

Phylogeny of the European *Collema* species (Peltigerales, Lecanoromycetes)

Alica Košuthová¹, Fredrik Jonsson², Ulrika Nordin², Mats Wedin¹

¹ Department of Botany, Swedish Museum of Natural History, P.O. Box 50007, SE-104 05 Stockholm, Sweden

² Alsens-Ede 227, SE-835 96 Trångsviken, Sweden

Corresponding author: Alica Košuthová (kosuthova.alica@gmail.com)

Abstract

The phylogenetic relationships and morphological diversity within European *Collema* s. str. species were investigated. A total of 104 new sequences (four molecular markers; mtSSU, b-tub, MCM7, and RPB2 genes) from 28 specimens were generated, and analysed and used for multi-locus phylogenetic analyses. Our results suggest that *Collema* is only monophyletic if *Collema glebulentum* is considered part of *Leptogium* s. str. where it originally was described. This is supported by its paraplectenchymatous thallus. Degelius' informal *Collema* "Flaccidum"- and "Nigrescens"-groups are not natural, as the "Flaccidum"-group is nested within the "Nigrescens"-group. Based on our findings, seven currently accepted *Collema* occur in Europe: *C. flaccidum*, *C. subflaccidum*, *C. curtisporum*, *C. furfuraceum*, *C. nigrescens*, *C. rysssoleum*, and *C. subnigrescens*. *Collema furfuraceum* is further non-monophyletic, suggesting a need for taxonomic revision.

Key words: Ascomycota, Collemataceae, cyanolichens, integrative taxonomy, systematics

Introduction

The generic delimitation of Collemataceae s. str. was for a long time unnatural, relying primarily on a single trait: the presence of a cellular cortex in *Leptogium* or its absence in *Collema* (Degelius 1954, 1974; Jørgensen 2007). Degelius (1954) had, however, already questioned whether *Collema* and *Leptogium* represented natural, monophyletic groupings. Subsequent molecular studies (Wiklund and Wedin 2003; Miadlikowska and Lutzoni 2004; Miadlikowska et al. 2014) supported this, revealing that the gelatinous genera with simple spores, traditionally assigned to Collemataceae, actually belonged to other families (Wedin et al. 2009; Otálora et al. 2010; Ekman et al. 2014; Weerakoon et al. 2018). It was not until the comprehensive analysis by Otálora et al. (2013a, 2013b) that a revised generic classification of Collemataceae s. str. was proposed. Their work confirmed the non-monophyletic nature of *Collema* and *Leptogium*, proposing the recognition of ten distinct monophyletic groups as separate genera. This revision retained the names *Collema* and *Leptogium* in restricted senses and reintroduced six older generic names (*Blennothallia*, *Enchylium*, *Lathagrium*, *Pseudoleptogium*, *Rostania*, and *Scytinium*), while also establishing two new genera, *Callome* and *Paracollema*.



Academic editor: Thorsten Lumbsch

Received: 17 December 2024

Accepted: 19 January 2025

Published: 18 March 2025

Citation: Košuthová A, Jonsson F, Nordin U, Wedin M (2025) Phylogeny of the European *Collema* species (Peltigerales, Lecanoromycetes). MycoKeys 115: 209–220. <https://doi.org/10.3897/mycokeys.115.144718>

Copyright: © Alica Košuthová et al.

This is an open access article distributed under terms of the Creative Commons Attribution License (Attribution 4.0 International – CC BY 4.0).

The European species of *Collema* in the sense of Otálora et al. (2013b), the focus of the present study, correspond to Degelius' "Flaccidum" and "Nigrescens"-groups (Degelius 1954). These groups are relatively small, with three species currently recognized in the "Flaccidum"-group and five in the "Nigrescens"-group. The two groups are closely related, sharing several characteristics, including their occurrence in sheltered, humid environments, and the two groups contain both corticolous and saxicolous species. Species within these groups are large in size, typically measuring about 2.0–8.0 cm in diameter, though some individuals may reach up to 15 cm. Their thalli are thin to moderately thin (50–150 µm), lacking both a paraplectenchymatous medulla, and a true cortex. The lobes are generally broad and rounded, measuring approximately 5.0–12.0 mm in diameter. The spores of both groups are ranging from 25–35 × 6–7 µm to 40–90 × 3–6 µm, and are narrowly ellipsoid to fusiform with transverse septation. Spores in the "Nigrescens"-group show greater polymorphism than those in the "Flaccidum"-group. A distinguishing feature between the two groups is that species in the "Flaccidum"-group lack pustules and ridges, whereas those are present in species of the "Nigrescens"-group. Both groups are distributed across Europe, with their distribution range and selected morphological features summarized in Table 1. Degelius (1954, 1974) accepted three European species in the "Flaccidum"-group: *Collema flaccidum* (Ach.) Ach., *C. glebulentum* (Cromb.) Degel., and *C. subflaccidum* Degel. previously known as *C. subfurvum* (Müll. Arg.) Degel. The five European species in the "Nigrescens"-groups include *C. curtisporum* Degel., *C. furfuraceum* (Arn.) DR. em. Degel., *C. nigrescens* (Huds.) DC., *C. rysssoleum* (Tuck.) Schneid., and *C. subnigrescens* Degel.

Otálora et al. (2013a, 2013b) included only four species (*C. curtisporum*, *C. flaccidum*, *C. furfuraceum* and *C. nigrescens*) of the European *Collema* s. str. in their phylogenies, with Miadlikowska et al. (2014) also including *C. subnigrescens*. As a result, the taxonomic positions of five out of eight species within this group have been tested using molecular methods, while the remaining three species have not yet been examined. As there is a substantial variation in shape and size of the thallus, lobes, apothecia, ascospores, and isidia among them, and as several former *Collema* species have been shown to belong elsewhere, the delimitation of the genus needs investigation. Here, we will test the current delimitation of *Collema* in Europe and propose a phylogenetic hypothesis of known species. Finally, we will note and comment on any indication of species non-monophyly in this genus.

Material and methods

Specimen selection and morphological observations

We sampled 28 specimens, representing species of the European "Flaccidum" and "Nigrescens"-groups from Scandinavia and including some extra-Scandinavian material for comparison. The collections resulting from our own recent fieldwork are deposited in the herbarium S, with additional material studied from GZU, hb. Malíček, TBS, UPS and S. Collections and the sequences used are summarized in Table 2. Herbarium acronyms follow Thiers (2018).

Table 1. Characteristic morphological features of European *Collema* species summarised according to Degelius (1954, 1974). Characters unique for the species in the “*Flaccidum*” and “*Nigrescens*” -groups are in bold.

| Group | Morphology | | | | | | | | | | | | | | |
|-------------------|-------------------------------|---------------------|------------------------------|-------------------------------------|---------------|---|-----------------------|--------------------------|-----------------------|---|---|---|-----------------|-------------------------|---------------------------------------|
| | Species (<i>Collema</i>) | Pustules/ ridges | Thallus thickness (µm) | Thallus colour | Lobes (cm) | Isidia | Isidia width (mm) | Isidia length (mm) | Apothecium (mm) | Apothecium disc | Spore measurements (µm) | Spore shape | Spore septas | Habitat | Distribution |
| <i>Flaccidum</i> | <i>flaccidum</i> | N | 70–170 | olive-green-blackish | 0.5–1.5(3) | squamiform (juv. globular) | 0.2–0.5 | min. 0.2–0.5 | rare, to 1.5 | pale red to dark red, sometimes pruinose | (20)26–34(45) x 6–6.5(8.5) | fusiform with acute ends | 4 to 6 | saxi- and corti-colours | wide, suboceanic |
| | <i>glebulentum</i> | N | (50)65–135(200) | light to dark olive-green-blackish | to 1 | teretiform, coralloid, capitate | 0.1 | 1.5 | N | N | N | N | N | saxicolous | arctic-alpine |
| | <i>subflaccidum</i> | N | 80–130 | olive-green-blackish | 0.5–1.5(3) | globular , old teretiform & coralloid | 0.05–0.1 | min. 0.3 | rare, to 1.5 | pale red to dark red, epruinose | 42–60(65) x (3)4.5–6.5(10.5) | broadly to narrowly fusiform, acicular | (4)6–8 | corti- and saxi-colours | oceanic |
| <i>Nigrescens</i> | <i>curtisporum</i> | Y | 65–106 | dark olive-green-blackish | 0.5–1 | N | N | N | 0.5–1.5 | pale-red-dark red-brown-blackish, epruinose | (18)26–34(40) x 3–4.5 | bacillar, straight or curved, obtuse ends | 4(5–6) | corticolous | boreal |
| | <i>furfuraceum</i> | Y | (50)60–105 | dark olive-green-blackish | 0.5–1 | teretiform, coralloid, branched (juv. globular) | 0.05–0.15(0.2) | 0.3 | very rare, 0.5–1.5 | pale-red-dark red-brown-blackish, epruinose | 40–80 x 3–6(8.5) | bacillar-narrowly fusiform with acute end, one end thicker and usually curved | 5 to 6 | corticolous | wide, suboceanic |
| | <i>nigrescens</i> | Y | (60)90–150 | dark olive-green, brownish-blackish | 0.5–1 | globular | 0.2 | 0.2 | often, 0.6–1 | pale-red-dark red-brown-blackish, epruinose | 50–90(112) x 3–4.5 | acicular to bacillar | (5)6–13 | corticolous | wide, suboceanic |
| <i>Nigrescens</i> | <i>ryssoleum</i> | Y | 50–85 | dark olive-green-blackish | 0.5–1.5 | N | N | N | 0.6–1(1.5) | dark red-blackish, epruinose | (22)26–40(47) x (4.5)5–8.5(10.5) | ellipsoid or fusiform, broader in middle , with acute ends | 4 to 6 | saxicolous | temperate - mediterranean zone |
| | <i>subnigrescens</i> | Y | 60–100 | dark olive-green-blackish | 0.5–1.5 | N | N | N | often, 1–1.5(2) | pale-red-dark red-brown-blackish, epruinose | (34)40–75 x 6–6.5(7) | narrowly fusiform with acute end, one end thicker and usually curved | 5 to 6 | corticolous | wide, suboceanic |

Table 2. Sequences utilized in this study. Newly produced sequences in bold with herbarium vouchers of the specimens given, remaining sequences downloaded from GenBank. For specimens of *Collema glebulentum*, origin of both, state and provinces are given.

| Specimen | DNA-voucher | Year | Locality & Herbarium voucher | mtSSU | BT | MCM7 | RPB2 5-7 | RPB2 7-11 |
|-------------------------------|-------------|------|--|-----------------|-----------------|-----------------|-----------------|-----------------|
| <i>Leptogium byssinum</i> | | | Norway: Westberg (S) | KT240180 | | KT240183 | | |
| <i>Leptogium terrenum</i> | | | Portugal: van den Boom 41781 (hb. van den Boom) | KT240181 | | KT240184 | | |
| <i>Collema glebulentum</i> | AL561 | 2015 | Sweden, Pite Lappmark (Arjeplog): Westberg 15-254 (S-F277955) | PQ932211 | PV021123 | | PV021148 | |
| <i>Collema glebulentum</i> | AL560 | 1988 | Sweden, Åsele Lappmark (Vilhelmina): Thor 7711 (S-L49768) | PQ932210 | | | | |
| <i>Collema glebulentum</i> | AL674 | 2018 | Sweden, Värmland (Säffle-Lurö): Košuthová et al. 401 (S-F492346) | PQ932212 | PV021124 | | PV021149 | |
| <i>Collema glebulentum</i> | AL366 | 2017 | Sweden, Närke (Örebro): Berglund (S-F492347) | PQ932209 | PV021122 | | | |
| <i>Leptogium azureum</i> | | | Chile: Cornejo 26507 (MA) | JX992939 | KC119021 | JX993002 | | |
| <i>Leptogium denticulatum</i> | | 2010 | Argentina: Wedin 8690 (S-F332474) | JX992947 | KC119025 | JX993012 | PV021147 | |
| <i>Collema subconveniens</i> | | 2010 | New Zealand: Wedin 9225 (S-F335747) | JX992937 | KC119019 | JX992996 | PV021150 | |
| <i>Collema leptaleum</i> | | | Argentina: Wedin 8822 (S) | JX992928 | KC119038 | JX992986 | | |
| <i>Collema flaccidum</i> | AL540 | 2016 | Sweden: Westberg et al. 244 (UPS-L872188) | PQ932216 | PV021129 | PV021173 | PV021154 | |
| <i>Collema flaccidum</i> | AL496 | 2017 | Sweden: Odelvik 17-523 (S-F317108) | PQ932215 | PV021128 | PV021172 | PV021153 | PV021189 |
| <i>Collema flaccidum</i> | AL531_AL494 | 2018 | Slovakia: Košuthová et al. 601 (S-F492348) | PQ932217 | PV021130 | PV021174 | PV021155 | PV021190 |
| <i>Collema subflaccidum</i> | AL649 | 2016 | Russia: Malíček et al. 10619 (S-F492349 & dupl. herb. Malíček) | PQ932214 | PV021132 | | PV021152 | |
| <i>Collema subflaccidum</i> | AL495 | 2018 | Norway: Aptroot 76306 (S-F492350) | PQ932213 | PV021131 | | PV021151 | PV021188 |
| <i>Collema nigrescens</i> | AL511 | 2018 | Spain: Westberg (UPS-L934034) | PQ932220 | PV021127 | | PV021158 | |
| <i>Collema nigrescens</i> | AL493 | 2018 | Slovakia: Košuthová et al. 600 (S-F492351) | PQ932219 | PV021126 | | PV021157 | |
| <i>Collema nigrescens</i> | AL603 | 2018 | Sweden: Košuthová & Arvidsson 571 (S-F492352) | PQ932218 | PV021125 | PV021175 | PV021156 | |
| <i>Collema curtisporum</i> | AL411 | 2017 | Sweden: Jonsson & U.Nordin FU6546 (S-F492353) | PQ932221 | PV021133 | | PV021159 | PV021191 |
| <i>Collema curtisporum</i> | AL568 | 1994 | Sweden: Hermansson 4603 (UPS-L111603) | PQ932222 | | PV021176 | PV021160 | |
| <i>Collema furfuraceum</i> | AL668_AL721 | 2002 | Sweden: Jonsson 2254 (S-F492354) | PQ932223 | | PV021177 | PV021161 | PV021192 |
| <i>Collema furfuraceum</i> | AL640_AL720 | 1998 | Sweden: Bergsten (S-F492355) | PQ932224 | | PV021178 | PV021162 | PV021193 |
| <i>Collema rysssoleum</i> | AL518 | 2001 | Italy: Trietach (TSB-35166) | PQ932231 | PV021135 | PV021184 | PV021168 | PV021196 |
| <i>Collema rysssoleum</i> | AL534_AL513 | 1974 | Spain: Tibell 5610 (UPS-L933969) | PQ932232 | PV021136 | | | |
| <i>Collema rysssoleum</i> | AL566 | 1994 | Madeira: Nordin 3524 (UPS-L178905) | PQ932233 | PV021134 | | | |
| <i>Collema subnigrescens</i> | AL407 | 2017 | Sweden: Jonsson & U.Nordin FU6531 (S-F492356) | PQ932227 | PV021140 | PV021180 | PV021165 | |
| <i>Collema subnigrescens</i> | AL500 | 2006 | Greece: Spribille 19637 (GZU66-201) | PQ932226 | PV021141 | | PV021164 | |
| <i>Collema subnigrescens</i> | AL570 | 2004 | Estonia: Odelvik 0485 (S-F57732) | PQ932228 | PV021142 | PV021181 | PV021166 | PV021195 |
| <i>Collema subnigrescens</i> | AL344 | 2017 | Sweden: Berglund (S-F492357) | PQ932225 | PV021139 | PV021179 | PV021163 | PV021194 |
| <i>Collema furfuraceum</i> | AL644 | 2018 | Scotland: Malíček 12545 (herb. Malíček) | PQ932230 | PV021143 | PV021183 | | |
| <i>Collema furfuraceum</i> | AL666 | 2012 | Italy: Jonsson A2 (S-F492358) | PQ932229 | | PV021182 | PV021167 | |
| <i>Collema furfuraceum</i> | AL712 | 2022 | Sweden: Westberg (UPS-L1049804) | PQ932235 | PV021138 | PV021186 | PV021170 | |
| <i>Collema furfuraceum</i> | AL536 | 2018 | Spain: Westberg (UPS-L934040) | PQ932234 | PV021137 | PV021185 | PV021169 | PV021197 |
| <i>Collema furfuraceum</i> | AL665 | 2012 | France: Jonsson A3 (S-F492359) | PQ932236 | PV021144 | PV021187 | PV021171 | |
| <i>Paracollema italicum</i> 3 | | | Croatia: Nordin 2763 (UPS) | JX992926 | | JX992985 | | |
| <i>Paracollema italicum</i> 1 | | | Croatia: Nordin 2708 (UPS) | JX992925 | KC119015 | JX992984 | | |
| <i>Callome multipartita</i> 1 | | | Norway: Haugan 7015 (O) | GQ259019 | | | | |
| <i>Callome multipartita</i> 2 | AL419 | 2009 | Austria: Hafellner 74818 (GZU-18-2009) | MK445271 | MK451935 | | PV021146 | |
| <i>Enchylium bachmanianum</i> | MWE111 | 1985 | Sweden: Nordin 1521 (UPS-L133627) | JX992914 | MK451936 | JX992974 | PV021145 | |

Anatomical features were studied using a light microscope on thin sections cut with a razor blade or squash preparations mounted in water. Measurements of mature spores were taken outside of the asci under $\times 1000$ magnification using oil immersion with a precision of 0.5 μm , or from calibrated digital

photographs using NIS-Elements (Nikon, Japan) with a precision of 0.1 μm . Spore measurements are presented in the format: (minimum value observed–) range including 80% of the observed values (–maximum value observed), with the mean of all observed values in the center and italicized (Table 1). Full lists of specimens examined in this study with DNA voucher codes and GenBank Accession numbers for newly generated sequences are given in Table 2.

DNA extraction, amplification and sequencing

Two apothecia with surrounding thalline parts, or a thallus fragment in the case of sterile samples, were selected for extraction. We extracted total DNA using the Plant DNA Mini Kit (Qiagen, Hilden, Germany) following the manufacturer's instructions except in order to increase the concentration of DNA, we used half the amount of Elution buffer in the last step. We amplified one mitochondrial ribosomal and three nuclear protein-coding genes. Approximately 0.8 kb of the small subunit of the mitochondrial rDNA (mtSSU) was amplified using the primers mrSSU1 and mrSSU3R (Zoller et al. 1999). Approximately 0.6 kb of Beta-tubulin (b-tub) was amplified and sequenced using the primers Bt3-LM5 and Bt10-LM3 (Myllys et al. 2001) and BetaCollF and BetaCollR (Otálora et al. 2013a). The PCR primers were used in the following combinations: Bt3-LM5 and Bt10-LM3, BetaCollF and BetaCollR, Bt3-LM5 and BetaCollR, BetaCollF and Bt10-LM3 (the best working combination), BetaCollF and BetaColl-intR, and BetaColl-intF2 and Bt10LM3. About 0.6 kb of mini-chromosome maintenance complex component 7 DNA replication licensing factor (MCM7) was amplified and sequenced using the primers Mcm7-709for and Mcm7-1348rev (Schmitt et al. 2009). The locus RNA polymerase II protein coding gene (RPB2, ~2 kb) was amplified as two parts: ~ 1 kb for RPB2 5–7 and ~ 1 kb for RPB2 7–11. Amplification was performed using the primers fRPB2-5F, fRPB2-7cF, fRPB2-7cR and fRPB2-11aR (Liu et al. 1999), which were also used as sequencing primers. PCR amplifications were performed using Illustra™ Hot Start PCR beads, according to manufacturer's instructions. PCR reactions were performed using the same settings as in previous studies (Košuthová et al. 2016, 2019; Otálora et al. 2013a).

Sequence editing, alignment and phylogenetic analyses

The generated sequences were assembled and edited using Geneious version R8 (<http://www.geneious.com>, Kearse et al. 2012). All edited sequences underwent initial identity verification through BLAST searches (Zhang et al. 2000). The alignment of these sequences was performed using the MUSCLE algorithm (Edgar 2004) in AliView 1.09 (Larsson 2014). An intron in the b-tub was excluded. Indels in the mtSSU were not deleted, allowing smaller gap positions within the final block. The five genetic regions defined above were analysed separately using Maximum Likelihood (ML). As no significant incongruence was detected, the alignments were concatenated. The final alignment has been deposited in TREEBASE (<http://www.treebase.org>) with accession number (<http://purl.org/phylo/treebase/phyloids/study/TB:S31975>). After concatenation, we inferred phylogenetic relationships using ML with the same settings used as in the individual gene analyses utilizing RAXML. Likelihood and ML bootstrapping were executed through RAXML 8 (Stamatakis 2014) implementing a general

time reversible (GTR) model of nucleotide substitution with gamma distributed rate heterogeneity (GTRGAMMA). One thousand bootstrap (BS) replicates were completed using the non-parametric BS algorithm of RAxML-HPC v.8 on XSEDE using the CIPRES Web Portal (Miller et al. 2010).

For the final concatenated dataset, we included 131 nucleotide sequences of mtSSU, b-tub, MCM7, and the two parts of RPB2 (5–7 and 7–11). This dataset encompassed 4140 nucleotide positions (843 bp for mtSSU, 708 bp for b-tub, 597 bp for MCM7, and 1074 bp for RPB2 5–7 and 918 bp for RPB2 7–11) with 39 terminals. It incorporates representatives from selected genera within the Collemataceae, in conjunction with additional data sourced from GenBank (utilized in Otálora et al 2013a, 2013b; Košuthová et al. 2019), with *Enchylium bachmanianum* as the outgroup. None of the sequences of the “*Nigrescens*” and “*Flaccidum*”-groups previously used in phylogenetic analyses and uploaded to GenBank (Otálora et al. 2010, 2013a, 2013b; Miadlikowska et al. 2014), were included in this study due to concerns about specimen misidentification. Sequences used in this study are summarized in Fig. 1, Table 2.

Results and discussion

Dataset and phylogenetic analyses

We generated 104 new sequences (Table 2), comprising 28 mtSSU, 23 b-tub, 16 MCM7, 27 RPB2. Our analyses resulted in a topology (Fig. 1) very similar to the findings of Otálora et al. (2013a, 2013b).

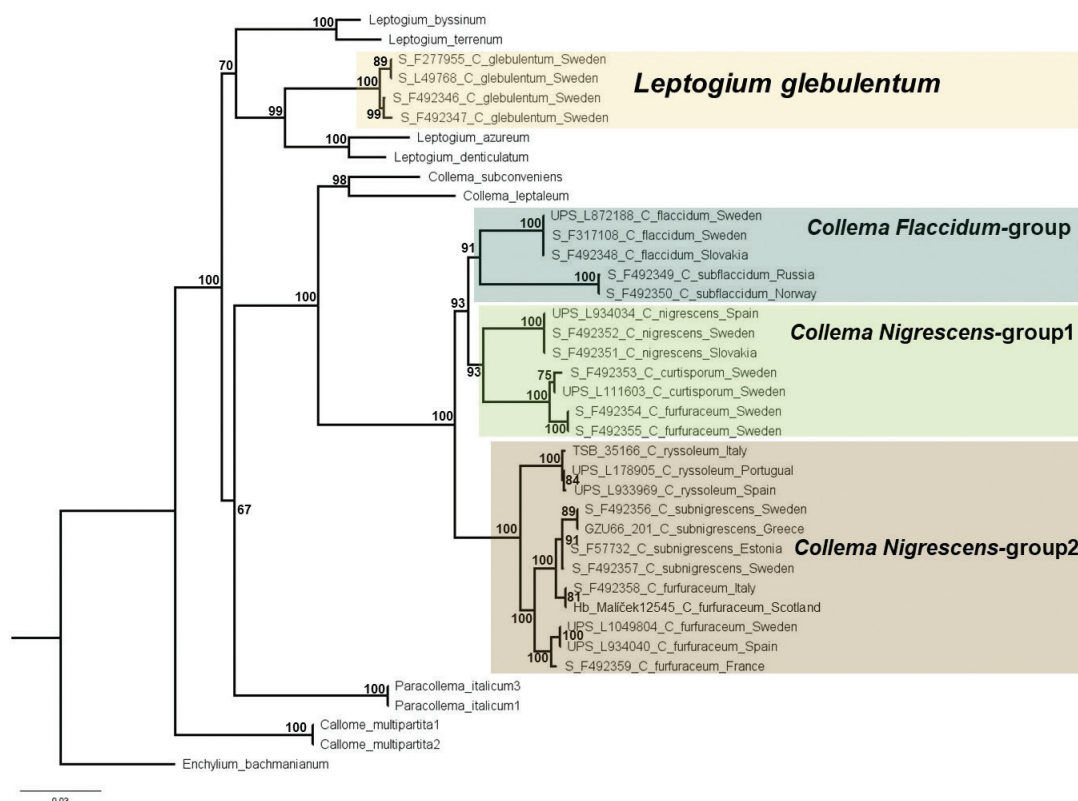


Figure 1. The most likely tree from RAxML analysis based on 4140 aligned characters of mtSSU, b-tub, MCM7 and RPB2 5–7 and 7–11 from 39 specimens. Support values (Likelihood BS) given when BS ≥ 60%. The scale bars indicate 0.03 substitution.

Our phylogenetic analysis reveals that European *Collema* should be treated as including *C. curtisporum*, *C. flaccidum*, *C. furfuraceum*, *C. nigrescens*, *C. rysssoleum*, *C. subflaccidum* and *C. subnigrescens*, but excluding *C. glebulentum* as this species clearly groups within *Leptogium* together with *Leptogium azureum*, the conserved type species of *Leptogium* (Fig. 1). The groups within *Collema* as informally circumscribed by Degelius (1954), are not supported by our phylogeny. The “*Flaccidum*”-group, consisting of *C. flaccidum* and *C. subflaccidum*, is nested within the “*Nigrescens*”-group. This indicates that these two informal groups are not useful for a phylogenetically based classification (Fig. 1). *Collema furfuraceum* is further non-monophyletic, suggesting the need for further taxonomic revision.

Morphological analyses

Degelius (1954) differentiated his informal groups in European *Collema* based on morphological characteristics, noting that species within the “*Flaccidum*”-group are all isidiate but not pustulate (Table 1, Fig. 2A–F). Isidia in *Collema flaccidum* are typically numerous and squamiform (flattened) when fully developed, rounded, and about 0.2–0.5 mm broad or larger. Larger isidia are often crenate or lobulate, with occasional teretiform isidia mixed in (Fig. 2B). In contrast, *C. subflaccidum* has smaller, globular isidia approximately 0.05–0.1 mm in diameter that become slightly teretiform as they mature, reaching at least 0.3 mm in length, with simple or slightly branched forms (Fig. 2C, D). *Collema glebulentum* (Fig. 2E, F) has, sometimes together with a primitive pseudocortex, a distinct typical pseudocortex, often developed on the lower surface, especially on smaller lobes. When a typical pseudocortex is present, it is composed of several cell layers where the cells can be quite large, reaching up to 15 µm in diameter (Fig. 3A). Parts of the thallus may have an entirely paraplectenchymatous structure. This characteristic, noted by Degelius (1954), actually supports the original classification of *C. glebulentum* in *Leptogium* (Fig. 3B). In areas where the thallus is not entirely paraplectenchymatous, its structure resembles that of other species in the “*Flaccidum*” and “*Nigrescens*”-groups. The entire thallus in species from these groups is composed of hyphae that are either loosely or compactly interwoven, or arranged distinctly perpendicular to the upper and lower cortices throughout the thallus (Fig. 3C).

Species within the “*Nigrescens*”-group are characterized by their pustulate and ridged thalli (Fig. 2G–P) and can be divided into non-isidiate species and those that produce isidia. Among the non-isidiate species, *Collema rysssoleum* (Fig. 2G, H) is distinct due to its spore morphology. The spores are short, measuring up to 40 µm in length, similar to those in *C. curtisporum*. However, unlike others in the group, *C. rysssoleum* spores are relatively wide (up to 8.5 µm) with acute ends (Table 1). Additionally, this species exhibits a unique Mediterranean distribution in the temperate zone and is adapted to a saxicolous habitat (Fig. 3D). *Collema subnigrescens* is characterized by an up to 20 cm large thallus (Fig. 2I) and by its narrowly fusiform to irregularly clavate spores with twisted, acute ends (5-celled, occasionally up to 12 cells; Fig. 3E). It closely resembles *C. curtisporum*, which, however, has a smaller, up to 4 cm wide thallus (Fig. 2J) and shorter, mostly 3-celled bacillariform spores with obtuse ends and a typical “curved-acute shape” (Fig. 3F). Spores in *C. nigrescens* differ from the other species in being acicular to bacillariform, longer and thinner than those in *C. subnigrescens* (usually around 60 µm long and less than 5 µm wide), 6–13 celled (Fig. 3G).

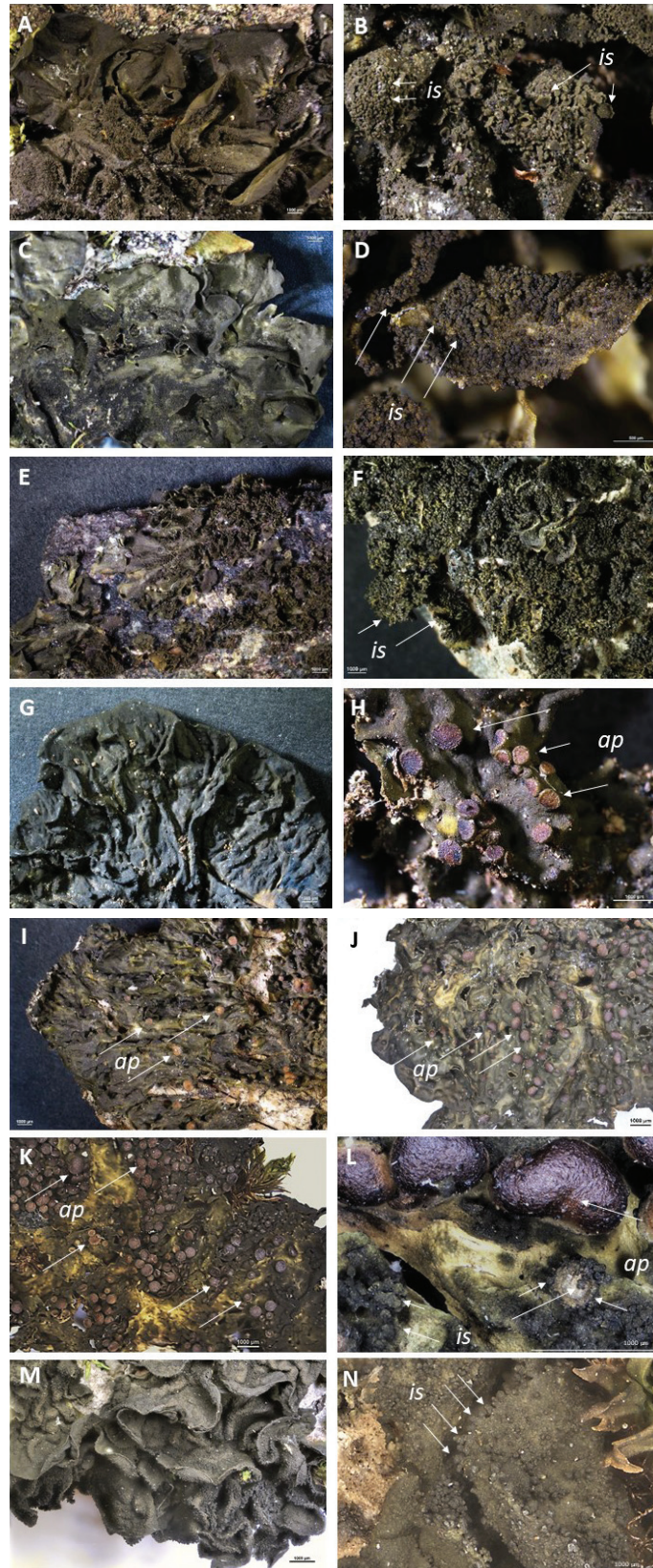


Figure 2. Thallus habitus and isidia of **A** *Collema flaccidum* (UPS-L872188) **B** *C. flaccidum* (UPS-L872188) **C** *C. subflaccidum* (S-F492349) **D** *C. subflaccidum* (S-F492350) **E** *Leptogium glebulentum* (S-L49768) **F** *L. glebulentum* – dwarf form (S-L49768) **G** *C. rysssoleum* (UPS-L933969) **H** *C. rysssoleum* (UPS-L933969) **I** *C. subnigrescens* (S-F57732) **J** *C. curtisporum* (UPS-L111603) **K** *C. nigrescens* (UPS-L934034) **L** *C. nigrescens* (UPS-L934034) **M** *C. furfuraceum* (UPS-L934040) **N** *C. furfuraceum* (UPS-L934040). **is** = isidia, **ap** = apothecium. Scale bar: **A–C, E–N**: 1 mm, **D**: 0.5 mm.

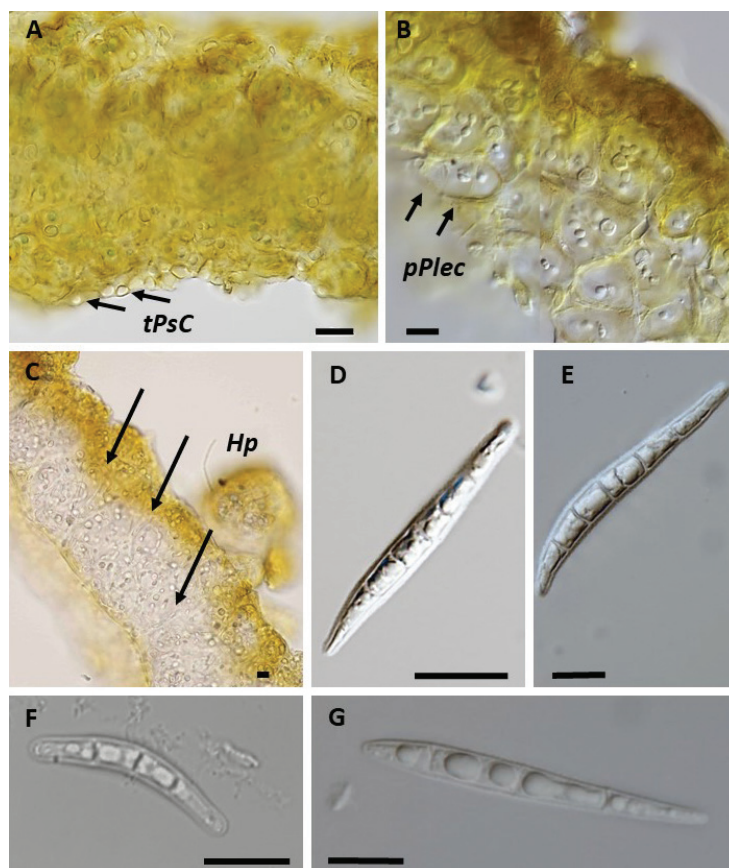


Figure 3. Thalli transversal cross-sections in water **A–C**, ascospores **D–G** **A** thallus with typical pseudocortex (*Leptogium glebulentum* S-L49768) **B** thallus paraplectenchymateous throughout (*Leptogium glebulentum* S-L49768), **C** thallus with hyphae which are perpendicular to the surface (*C. furfuraceum* S-F492354) **D** spore of *C. ryssoleum* (UPS-L178905) **E** spore of *C. subnigrescens* (GZU66-201) **F** spore of *C. curtisporum* (UPS-L111603) **G** *C. nigrescens* (S-F492352), **tPsC** = typical pseudocortex, **Hp** = hyphae, **pPlect** = paraplectenchyma. Scale bar: 10 µm.

Among the isidiate species, *Collema nigrescens* is notable for its globular isidia (ca 0.2 mm in diameter; Fig. 2K, L), whereas *C. furfuraceum* has thinner, teretiform isidia (ca 0.05–0.15 mm wide) that become coralloid and reach up to 0.3 mm in length (Fig. 2M, N). Degelius (1974) later included *Collema luzonense* Räs. from the Philippines as a variety of *C. furfuraceum* and the var. *luzonense*, was distinguished by its euparaplectenchymatous excipulum proprium and frequent presence of white-pruinose apothecia. Another distinguishing feature of var. *luzonense* is its thicker lobes, reaching up to 300 µm, compared to 100 µm in var. *furfuraceum*. The var. *luzonense* has not been reported from Europe and the thallus thickness of the samples (which were all sterile) in our study suggests that they belong in var. *furfuraceum*.

Given the results of our phylogenetic analysis, it is crucial to study *C. furfuraceum* further. This study is under way, whereby we also plan to investigate the potential role of isidia as a distinguishing feature for species identification.

Conclusion

This study provides new insights into the phylogenetic relationships and morphological diversity among the European *Collema* species. Our results show that *C. glebulentum* belong in *Leptogium* s. str., which is supported by its

paraplectenchymatous thallus structure. Phylogenetic analysis reveals that Degelius' (1954) *Collema* "Flaccidum" and "Nigrescens"-groups are not supported by the actual phylogenetic relationships. The study indicates that there are still substantial taxonomic issues worth studying and clarifying within European *Collema* s. str. We will present the results of such a study in a forthcoming publication.

Acknowledgements

We gratefully acknowledge Lars Arvidsson, Barbara Klein, Martin Westberg, Martin and Maroš Dinga for their kind help during fieldwork. We are grateful to Björn Owe-Larsson for organizing an excursion to the Swedish island Lurö. We further thank Andre Aptroot, Toni Berglund, Johannes Bergsten, Robin Isaksson, Göran Odelvik and Jirka Malíček, for providing samples. Länsstyrelsen of Uppsala län provided a collecting permit for Fiby urskog Nature Reserve. The staff of the herbaria BG, GZU, TSB, S and UPS kindly provided loans. We are grateful to the Department of Bioinformatics and Genetics of the Swedish Museum of Natural History, in particular Bodil Cronholm, for her skillful lab assistance.

Additional information

Conflict of interest

The authors have declared that no competing interests exist.

Ethical statement

No ethical statement was reported.

Funding

We received generous funding from the Swedish Taxonomy Initiative (Svenska Artprojektet) administered by the Swedish Species Information Center (ArtDatabanken; grants 2016-207 4.3 and SLU.dha.2019.4.3-48), and the Stenholms donationsfond (GBT 2018-00002-26).

Author contributions

Conceptualization: MW, AK. Data curation: AK. Formal analysis: AK. Funding acquisition: AK, MW. Investigation: UN, FJ. Methodology: AK. Validation: FJ, UN, MW. Visualization: UN. Writing - original draft: AK. Writing - review and editing: UN, FJ, MW.

Author ORCIDs

Alica Košuthová  <https://orcid.org/0000-0001-5991-7444>

Mats Wedin  <https://orcid.org/0000-0002-8295-5198>

Data availability

All of the data that support the findings of this study are available in the main text.

References

- Degelius G (1954) The lichen genus *Collema* in Europe. *Symbolae Botanicae Upsalienses* 13: 1–499.
- Degelius G (1974) The lichen genus *Collema* with special reference to the extra-European species. *Symbolae Botanicae Upsalienses* 20: 1–215.

- Edgar RC (2004) MUSCLE: Multiple sequence alignment with high accuracy and high throughput. *Nucleic Acids Research* 32(5): 1792–1797. <https://doi.org/10.1093/nar/gkh340>
- Ekman S, Wedin M, Lindblom L, Jørgensen PM (2014) Extended phylogeny and a revised generic classification of the Pannariaceae (Peltigerales, Ascomycota). *Lichenologist* 46(5): 627–656. <https://doi.org/10.1017/S002428291400019X>
- Jørgensen PM (2007) Collemataceae. *Nordic Lichen Flora* 3: 14–45.
- Kearse M, Moir R, Wilson A, Stones-Havas S, Cheung M, Sturrock S, Buxton S, Cooper A, Markowitz S, Duran C, Thierer T, Ashton B, Meintjes P, Drummond A (2012) Geneious Basic: An integrated and extendable desktop software platform for the organization and analysis of sequence data. *Bioinformatics* 28(12): 1647–1649. <https://doi.org/10.1093/bioinformatics/bts199>
- Košuthová A, Fernández-Brime S, Westberg M, Wedin M (2016) *Collolechia* revisited, and a re-assessment of ascus characteristics in Placynthiaceae (Peltigerales, Ascomycota). *Lichenologist* 48(1): 3–12. <https://doi.org/10.1017/S0024282915000432>
- Košuthová A, Westberg M, Otálora MAG, Wedin M (2019) *Rostania* revised: Testing generic delimitations in Collemataceae (Peltigerales, Lecanoromycetes). *MycKeys* 47: 17–33. <https://doi.org/10.3897/mycokeys.47.32227>
- Larsson A (2014) AliView: A fast and lightweight alignment viewer and editor for large datasets. *Bioinformatics* 30(22): 3276–3278. <https://doi.org/10.1093/bioinformatics/btu531>
- Liu YJ, Whelen S, Hall BD (1999) Phylogenetic relationships among ascomycetes: Evidence from an RNA polymerase II subunit. *Molecular Biology and Evolution* 16(12): 1799–1808. <https://doi.org/10.1093/oxfordjournals.molbev.a026092>
- Miadlikowska J, Lutzoni F (2004) Phylogenetic classification of peltigeralean fungi (Peltigerales, Ascomycota) based on ribosomal RNA small and large subunits. *American Journal of Botany* 91(3): 449–464. <https://doi.org/10.3732/ajb.91.3.449>
- Miadlikowska J, Kauff F, Högnabba F, Oliver JC, Molnár K, Fraker E, Gaya E, Hafellner J, Hofstetter V, Gueidan C, Otálora MAG, Hodkinson B, Kukwa M, Lücking R, Björk C, Sipman HJM, Burgaz AR, Thell A, Passo A, Myllys L, Goward T, Fernández-Brime S, Hestmark G, Lendemer J, Lumbsch HT, Schmall M, Schoch CL, Sérusiaux E, Maddison DR, Arnold AE, Lutzoni F, Stenroos S (2014) A multigene phylogenetic synthesis for the class Lecanoromycetes (Ascomycota): 1307 fungi representing 1139 infrageneric taxa, 312 genera and 66 families. *Molecular Phylogenetics and Evolution* 79: 132–168. <https://doi.org/10.1016/j.ympev.2014.04.003>
- Miller MA, Pfeiffer W, Schwartz T (2010) Creating the CIPRES Science Gateway for inference of large phylogenetic trees. *Proceedings of the Gateway Computing Environments Workshop (GCE)*. New Orleans, LA, 1–8. <https://doi.org/10.1109/GCE.2010.5676129>
- Myllys L, Lohtander K, Tehler A (2001) β -Tubulin, ITS and group I intron sequences challenge the species pair concept in *Physcia aipolia* and *P. caesia*. *Mycologia* 93(2): 335–343. <https://doi.org/10.1080/00275514.2001.12063165>
- Otálora MAG, Aragón G, Molina MC, Martínez I, Lutzoni F (2010) Disentangling the *Collema-Leptogium* complex through a molecular phylogenetic study of the Collemataceae (Peltigerales, lichen-forming Ascomycota). *Mycologia* 102(2): 279–290. <https://doi.org/10.3852/09-114>
- Otálora MAG, Aragón G, Martínez I, Wedin M (2013a) Cardinal characters on a slippery slope—a re-evaluation of phylogeny, character evolution, and evolutionary rates in the jelly lichens (Collemataceae s. str.). *Molecular Phylogenetics and Evolution* 68(2): 185–198. <https://doi.org/10.1016/j.ympev.2013.04.004>

- Otálora MAG, Jørgensen PM, Wedin M (2013b) A revised classification of the jelly lichens, Collemataceae. *Fungal Diversity* 64[2014]: 275–293. <https://doi.org/10.1007/s13225-013-0266-1>
- Schmitt I, Crespo A, Divarak PK, Fankhauser JD, Herman-Sackett E, Kalb K, Nelsen MP, Nelsen NA, Rivas-Plata E, Shimp AD, Widhelm T, Lumbsh HT (2009) New primers for promising single-copy genes in fungal phylogenetics and systematics. *Persoonia* 23(1): 35–40. <https://doi.org/10.3767/003158509X470602>
- Stamatakis A (2014) RAxML version 8: A tool for phylogenetic analysis and post-analysis of large phylogenies. *Bioinformatics* 30(9): 1312–1313. <https://doi.org/10.1093/bioinformatics/btu033>
- Thiers B (2018) [Continuously updated] Index Herbariorum: A Global Directory of Public Herbaria and Associated Staff. New York Botanical Garden's Virtual Herbarium. <http://sweetgum.nybg.org/science/ih/>
- Wedin M, Wiklund E, Jørgensen PM, Ekman S (2009) Slippery when wet: Phylogeny and character evolution in the gelatinous cyanobacterial lichens (Peltigerales, Ascomycetes). *Molecular Phylogenetics and Evolution* 53(3): 862–871. <https://doi.org/10.1016/j.ympev.2009.08.013>
- Weerakoon G, Aptroot A, Wedin M, Ekman E (2018) *Leightoniella zeylanensis* belongs to the Pannariaceae. *Nordic Journal of Botany* 36(7): 1–6. <https://doi.org/10.1111/njb.01880>
- Wiklund E, Wedin M (2003) The phylogenetic relationships of the cyanobacterial lichens in the Lecanorales suborder Peltigerineae. *Cladistics* 19(5): 419–431. <https://doi.org/10.1111/j.1096-0031.2003.tb00312.x>
- Zhang Z, Schwartz S, Wagner L, Miller W (2000) A greedy algorithm for aligning DNA sequences. *Journal of Computational Biology* 7(1–2): 203–214. <https://doi.org/10.1089/10665270050081478>
- Zoller S, Scheidegger C, Sperisen C (1999) PCR primers for the amplification of mitochondrial small subunit ribosomal DNA of lichen-forming ascomycetes. *Lichenologist* 31(5): 511–516. <https://doi.org/10.1006/lich.1999.0220>