Chorologisch-soziologischer Abriss. *Zeitschrift Courier Forschungsinstitut Senckenberg* **129**, 91–102.
- Gardes M and Bruns TD (1993) ITS primers with enhanced specificity for basidiomycetes – application to the identification of mycorrhizae and rusts. *Molecular Ecology* **2**, 113–118.
- Grube M, Muggia L, Baloch E, Hametner C and Stocker-Wörgötter E (2017) Symbioses of lichen-forming fungi with trentepohlialean algae. In Grube M, Seckbach J and Muggia L (eds), *Algal and Cyanobacteria Symbioses*. London: World Scientific Publishing Europe, pp. 85–110.
- Guindon S and Gascuel O (2003) A simple, fast, and accurate algorithm to estimate large phylogenies by maximum likelihood. *Systematic Biology* **52**, 696–704.
- Hall TA (1999) BioEdit: a user-friendly biological sequence alignment editor and analysis program for Windows 95/98/NT. *Nucleic Acids Symposium Series* **41**, 95–98.
- Kalb K and Hafellner J (1992) Bemerkenswerte Flechten und lichenicole Pilze von der Insel Madeira. *Herzogia* **9**, 45–102.
- Kantvilas G, Rivas Plata E and Lücking R (2018) The lichen genus *Coenogonium* in Tasmania. *Lichenologist* **50**, 571–582.
- Katoh K and Standley DM (2013) MAFFT multiple sequence alignment software version 7: improvements in performance and usability. *Molecular Biology and Evolution* **30**, 772–780.
- Katoh K, Kuma K, Toh H and Miyata T (2005) MAFFT version 5: improvement in accuracy of multiple sequence alignment. *Nucleic Acids Research* **33**, 511–518.
- Lücking R, Hodkinson BP and Leavitt SD (2017) The 2016 classification of lichenized fungi in the *Ascomycota* and *Basidiomycota* – approaching one thousand genera. *Bryologist* **119**, 361–416.
- Ludwig LL (2014) *Coenogonium fruticulosum*, a new isidiate species from New Zealand. *Australasian Lichenology* **75**, 18–27.
- McCarthy PM and Kantvilas G (2017) A new species of *Porina* (lichenized *Ascomycota*, *Porinaceae*) from Tasmania. *Telopea Journal of Plant Systematics* **20**, 109–113.
- Miadlikowska J, Kauff F, Högnabba F, Oliver JC, Molnár K, Fraker E, Gaya E, Hafellner J, Hofstetter V, Gueidan C, *et al.* (2014) A multigene phylogenetic synthesis for the class *Lecanoromycetes* (*Ascomycota*): 1307 fungi representing 1139 infrageneric taxa, 317 genera and 66 families. *Molecular Phylogenetics and Evolution* **79**, 132–168.
- Nelsen MP, Plata ER, Andrew CJ, Lücking R and Lumbsch HT (2011) Phylogenetic diversity of trentepohlialean algae associated with lichen-forming fungi. *Journal of Phycology* **47**, 282–290.
- Orange A, James PW and White FJ (2010) *Microchemical Methods for the Identification of Lichens*. London: British Lichen Society.
- Orange A, Palice Z and Klepsland J (2020) A new isidiate saxicolous species of *Porina* (*Ascomycota*, *Ostropales*, *Porinaceae*). *Lichenologist* **52**, 267–277.
- Rivas Plata E, Lücking R, Aptroot A, Sipman HJM, Chaves JL, Umaña L and Lizano D (2006) A first assessment of the Ticolichen biodiversity inventory in Costa Rica: the genus *Coenogonium* (*Ostropales: Coenogoniaceae*), with a world-wide key and checklist and a phenotype-based cladistic analysis. *Fungal Diversity* **23**, 255–321.
- Rodrigues AFF and Aptroot A (2005) New data and corrections to the list of lichens and lichenicolous fungi from the Azores. In Borges PAV, Cunha R, Gabriel R, Martins AF, Silva L and Vieira V (eds), *A List of the Terrestrial Fauna and Flora from the Azores*. Horta: Direcção Regional do Ambiente and Universidade dos Açores, pp. 231–247.
- Ronquist F and Huelsenbeck JP (2003) MrBayes 3: Bayesian phylogenetic inference under mixed models. *Bioinformatics* **19**, 1572–1574.
- Ronquist F, Teslenko M, van der Mark P, Ayres DL, Darling A, Höhna S, Larget B, Liu L, Suchard MA and Huelsenbeck JP (2012) MrBayes 3.2: efficient Bayesian phylogenetic inference and model choice across a large model space. *Systematic Biology* **61**, 539–542.
- Sérusiaux E (1991) *Porina rosei* sp. nov., une espèce nouvelle d'Europe occidentale. *Cryptogamie, Bryologie et Lichénologie* **12**, 31–39.

- Škaloud P and Rindi F** (2013) Ecological differentiation of cryptic species within an asexual protist morphospecies: a case study of filamentous green alga *Klebsormidium* (Streptophyta). *Journal of Eukaryotic Microbiology* **60**, 350–362.
- Škaloud P, Rindi F, Boedeker C and Leliaert F** (2018) *Freshwater Flora of Central Europe, Vol 13: Chlorophyta: Ulvophyceae*. Berlin, Heidelberg: Springer.
- Swofford DL** (2003) *PAUP*. Phylogenetic Analysis Using Parsimony (*and Other Methods), Version 4*. Sunderland, Massachusetts: Sinauer Associates.
- Tavares CN** (1952) Contributions to the lichen flora of Macaronesia. I. Lichens from Madeira. *Portugaliae Acta Biologica* **3**, 308–391.
- Thor G and Vězda A** (1984) Einige neue oder bemerkenswerte Flechten mit gyalectoiden Apothecien von Nord-Indien und Nepal. *Folia Geobotanica et Phytotaxonomica* **19**, 71–82.
- White TJ, Bruns TD, Lee S and Taylor J** (1990) Amplification and direct sequencing of fungal ribosomal RNA genes for phylogenetics. In Innis MA, Gelfand DH, Sninsky JJ and White TJ (eds), *PCR Protocols: a Guide to Methods and Applications*. San Diego: Academic Press, pp. 315–322.
- Zhu H, Zhao Z, Xia S, Hu Z and Liu G** (2015) Morphological examination and phylogenetic analyses of *Phycopeltis* spp. (*Trentepohliales, Ulvophyceae*) from tropical China. *PLoS ONE* **10**, e0114936.
- Zoller S, Scheidegger C and Sperisen C** (1999) PCR primers for the amplification of mitochondrial small subunit ribosomal DNA of lichen-forming ascomycetes. *Lichenologist* **31**, 511–516.
- Zwickl DJ** (2006) *Genetic algorithm approaches for the phylogenetic analysis of large biological sequence datasets under the maximum likelihood criterion*. Ph.D. thesis, University of Texas at Austin.