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*Inocybe praetervisa* group — A clade of four closely related species with partly different geographical distribution ranges in Europe

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**Abstract**

Sequence data from a broad geographical region and different habitats show that the *Inocybe praetervisa* clade is comprised of four closely related species. These species of section *Marginatae* are characterized by having nodulose spores and a stipe that is abundantly pruinose only in the upper half. *Inocybe praetervisa* occurs in Southern Europe in mountainous mixed coniferous forests, and is not confirmed from Northern Europe. *Inocybe rivularis* occurs in northern boreal forests up to the lower alpine zone, associated with *Betula* in moist habitats, and is not confirmed from Southern Europe. *Inocybe taxocystis* is confirmed as a later synonym of *I. favrei*. The species has a wide geographical distribution range in Europe, mainly restricted to the alpine zone and moist soils, associated with *Salix herbacea*. *Inocybe arctica* is here described as a new species. It occurs in the arctic and higher alpine zones, associated with *Dryas octopetala, Salix polaris, S. reticulata* and *S. herbacea*. All species except *I. arctica* are shown to have an intercontinental distribution range and are confirmed from North America. Sequence data suggest the occurrence of one additional species in the alpine zone of China. A key to the species in the *I. praetervisa* group is provided.

1. Introduction

*Inocybe* section *Marginatae* Kühner (1933) is known to be comprised of several species complexes where names have been interpreted differently and often misapplied (Esteve-Raventós, Moreno, Bizio, & Alvarado, 2015, 2016; Jacobsson & Larsson, 2012; Vauras & Larsson, 2015).

The species in *Marginatae* are usually characterized by the absence of a cortina, having nodulose spores and caulocystidia descending to the base. However species in the *I. praetervisa* clade, as defined by Esteve-Raventós et al. (2016), have abundant caulocystidia in the upper half that are sparse or absent in lower half of the stipe. In this recent paper dealing with the taxonomy of the *Inocybe praetervisa* group (Esteve-Raventós et al., 2016) a lectotype for *I. praetervisa* Quél. was selected. In addition, an epitype was designated, from a representative specimen of the Bresadola herbarium collected from the type locality in Italy in order to fix the name also by sequence data of the internal transcribes spacer (ITS) region (Ariyawansa et al., 2014).

In morphology, *I. praetervisa* is characterized by having a smooth yellowish-brown pileus that becomes darker brown with age and a margin that is rimose. The lamellae are at first pale white with a greyish tint, and are later more brown. The stipe is pale whitish at first and later yellowish brown, possesses a more or less marginate bulbous stipe base, and is densely pruinose in the upper half. Spores are heterodiametric and distinctly nodulose. Cheilo- and pleurocystidia are utriform to broadly fusiform, and rather thick-walled with crystals at the apex. Specimens become darker brown upon drying but are not blackening. The selected epitype, collected by Bresadola, originates from a coniferous forest. For a more detailed description see Esteve-Raventós et al. (2016).

In the Esteve-Raventós et al. (2016) study, phylogenetic analyses based on ITS sequences were performed to infer phylogenetic relationships among the species. In the phylogeny presented, the sequence of a paratype of *I. rivularis* Jacobsson & Vauras fell within the *I. praetervisa* clade with strong support, suggesting it should be a later synonym of *I. praetervisa*. Accordingly, based on the
Inocybe rivularis was regarded as a synonym of *I. praetervisa* (Esteve-Raventós et al., 2016).

Inocybe rivularis was described by Jacobsson and Vauras (1989) from moist forest habitats in the boreal zones of Fennoscandia, likely associated with *Betula* spp., *Picea abies* (L.) H. Karst. and *Salix* spp. The species is considered to have a rather wide distribution and is common in suitable habitats, usually close to small spring brooks. In morphology it is rather similar to the description of *I. praetervisa* (Esteve-Raventós et al., 2016) but was characterized as having a cortina in young specimens that soon disappears leaving no remnants on the stipe. The stipe base is clavate to bulbous but not marginately bulbous. As the species was densely pruinose only in the upper part of the stipe it was at first thought to belong in section *Cortinatae* Kühner.

A species with similar morphology is *I. taxocystis* (Fr.) Fr. Singer. The species was first described by Favre (1955) as a variety of *I. decipientoides* Peck (*I. decipientoides var. taxocystis* Favre), from the Swiss alpine zone associated with *Salix herbacea* L. and *Dryas octopetala*. Favre’s variety was characterized by having a stipe that is pruinose at least to half the stipe, glabrous below, with a bulbous but not marginate stipe base. It was first raised to species level by Singer (1986), but without selecting a lectotype. It was also described as the new species *Asterosporina taxocystis* J. Favre & E. Horak by Horak (1987), and later moved to *Inocybe* (Fr.) Fr. by Sinn-Irlet (1992).

Another species with similar morphology and habitat was published as *I. favrei* Bon (1985). The species was described with a stipe that is more or less pruinose particularly in the upper half and with an obtusely marginate stipe base, and is associated with *S. herbacea*, *S. reticulata* L. and *Dryas*. The species was described by Bon to accommodate the description of the species *I. decipiens* Bres. sensu Kühner by Favre from the alpine zone (Favre, 1955). Bon designated a holotype (Bon 84095) from the alpine zone in France associated with *S. herbacea*.

Analyses of ITS sequence data have become an important tool for evaluating the morphological characters used to determine and diagnose species. This has been especially valuable for groups with morphologically similar characters and seemingly broad ecology, as often encountered within *Inocybe* (Cripps, Larsson, & Horak, 2010; Kokkonen & Vauras, 2012; Larsson, Vauras, & Cripps, 2014; Vauras & Larsson, 2015). The ITS region is now agreed upon as the bar code marker for fungi (Schoch et al., 2012) and has been found to be suitable and to work well for delimitation and segregation of species in the majority of the Basidiomycetes (Köljalg et al., 2013).

In the present study, we compare morphology and sequence data of a broader sampling of specimens described as *I. rivularis*, *I. taxocystis* and *I. favrei*, to get a better picture of *I. praetervisa* sensu Esteve-Raventós et al. (2016).

## 2. Material and methods

### 2.1. Morphological methods

The majority of the sequenced specimens were collected by the authors. Additional material received as loans from different herbaria were studied (AH, G, LIP, Q, TUR, S, UPS). Abbreviation of herbaria follows Index Herbariorum (http://sweetgum.nybg.org/science/ih/).

Macro-morphological characters were noted and cross-sections drawn from some fresh basidiomata. The color codes refer to Cailleux (1981). Micro-morphological characters were measured and drawn from dried material mounted in 10% NH4OH solution following the methodology described in Vauras and Kokkonen (2009). Photos of micro-morphological characters were made using an Axioskop 2 (Zeiss, Oberkochen, Germany) light microscope and the AxioVision software (http://www.zeiss.com/microscopy/int/home.html).

### 2.2. Molecular methods

Sixty-three specimens from Northern Europe, Svalbard, and North America were targeted for sequencing in this study, including the holotype of *I. favrei* (Bon 84095, LIP!). In addition the ITS sequence of the lectotype of *I. taxocystis* (JNS80884) and paratype of *I. rivularis* (JNS80885) from the study by Kokkonen and Vauras (2012), the epitype of *I. praetervisa* (KT201675), *I. tabacina* Furrer-Ziogas and *I. phaeosticta* Furrer-Ziogas from the study by Esteve-Raventós et al. (2016) were included. Also a few additional ITS and LSU sequences of *Inocybe* section *Marginatae* published in Ryberg, Larsson, and Jacobsson (2010) were included. For comparison and rooting of trees the sampling was selected to correspond with Clade B in Esteve-Raventós et al. (2016). Sequence data of the ITS and LSU regions of *I. fibrosilvae* Kühner (EL51–14; Genbank KY033846) and *I. decemgibbosa* (Kühner) Vauras (JV8115; Genbank KY033847) were generated. Each target ITS sequence type was also blasted in GenBank (Clark, Karsch-Mizrachi, Ostell, & Sayers, 2016) and the UNITE database (Köljalg et al., 2013) to search for additional data generated from specimens, environmental samples as soil and ectomycorrhizae. In all 13 sequences were found and added to the data set.

Sequences from the complete internal transcribed spacer (ITS) region and about 1400 or 375 base pairs (bp) of the 5′ end of the large subunit (LSU) of the nuclear ribosomal DNA were generated. DNA extractions were performed using DNeasy Mini Kit (Qiagen, Hilden, Germany), PCR reactions using Illuma PuReTaq ReadyToGo PCR beads (GE Healthcare, Buckinghamshire, United Kingdom). For PCR clean up QiAquick PCR purification kit (Qiagen, Hilden, Germany) was used. The DNA from the type specimen was extracted using a CTAB method, and primers for PCR followed protocols described in Larsson and Jacobsson (2004). Primers used to amplify the complete ITS region and the 5′ end of the LSU region were ITS1F (Gardes & Bruns, 1993) and LR21, LR0R, and LR7 (Hopple & Vilgalys, 1999). Primers used for sequencing were ITS1, ITS4 (White, Bruns, Lee, & Taylor, 1990), Ctb6 (https://nature.berkeley.edu/bruns/lab), LR3R and Lr5 (Hopple & Vilgalys, 1999).

Sequences were edited and assembled using Sequencher 5.1 (Gene Codes, Ann Arbor, Michigan). Sequences generated for this study have been deposited in GenBank and accession numbers given (KY033785–KY033847). Sequences sequenced in this study are indicated with an asterix (*) and the GenBank number is noted in the list of specimens examined.

Alignment was performed using the L-INS-i strategy as implemented in MAFFT v. 7.017 (Katoh and Standley, 2013). The alignment was adjusted using Alview 1.171 (Larsson, 2014). For inferring phylogenetic relationships among species heuristic searches for the most parsimonious trees were performed using PAUP* (Swofford, 2003). All transformations were considered unordered and equally weighted. Variable regions with ambiguous alignment were excluded and gaps were treated as missing data. Heuristic searches with 1000 random-addition sequence replicates and TBR branch swapping were performed, saving at most 25 trees in each replicate. Relative robustness of clades was assessed by the bootstrap method using 1000 heuristic search replicates with 100 random taxon addition sequence replicates and TBR branch swapping, the latter saving at most 100 trees in each replicate. In addition a Bayesian analysis was carried out in MrBayes 3.2.6 (Ronquist & Huelsenbeck, 2012), with a best-fit model of nucleotide evolution supplied by MrModeltest 2.2 (Nylander, 2004).
3. Results

The aligned complete dataset consisted of 95 sequences and 2095 characters. After exclusion of ambiguous regions mainly from the beginning and the end of the data set 1977 characters remained for the analysis. Of these, 1670 were constant, 57 were variable but parsimony uninformative, and 250 (12.6%) were parsimony informative. The maximum parsimony analysis yielded 18,950 equally most parsimonious trees (length = 460 steps, CI = 0.8217, and RI = 0.9359). The 50% majority-rule consensus bootstrap tree is presented in Fig. 1. The bootstrap analysis recovered the *I. praetervisa* complex with 100% support. Five supported clades with low to strong bootstrap support were found within the *I. praetervisa* complex, corresponding to *I. rivularis* and *I. favrei* (see taxonomy section, 83%). The four last clades form a subclade within the complex with a support value of 71%.

As suggested by MrModeltest, the nucleotide evolution model HKY + G was used for the ITS1 spacer; GTR + I was used for the 5.8S gene; HKY + G was used for the LSU in the Bayesian analysis. The MCMC analysis converged well in advance of the burn-in threshold and chain mixing was good. The MCMC analysis converged well in advance of the burn-in threshold and chain mixing was good. The MCMC analysis converged well in advance of the burn-in threshold and chain mixing was good. The MCMC analysis converge...
GenBank KY033831); Svalbard, Nordenskiöld Land, Longyearbyen. 2002, B. Gahne 02-96 (GB; GenBank AM882980); Svalbard, Nordenskiöld Land, Bolterdalen, moist with Salix polaris, 13 Aug 2015, A. Molia 25-2015 (O; NOBAS2299, KY033835). SWEDEN, Torne Lappmark, Jukkasjärv, Latnjajure, moist with Salix reticulata, 10 Aug 2007, E. Larsson 72-07* (GB; TUR-A; GenBank KY033839); 7 Aug 2006, with Salix polaris, E. Larsson 71-06* (GB; TUR-A; GenBank KY033842); Lule lappmark, jokkmokk, padelanta NP, vielgggisbakte on NW side of Vastenjaure Lake, near Salix herbacea, 11 Aug 2016, J. Vauras 31471* (TUR-A; GB; GenBank KY033840); Lule lappmark, jokkmokk, padelanta NP, Arranoojvve on N side of the Vastenjaure Lake, near Salix herbacea, 11

**Fig. 1.** The 50% majority-rule bootstrap consensus tree from the parsimony analysis. Parsimony bootstrap values and Bayesian posterior probabilities are indicated on branches. Clades discussed in the text are indicated with bars and species epithets. Sequences originating from type specimens are marked. Clade B refers to Esteve-Raventós et al. (2016).
Aug 2016, J. Vauras 31460* (TUR-A; GB; GenBank KY033841); Lule lappmark, Jukkasjärvi, Padjelanta NP, Slaphejaure NO, with Salix herbacea, 14 Aug 2016, E. Larsson 117-16* (GB; GenBank KY033845).

*Inocybe praetervisa* Qué., in Bresadola, Fung. Trident. 1:35, 1881. Fig. 4E, F.

For the description of *I. praetervisa*, we refer to Esteve-Raventós *et al.* (2016). The species resembles *I. rivularis*, but differs somewhat by having smaller spores often with a long apical node, on average smaller nodules, and in the form of the pleurocystidia, which tend to attenuate at the apex on average more than in *I. rivularis*.

Micro-morphological measurements: Spores yellow-brown, (8.8–)9.1–11.4(–12.7) × (6.0–)6.2–7.4(–8.1) μm, on average 10.2 × 6.8 μm, range of mean values 9.8–10.4 × 6.5–6.9 μm, Q = (1.2–)1.3–1.7(–2.0), on average 1.50, range of mean Q-values 1.49–1.52 (80 spores from 3 collections). Basidia 31–38 × 10–12 μm, on average 34 × 11 μm, generally 4-spored, clavate (8 basidia from 1 collection). Pleurocystidia (55–)59–83(–92) × (15–)18–28(–29) μm, on average 69 × 22 μm, range of mean values 67–71 × 21–24 μm (54 pleurocystidia from 3 collections), mainly fusiform, thick-walled, with up to 4 μm pale yellow to yellow wall. Caulocystidia abundant at upper part of stipe, very few or not in the lower half, variable, thick-walled, some cauloparacystidia thick-walled.

Specimens examined: ITALY, Bolzano, NE of Campo Túres, Riva di Túres, coniferous forest, ca. 1600 m, 23 Jul 2006, G. Verol (J. Vauras 23918*; TUR-A; GB; GenBank KY033785); Trentino, Rabbi, Jul 1909, in coniferous forest, G. Bresadola (S-F29598, epitype); SPAIN, Madrid, Rascafria, *Pinus sylvestris* and *Salix* sp., 8 Jul 2013, leg. F. Pancorbo, det. F. Esteve-Raventós AH44415 (AH, GB).


For the description of *I. rivularis* we refer to Jacobsson and Vauras (1989), repeating here only the sizes of spores and pleurocystidia, and the distribution of caulocystidia. Spores (9–)10–12.5(–13.5) × 6.5–8.5(–9) μm, on average 11.1 × 7.8 μm, range of mean values 10.5–12.3 × 7.1–8.1 μm, Q = 1.2–1.6(–1.8), average 1.43, range of mean Q-values 1.35–1.6 (300 spores from 15 collections). Pleurocystidia 50–76(–90) × 14–28(–33) μm. With further experience we complement the description and add that sometimes the species can have fairly abundant velipellis on the pileus (Fig. 5C), and with few thick-walled caulocystidia down to base of stipe. Further, it can be found even in the low-alpine zone with *Betula nana*. Even though this quite large species is fairly common in suitable habitats in northern part of Fennoscandia, there are rather few reports of it (Alpago Novello, 2002). It has been selected as one of the indicator fungi for forests of Finland, associated with spruce mires, brook-side forests and spring habitats (Bonsdorff *et al.*, 2014).

Specimens examined: CANADA, Quebec, Schefferville area, Snowy Channel, alpine region, moist depression, growing by water, 14 Aug 1963, leg. P. Kallio 482*, det. J. Vauras (TUR; GenBank KY033825); 14 Aug 1963, leg. P. Kallio 489* (TUR); ESTONIA, Hiium, Käina, moist forest with *Picea abies*, *Betula sp.*, *Alnus glutinosa* (L.) Gaertn., *Populus tremula* L., *Pinus sylvestris* L. and *Fraxinus excelsior* L., 14 Sep 2010, J. Vauras 27165 (TUR-A; TU: TAAI01616, UNITE UDB011734); FINLAND, Oulu Pohjanmaa, Oulu, Sanginsuus, Hakala, 4 Aug 1986, J. Ruotsalainen and J. Vauras 2152F (TUR! Holotype; GenBank JN580885); Inarin Lappi, Utsjoki, Kevo, around Kevojärvi, subarctic birch forest with *Salix* spp., close to the lake. 30 Aug 2012, E. Larsson 178–12*(GB; GenBank KY033823); 27 Aug 2012, subarctic birch forest with *Salix* spp., E. Larsson 160–12* (GB; GenBank KY033824); Inarin Lappi, Utsjoki, Kevo, Kevojärvi, W. slope of Korkea-Jehkas, SE of Jehkasjärvi, low alpine slope with small brooks, near *Betula nana* and *Salix* spp., amongst moist mosses, 24 Aug 2014, J. Vauras 30708* (TUR-A; GenBank KY033802); Enontekion Lappi, Enontekiö, Kilpisjärvi, Malla Strict Reserve, Mountain Pikkut-Malla, alpine site with *Salix* spp. and *Betula nana*, close to snow bed, moist, gently sloping site with mosses and stones, 7 Aug 1985, J. Ruotsalainen and J. Vauras 1929* (TUR-A; GenBank KY033804); Enontekion Lappi, Enontekiö, Kilpisjärvi, mountain birch forest near small river Tsahkaljohka, rather moist site by path, 13 Aug 1990, J. Vauras 4744* (TUR-A; GenBank KY033808); Enontekiö Lappi, Enontekiö, Kilpisjärvi, E side slope of Jehkas, snow bed area, moist with *Salix herbacea*, 29 Aug 2013, E. Larsson 288–13* (GB; GenBank KY033815); Kainuu, Ristijärvi, church village, Lukkarila, camping place Ristikärven Pirrit, open.

**Fig. 2.** *Inocybe arctica.* Basidiomata of the holotype growing with *Salix polaris* and *Bistorta vivipara* (J. Vauras 2238F).
park lawn with Betula sp., 27 Aug 2016, J. Vauras 31686* (TUR-A; GenBank KY033810); Pohjois-Häme, Multia, Lehtomäki, E of the farm house, pastured forest on E slope with Picea abies and Betula sp., on mull soil, 8 Sep 2013, J. Issakainen and J. Vauras 30173* (TUR-A; GenBank KY033813); 8 Sep 2013, E of the farm house, pastured forest on E slope with Picea abies and Betula sp., Pinus sylvestris, Alnus incana (L.) Moench., Populus tremula and Salix caprea L., near brook, J. Issakainen and J. Vauras 30172* (TUR-A; GenBank KY033820); Perä-Pohjanmaa, Rovaniemi, Meltau, N of the Meltaus – Marrasjärv, SE of Sortovaara, Kenkäkorpi, herb-rich forest with Picea abies, Populus tremula, Betula sp., Alnus incana, Salix caprea and Pinus sylvestris, with moist depressions, 23 Aug 1999, J. Vauras 15316*(TUR-A; GenBank KY033822); SWEDEN, Lycksele lappmark, Tärna, Hemavan, Porthron, Jóbäcken, subalpine Betula pubescens forest with Salix spp., 27 Jul 2014, E. Larsson 24-14* (GB; GenBank KY033803); Lule lappmark, Jukkasjärv, Padjelanta NP, Huorso, Vastejaure, among Salix spp., 1 Aug 2001, S. Kuoljok 0125* (GB; GenBank KY033805); 5 Aug 2000, torvkulle med Betula nana, S. Kuoljok 0052* (GB; GenBank KY033809); Lule lappmark, Jukkasjärv, Padjelanta NP, Njoammeljaure NO, moist with Betula nana and Salix spp., 18 Aug 2016, E. Larsson 241-16* (GB; GenBank KY033806); Lule lappmark, Jukkasjärv, Padjelanta NP, Njoammeljaure, moist depression on S slope, near Salix spp. and Betula nana, 18 Aug 2016, J. Vauras 31568* (TUR-A; GenBank KY033801); Lule lappmark, Jukkasjärv, Padjelanta NP, Slahpejaure NO, moist area with small brook, with Betula nana and Salix spp., 14 Aug 2016, E. Larsson 129-16* (GB; GenBank KY033807); Lule lappmark, Jukkasjärv, Padjelanta NP, Unna Duvge, moist area with Betula nana and Salix spp., 15 Aug 2016, E. Larsson 158-16* (GB; GenBank KY033812); Torne lappmark, Jukkasjärv, Latnajaure, moist with Salix herbacea and Betula nana, 2 Aug 2006, E. Larsson 23-06* (GB; GenBank KY033814); Härjedalen, Tännäs, Andersborgsvägen, moist

area with brooks, *Betula pubescens*, *B. nana* and *Salix* spp., 15 Aug 2006, E. Larsson 82-06* (GB; GenBank KY033817); 15 Aug 2006, E. Larsson 82B-16* (GB; GenBank KY033816); Värmland, Sunne, Gärdesrud, 26 Aug 1990, B. Jansson* (GB; GenBank KY033818); NORWAY, Troms, Storfjord, Lulleslåttan, mossy forest with *Pinus sylvestris*, *Betula* sp. and scattered *Picea abies*, on tractor track, 22 Aug 2014, J. Vauras 30676* (TUR-A; GenBank KY033819); Oppland, Valdres, W of Fisketjernsknauen, fjällhed, 7 Jul 1995, M. Jeppson 3722* (GB; GenBank FN550887).

**Fig. 4.** Comparison of basidiospores and pleurocystidia of *Inocybe* spp. A, B: *I. arctica* (EL 34-09; A basidiospores, B pleurocystidia). C, D: *I. favrei* (JV 31508F; C basidiospores, D pleurocystidia). E, F: *I. praetervisa* (JV 23918; E basidiospores, F pleurocystidia). G, H: *I. rivularis* (EL & JV 30072F; G basidiospores, H pleurocystidia). Bars: 10 μm.


For the description of *I. favrei* we refer to Bon (1985) and Favre (1955). In macro-morphology, *I. favrei* resembles *I. rivularis* and *I. arctica*. In micro-morphology, it is characterized by having cystidia (both pleurocystidia, cheilocystidia and caulocystidia) that are ventricose and are fairly broad at the apex (see Figs. 4–6). Further, the spores of *I. favrei* are on average shorter and wider compared with *I. arctica*. The species is most common in the low alpine zone, but has been found on few occasions in forests of the northern boreal zone on calcareous soil.

Micro-morphological measurements: Spores nodulose, but variable in form and size, fairly dark yellow-brown, thick-walled, (9.7–11.3) × (6.4–6.6–8.4–8.7) μm, on average 10.9 × 7.3 μm, range of mean values 10.9–11.4 × 7.1–7.5 μm, Q = (1.25–1.4–1.7–1.75), on average 1.52, range of mean Q-values 1.50–1.59 (100 spores from 5 collections). Basidia (28–40) × (10–14–14–14 μm, on average 33 × 13 μm, generally 4-spored, clavate (17 basidia from 4 collections). Pleurocystidia (54–158–74–80) × (11–)14–26 (–30) μm, on average 66 × 20 μm, range of mean values 63–70 × 16–25 μm (83 pleurocystidia from 5 collections), mainly ventricose, thick-walled, with up to 4 μm pale

yellow to yellow wall. Cheilocystidia 44–65 × 14–25 μm, on average 56 × 19 μm (n = 15), mainly ventricose, abundant, paracystidia pyriform, some thick-walled. Caulocystidia abundant at stipe apex, (40–45–66–68) × (13–16–24–27) μm, on average 54 × 20 μm (n = 20), more rare but present to stipe base, variable, thick-walled, some of the cauloparacystidia thick-walled.

Specimens examined: FINLAND, Enontekiö Lappi, Enontekiö, Kilpisjärvi, at border of Finland and Norway, W of the main road, low alpine site at small brook, near Betula nana and Salix spp., 20 Aug 2014, J. Vauras 30653F* (TUR-A; GB; GenBank KY033797); Koillismaa, Kuusamo, Jakälniemi, fairly moist forest with Picea abies, Pinus sylvestris and Betula sp., 28 Aug 2011, J. Vauras 28431* (TUR-A; GenBank KY033795); FRANCE, Savoie, Haute-Maurienne, Termignon, Bellecombe, parc national de la Vanoise, 30 Aug 1984, Bon 84095* (LIP; Holotype; GenBank KY033786); Savoie, Bourg-Saint-Maurice, Arc 2000, lac Marloup, 21 Aug 1993, with Salix herbacea on wet soil along a small lake, higher alpine zone, 2500 m, leg. P.-A. Moreau, PAM93082101* (LIP; GenBank KY033799); NORWAY, Troms, Storfjord, near the border of Finland, Bossovarri, low alpine heath, near dried pond, Salix spp. and Betula nana, 21 Aug 2014, J. Vauras 30673F* (TUR-A; GenBank KY033798); SWEDEN, Torne lappmark, Jukkasjärvi, Latnajaure, with Salix herbacea, 6 Aug 2006, E. Larsson 57-06 (GB; TUR-A; GenBank FN550886); 7 Aug 2006, moist place with Salix herbacea and Bistorta vivipara, E. Larsson 78-06* (GB; GenBank KY033800), 5 Aug 2011, with Salix herbacea, E. Larsson 105-11* (GB; GenBank KY033796); Lule lappmark, Jokkmokk, Padjelet NA, Arralábbdâ, fjällhed med Salix herbacea, 11 Aug 2016, E. Larsson 41-16* (GB; GenBank KY033794); Lule lappmark, Jokkmokk, Padjelet NA, Vieggbisbakte, in snow bed area with Salix herbacea, 12 Aug 2016, E. Larsson 69B-16* (GB; GenBank KY033793); Lule lappmark, Jukkasjärvi, Padjelet NA, Slahejare NO, in snow bed area with Salix herbacea, 14 Aug 2016, E. Larsson 124-16* (GB; GenBank KY033792), 14 Aug 2016, on S slope, moist boggy meadow near Salix polaris, S. reticulata, J. Vauras 31508* (TUR-A; GenBank KY033789); 14 Aug 2016, on S slope near Salix herbacea, J. Vauras 31515* (TUR-A; GenBank KY033788); Lule lappmark, Jukkasjärvi, Padjelet NA, Njoamnejeare NO, with Salix herbacea and Bistorta vivipara, 18 Aug 2016, E. Larsson 229-16* (GB; GenBank KY033791); 16 Aug 2016, snow bed area with Salix herbacea, E. Larsson 231-16* (GB; GenBank KY033790); 16 Aug 2016, at dried pond near Salix and Betula nana, J. Vauras 31579 (TUR-A; GB; GenBank KY033787); SWITZERLAND, Kt. Graubünden, Tarasp, Munt de la Bescha, Salix herbacea, 8 Sep 1945, J. Favre 122a* (G; Lectotype; GenBank JN580884); Val S-charl, Mount Plazer, on tapis de Salix herbacea, sol gneissique, 20 Aug 1952, J. Favre 122c (G); 26 Aug 1952, J. Favre 122-1 (G).
5. Key to the species in the *I. praetervisa* clade

1a. Alpine/arctic ecosystems

1b. Forest ecosystems

2a. High alpine to arctic, spores on average 11.5–12.2 × 7.2–7.9 μm, pleurocystidia on average 65–77 × 14–20 μm

2b. Sub- to low alpine, spores on average 10.9–11.4 × 7.1–7.5 μm, pleurocystidia on average 63–70 × 16–25 μm

3a. N. Europe and Canada, with *Betula*, spores on average 10.5–12.3 × 7.1–8.1 μm

3b. S. Europe and N. America, in coniferous forests, spores on average 9.8–10.4 × 6.5–6.9 μm

6. Discussion

With the support from phylogenetic analyses of sequence data we show that the *I. praetervisa* clade is comprised of at least four morphologically and genetically similar but separate species. In addition, two sequences from ectomycorrhizal (ECM) root tips isolated from *Kobresia* Willd. in the alpine zone of China form a sub-clade within this group, indicating the occurrence of one additional species in Asia. The species show overlapping but partly different geographical distribution patterns in Europe.

*Inocybe praetervisa* is confirmed to have an intercontinental distribution range. According to our data it occurs in the mountains of Southern Europe then associated with coniferous mixed forests and in Canada associated with *Pseudotsuga* Carrière. Also, seven sequences deposited in GenBank under various names for collections from British Columbia, lacking information on host or forest type, are 100% identical with the epitype ITS sequence. So far, we have no specimens or evidence from sequence data of environmental samples that confirm the occurrence of the species in Northern Europe, although it has frequently been reported. However, these reports are based on misinterpretations of the species concept ([Estèves-Raventós et al., 2016]). It likely reflects the fact that there are rather many species with similar morphology and traditionally *I. praetervisa* has been treated as a species with a broad species concept in the Nordic countries ([Jacobsson & Larsson, 2012]). It was thought to be a species with a more or less completely pruinose stipe, a marginate bulbous stipe base and a pileus and stipe which darken upon drying. All four species treated completely pruinose stipe, a marginate bulbous stipe base and a brown to dark brown pileus, occasionally with a reddish tint, an absence of veil, and a pruinose stipe with a marginate bulb at the stipe base. In addition, at pileus and stipe which darken upon drying. All four species treated have a more or less bulbous, and coniferous mixed forests. The residual specimens originate from northern boreal forests to the alpine zone, in association with *Betula nana*, various *Salix* shrubs, *S. herbacea* and *S. reticulata* on moist places as close to small brooks and in snow bed areas. The three sequences from Alaska, downloaded from Genbank, are isolated from soil and from ECM of *Picea mariana*, and suggest that it also occurs in North America with a northern distribution in the taiga and boreal forest ecosystems.

In the molecular data, *I. rivularis* shows a much larger sequence variation than *I. favrei*. The herein recognized *I. rivularis* clade receives a week bootstrap support and eight of the sequences form a sub-clade with 84% support value (Fig. 1). The collections within the sub-clade are mainly collected in boreal to subalpine regions, likely associated with *Betula pendula* Roth., *P. pubescens* or *B. nana*. One of the specimens originates from Estonia and was collected in moist mixed forest. The residual specimens originate from northern boreal forests to the alpine zone, in association with *P. pubescens*, *B. nana*, various *Salix* shrubs and *S. herbacea* in moist habitats. The species seems to have a northern distribution within Europe, with the most southern confirmed specimens from Estonia. Sequence data of one specimen collected in Canada (Quebec) was included and confirms the species also from North America. With the data at hand the genetic differences cannot be correlated with differences in ecology, geographic distribution or morphology. So, we choose to recognize the major clade despite the low support value.

*Inocybe arctica* is here described and represents the recovered clade within the complex without a name. Our specimens originate from Svalbard and Fennoscandia, mainly from the arctic and higher alpine zones. On Svalbard it is one of the most common *Inocybe* species found in mats of *Dryas octopetala* and with *Salix polaris*. Specimens from the higher alpine zone in Fennoscandia are also found with *D. octopetala* and *S. polaris* but also with *S. reticulata* and *S. herbacea*. In comparison to *I. rivularis* and *I. favrei*, seems to have preferences for more rich and calcareous soils. *Inocybe arctica* seems to occur in higher alpine regions in Northern Fennoscandia than *I. favrei*. *Inocybe favrei* is not confirmed from Svalbard and
I. arctica has not yet been confirmed from North America.

Disclosure

The authors declare no conflict of interest.

Acknowledgments

Curators of herbaria G, LIP, AH, O are gratefully acknowledged for arranging loans, and Pierre-Arthur Moreau, Gro Gulden, Fernando Esteve-Raventós and Fermin Pancorbo for sharing interesting collections and photos. Seppo Huhtinen and reviewers for valuable comments on the ms. Financial support was received from The Swedish Taxonomy Initiative, ArtDBanken SLU Uppsala, Anna och Gunnar Vidfelts Fond, Stiftelsen Wilhelm och Martina Lundgrens Vetenskapsfond, and the FinBOL-project funded by the Finnish Cultural Foundation and Kone Foundation.

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