

Microbotryum silenes-saxifragae sp. nov. sporulating in the anthers of *Silene saxifraga* in southern European mountains

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Abstract: Currently, the monophyletic lineage of anther smuts on *Caryophyllaceae* includes 22 species classified in the genus *Microbotryum*. They are model organisms studied in many disciplines of fungal biology. A molecular phylogenetic approach was used to resolve species boundaries within the caryophyllaceous anther smuts, as species delimitation based solely on phenotypic characters was problematic. Several cryptic species were found amongst the anther smuts on *Caryophyllaceae*, although some morphologically distinct species were discernible, and most species were characterized by high host-specificity. In this study, anther smut specimens infecting *Silene saxifraga* were analysed using rDNA sequences (ITS and LSU) and morphology to resolve their specific status and to discuss their phylogenetic position within the lineage of caryophyllaceous anther smuts. The molecular phylogeny revealed that all specimens form a monophyletic lineage that is supported by the morphological trait of reticulate spores with tuberculate interspaces (observed in certain spores). This lineage cannot be attributed to any of the previously described species, and the anther smut on *Silene saxifraga* is described and illustrated here as a new species, *Microbotryum silenes-saxifragae*. This species clusters in a clade that includes *Microbotryum* species, which infect both closely and distantly related host plants growing in diverse ecological habitats. It appears possible that host shifts combined with changes to ecological host niches drove the evolution of *Microbotryum* species within this clade.

Key words:

Anther smuts
Caryophyllaceae
Microbotryales
Microbotryum violaceum complex
Molecular phylogenetics
Plant pathogens
Pseudo-cryptic species

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INTRODUCTION

Plant parasitic fungi sporulating in the anthers of their hosts evolved independently in several genera/species of two major phylogenetic basidiomycetous lineages, including the pucciniomycotinous genera *Bauerago* (Vánky 1999, 2012) and *Microbotryum* (Vánky 1998, 2012, Kemler *et al.* 2006, 2009), and the ustilaginomycotinous genera *Antherospora* (Bauer *et al.* 2008, Piątek *et al.* 2011, 2013), *Thecaphora* (Roets *et al.* 2008, 2012, Vánky & Lutz 2007) and *Urocystis* (Vánky 2012). The anther smuts of *Caryophyllaceae*, commonly referred to as the *Microbotryum violaceum* complex, form a monophyletic lineage within the genus *Microbotryum*. They are model organisms studied in many disciplines of fungal biology, for example, ecology (Thrall *et al.* 1993), genomics (Hood 2005, Yockteng *et al.* 2007), population studies (Lee 1981, Alexander & Antonovics 1995, Alexander *et al.* 1996), life cycle studies (Schäfer *et al.* 2010),

geographic distribution (Hood *et al.* 2010, Fontaine *et al.* 2013), phylogeography (Vercken *et al.* 2010), evolutionary history (López-Villavicencio *et al.* 2005, Refrégier *et al.* 2008), speciation (Devier *et al.* 2010, Gladioux *et al.* 2011), and phylogeny and systematics (Lutz *et al.* 2005, 2008, Denchev *et al.* 2009, Piątek *et al.* 2012, Kemler *et al.* 2013).

The assignment of organisms to the appropriate species is critically important in every biological discipline, however challenging as in cases of complexes of morphologically similar species. Delimitation of species within the *Microbotryum violaceum* complex is a good example where morphology (of spores) alone is inadequate. The vast majority of species and specimens within this complex have reticulate spores of similar size, with only a few exceptions from this general morphological pattern (Vánky 2004, 2012). The oldest available species name for anther smuts on *Caryophyllaceae*, that is, *Microbotryum violaceum* (syn. *Ustilago violacea*), has been usually assigned to morphologically similar specimens

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on diverse host plants worldwide. However, cross infection experiments (Zillig 1921a, Liro 1924) and molecular analyses (Bucheli *et al.* 2000, Freeman *et al.* 2002, Le Gac *et al.* 2007) have indicated that many of such specimens are biologically and genetically distinct.

In order to break the limitation of phenotype-based species identification, Lutz *et al.* (2005) provided a robust phylogenetic framework for species delimitation based on the nuclear ribosomal ITS region, which recently was proposed as barcode marker for *Fungi* (Schoch *et al.* 2012). ITS well resolves species boundaries in the *Microbotryum violaceum* complex, and the obtained resolution agrees well with that obtained with other phylogenetic markers (β -tub, γ -tub, Ef1 α , pheromone receptors pr-MatA1, pr-MatA2) (Le Gac *et al.* 2007, Refrégier *et al.* 2008, Devier *et al.* 2010). This phylogenetic framework has subsequently been improved by adding further species and specimens from diverse host plants and incorporating the nuclear LSU rDNA, combined with the ITS, as an additional phylogenetic marker (Lutz *et al.* 2008, Piątek *et al.* 2012). The resultant molecular phylogeny and genetic divergences between specimens on different hosts, together with ecological and, if available, morphological data confirm that multiple species are hidden within the *Microbotryum violaceum* morphotype, with most specific to single host species. ITS and LSU sequences are available for 18 out of 22 recognized *Microbotryum* species in the anthers of caryophyllaceous plants. One species (*Microbotryum savilei*) is not sequenced yet, and for three species (*M. carthusianorum*, *M. coronariae*, *M. dianthorum s. str.*) sequences are available for some nuclear DNA regions (β -tub, γ -tub, Ef1 α , ITS, pheromone receptors pr-MatA1, pr-MatA2) (Lutz *et al.* 2005, Le Gac *et al.* 2007, Refrégier *et al.* 2008, Denchev *et al.* 2009, Devier *et al.* 2010, Kemler *et al.* 2013), but not for the LSU. *Microbotryum violaceum s. str.* is currently restricted to *Silene nutans* and its taxonomy is stabilized by the sequenced neotype specimen (ITS and LSU) from material collected in Germany (Lutz *et al.* 2008). It is likely that many undescribed species of anther smuts remain to be discovered amongst the large number of specimens reported from different hosts worldwide, especially considering that anther smuts from 108 different caryophyllaceous hosts listed in recent smut world monograph (Vánky 2012) are still not analysed with molecular methods. The re-collection of fresh materials is desirable since many of herbarium materials are too old for effective isolation of DNA.

The anther smut on *Silene saxifraga* (incl. *S. hayekiana*, Tutin *et al.* 1993) reported from several European countries (Zillig 1921b, Zundel 1953, Scholz & Scholz 1988, Vánky 1994, 2012, Almaraz & Durrieu 1997, Zwetko & Blanz 2004, Lutz & Vánky 2009, as *Microbotryum violaceum*, *M. violaceum s. l.* or *Ustilago violacea*) is a putative distinct species. In the molecular studies of Lutz *et al.* (2005, 2008), the sequences from two specimens of the anther smut on *Silene saxifraga* (as *S. saxifraga* subsp. *hayekiana*) clustered together in a sister position to the lineage containing *Microbotryum silenes-inflatae* on *Silene maritima* and *S. vulgaris*, and *M. aff. violaceum* on *Lychnis flos-cuculi* (as *S. flos-cuculi*) and *S. dioica*, the latter smut now referred to as *M. coronariae*. The limited number of samples was the main reason why the new species for the anther smut on *Silene saxifraga* was not

described at that time. Additionally, only LM morphology was assessed, and SEM studies were not conducted for these two specimens.

The present study aims to resolve the specific status of the anther smut on *Silene saxifraga* using molecular phylogenetic analyses of concatenated ITS + LSU rDNA sequences as well as light and scanning electron microscope examination of specimens from several populations. A further aim is to discuss the phylogenetic position of the anther smut on *Silene saxifraga* within the lineage of anther smuts on *Caryophyllaceae*, and to expand the number of ITS and LSU sequences available for genetic analyses and comparisons.

MATERIALS AND METHODS

Host plant nomenclature, specimen sampling and documentation

In accordance with Tutin *et al.* (1993), and supported by molecular phylogenetic data (Kemler *et al.* 2013), the host species names *Silene saxifraga* and *S. hayekiana* are accepted as single species *Silene saxifraga*. All examined host plant specimens were assigned to this species.

This study is based on phylogenetical and/or morphological analyses of specimens of *Microbotryum* sp. on *Silene saxifraga* originating from nine populations in three main European high mountain ranges, namely the Alps, the Dinaric Alps, and the Pyrenees. Six specimens were freshly collected in the field, three were received from fungal herbaria, and three were found by screening the sheets with *Silene saxifraga* (eight sheets labelled as *S. hayekiana* – none infected, 40 sheets labelled as *S. saxifraga* – three infected) preserved in the phanerogamic Herbarium of the W. Szafer Institute of Botany, Polish Academy of Sciences, Kraków, Poland (KRAM). Additionally, the LSU and ITS + LSU, respectively, of two specimens of *Microbotryum coronariae* on *Lychnis flos-cuculi* were newly sequenced for phylogenetic analyses. The voucher specimens are deposited in KR-M, KRAM, KRAM F, TUB, and H.U.V. (Table 1). The latter abbreviation refers to the personal collection of Kálmán Vánky, “Herbarium *Ustilaginales* Vánky” currently held at his home (Gabriel-Biel-Straße 5, D-72076 Tübingen, Germany). Nomenclatural novelty was registered in MycoBank (www.MycoBank.org, Crous *et al.* 2004). The genotype concept follows the proposal of Chakrabarty (2010).

Morphological examination

Dried fungal spores of the investigated specimens were mounted in lactic acid, heated to boiling point, and then examined under a Nikon Eclipse 80i light microscope at a magnification of $\times 1000$, using Nomarski optics (DIC). Spores were measured using NIS-Elements BR 3.0 imaging software. The extreme measurements were adjusted to the nearest 0.5 μm . The spore size range, mean and standard deviation of 50 spore measurements from each specimen are shown in Table 1. The species description includes combined values from all measured specimens. LM micrographs were taken with a Nikon DS-Fi1 camera. The infected anthers of *Silene saxifraga* (KRAM F-49440), and the spore ornamentation in

specimens from different populations (H.U.V. 19570, KR-M-23890, KRAM 1760, KRAM F-49439, 49440, TUB 11790) were analysed using scanning electron microscopy (SEM). For this purpose, infected anthers and dry spores were mounted on carbon tabs and fixed to an aluminium stub with double-sided transparent tape. The tabs were sputter-coated with carbon using a Cressington sputter-coater and viewed with a Hitachi S-4700 scanning electron microscope, with a working distance of ca. 12 mm. SEM micrographs were taken in the Laboratory of Field Emission Scanning Electron Microscopy and Microanalysis at the Institute of Geological Sciences, Jagiellonian University, Kraków (Poland).

DNA extraction, PCR, and sequencing

The methods of isolation of fungal material, DNA extraction, amplification of the ITS 1 and ITS 2 regions of the rDNA including the 5.8S rDNA (ITS, about 690 bp) and the 5'-end of the nuclear large subunit ribosomal DNA (LSU, about 625 bp), purification of PCR products, sequencing, and processing of the raw data followed Lutz *et al.* (2004) and Piątek *et al.* (2012). DNA sequences determined for this study were deposited in GenBank. GenBank accession numbers are given in Table 1 and Fig. 1.

Phylogenetic analyses

The *Microbotryum* specimens examined in this study are listed in Table 1. To elucidate the phylogenetic position of the *Microbotryum* specimens on *Silene saxifraga* their concatenated ITS + LSU sequences were analysed within a dataset that covered all Caryophyllaceae anther smut species of which ITS and LSU sequences were available (Freeman *et al.* 2002, Lutz *et al.* 2005, 2008, Hood *et al.* 2010, Sloan *et al.* 2008, Piątek *et al.* 2012) or that were sequenced in the present study (*Microbotryum coronariae*, Table 1), comprising 19 of the 22 currently recognized species. For the final analysis the dataset was reduced to a maximum of two sequences per species. All sequences available in GenBank that clustered both within the *Microbotryum* sp. on *Silene saxifraga* clade and its sister clade (*M. coronariae*, *M. silenes-inflatae*) and which could not be assigned to any *Microbotryum* species (compare Fig. 1) were kept.

Sequence alignment was obtained using MAFFT 6.845b (Kato *et al.* 2002, Kato & Toh 2008) and the L-INS-I option. In the final alignment we retained conserved alignment positions using GBLOCKS (Castresana 2000) with the following options: 'Minimum Number of Sequences for a Conserved Position': 25, 'Minimum Number of Sequences for a Flank Position': 25, 'Maximum Number of Contiguous Non-conserved Positions': 8, 'Minimum Length of a Block': 5 and 'Allowed Gap Positions' to 'With half'.

The resulting alignment [new number of positions: 1276 (57% of the original 2232 positions) variable sites: 212] was used for phylogenetic analyses. Bayesian Analysis (BA) was performed using MrBayes 3.1.2 (Huelsenbeck & Ronquist 2001, Ronquist & Huelsenbeck 2003) applying the same settings as in Piątek *et al.* (2012). Four incrementally heated chains were run for 10,000,000 generations, sampled every 100th generation, thereby resulting in 100,001 trees of which the first 25,001 sampled trees were discarded. Maximum Likelihood (ML) was performed using RAxML 7.2.6

(Stamatakis 2006) via the raxmlGUI (Silvestro & Michalak 2012). We used the GTRGAMMA and rapid bootstrap option (Stamatakis *et al.* 2008). Trees were rooted with *Microbotryum scabiosae* following Kemler *et al.* (2006).

RESULTS

Morphology

The specimens on *Silene saxifraga* developed sori in all anthers of an inflorescence and within a plant clump most (but not all) flowers contained anthers with smut spores (Fig. 2). The smut sporulated inside the pollen sacs, which at first were completely covered by the anther's epidermis and later split longitudinally by the stomia revealing a dark brownish violet, powdery mass of spores. Pollen was not produced by infected anthers (Fig. 2). The spores in all specimens were reticulate under the light microscope, regular in shape and uniform in size within each collection, and highly uniform in shape, size range and average size between different collections (Fig. 3, Table 1). In scanning electron microscope, the spores were reticulate with variably ornamented interspaces. The interspaces usually ranged from almost smooth to rough or verruculose, but certain spores had more or less well-developed tuberculate warts on the interspaces or lower parts of the muri (Figs 3–4). The spores with tuberculate warts constituted a small but regular fraction of spores. Tuberculate warts were most apparent in the material from Kanzianiberg (Austria) that is designated here as holotype of the new species.

Phylogenetic analyses

For both the ITS and LSU, the sequences of the *Silene saxifraga* anther smut specimen from Montenegro (KRAM F-49440) differed in one position from the remaining sequences, which were identical among each other.

The different runs of the BA that were performed and the ML analyses yielded consistent topologies. To illustrate the results, the consensus tree of one run of the BA is presented (Fig. 1).

In all analyses, the known species were inferred with high support values except for *Microbotryum dianthorum* s. l., for which the sequences clustered in two different lineages. With high to moderate support in all analyses the sequences of anther smut specimens on *Silene saxifraga* clustered together, forming the sister lineage to *Microbotryum* sp. on *S. campanula*. That clade formed a monophylum with *M. coronariae*, *M. silenes-inflatae*, *M. sp.* on *S. ciliata*, and *M. violaceum* s. str. However the phylogenetic relations between those taxa received only low support values. Within the cluster of anther smut on *Silene saxifraga*, the specimen from Montenegro (KRAM F-49440) was revealed in a sister position to the remaining *S. saxifraga* anther smut specimens, which were identical among each other.

Considering groups that received considerable support in all analyses the phylogenetic relationships between the species inferred here were in contrast to the results discussed by Piątek *et al.* (2012) in two aspects: *Microbotryum violaceo-verrucosum* clustered as sister taxon to *M. heliospermae* and *M. lagerheimii*, and *M. saponariae* was revealed as sister taxon to the *Dianthus* and *Petrorhagia* anther smuts.

Table 1. List of examined *Microbotryum* specimens, with host plants, GenBank accession numbers, spore size range, mean spore sizes with standard deviation and reference specimens.

Species	Host plants	GenBank acc. no.	Spore size range (µm)	Mean spore size with standard deviation (µm)	Reference specimens ¹
<i>Microbotryum coronariae</i>	<i>Lychnis flos-cuculi</i>	ITS: KC684887 LSU: KC684886	Not analysed	Not analysed	Germany, Bayern, Allgäu, Ks. Oberallgäu, Oberjoch, Kematsriedmoos, Westteil, ca. 1150 m a.s.l., 25 Jun. 2008, <i>M. Scholler</i> , KR-M-23797
<i>Microbotryum coronariae</i>	<i>Lychnis flos-cuculi</i>	² ITS: AY877417 LSU: KC684885	Not analysed	Not analysed	Norway, Kristiansund, Farstad, 14 Aug. 2002, <i>M. Lutz</i> , TUB 012115
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	6.5–9.5(–10.5) × 6.0–8.5	7.6 ± 0.9 × 7.1 ± 0.6	Austria, Carinthia, Karawanken, 7 km WSW of Bad Eisenkappel, Trögern valley, 22 Jul. 1962, <i>H. Teppner</i> , KR-M-34470 (Dupla Graecensia Fungorum 237)
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: AY588102 LSU: JN000077	5.5–8.5(–9.5) × 5.0–6.5(–7.0)	6.7 ± 0.8 × 6.0 ± 0.5	Austria, Carinthia, Villach, Finkenstein, Kanzianiberg, 18 Jun. 2003, <i>M. Lutz</i> , TUB 11791
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: JN000073 LSU: JN000079	5.0–8.5 × 5.0–8.0	6.7 ± 0.8 × 6.2 ± 0.6	Austria, Carinthia, Villach, Finkenstein, nort of Kanzianiberg, 7 Jul. 2005, <i>M. Lutz</i> , KR-M-23889
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: JN000071 LSU: JN000075	5.0–8.0(–9.0) × (4.5–)5.0–7.5	6.6 ± 0.8 × 6.1 ± 0.8	Austria, Carinthia, Villach, Finkenstein, southern part of the Kanzianiberg, near the church, 24 Jun. 2006, <i>M. Lutz</i> , KR-M-23890 – holotype
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	(5.5–)6.0–7.5(–8.5) × 5.0–7.5	6.8 ± 0.6 × 6.2 ± 0.6	France, Central Pyrenees, rocks between Gavarnie village and Cirque de Gavarnie, 11 Jul. 1961, <i>S. Batko</i> , KRAM 1762
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: JN000074 LSU: JN000078	5.5–7.5 × 5.0–6.5 (–7.0)	6.4 ± 0.5 × 5.8 ± 0.5	Germany, Baden-Württemberg, Tübingen, Botanical Garden, cultivated (originating from Slovenia, Bovec, Vas na Skali, 17 Jul. 1994), 11 Jun. 1999, <i>C. Vánky</i> & <i>K. Vánky</i> , H.U.V. 19570
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	6.0–7.5 × 5.5–7.5	6.7 ± 0.4 × 6.3 ± 0.5	Germany, Baden-Württemberg, Tübingen, Botanical Garden, cultivated (originating from Slovenia, Bovec, Vas na Skali, 17 Jul. 1994), 24 May 2011, <i>M. Lutz</i> , KRAM F-49439
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	6.5–8.5(–9.5) × 6.0–8.0	7.4 ± 0.7 × 6.9 ± 0.5	Italy, Tridentum, Doss Trento, 25 May 1893, <i>Evers</i> , KRAM 1760
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	6.0–8.0(–8.5) × 6.0–7.5	7.0 ± 0.5 × 6.5 ± 0.5	Italy, Alpi Maritime, Valle de Gosso, 7 Jun. 1992, <i>M. Schubert</i> , KR-M-23949
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: JN000072 LSU: JN000080	6.0–8.5(–9.5) × 5.5–8.5(–9.0)	7.0 ± 0.8 × 6.5 ± 0.8	Montenegro, Dinaric Alps, Durmitor Mts, along trail Sedlo-Bobotov Kuk, Surutka valley, 10 Aug. 2009, <i>A. Ronikier</i> & <i>M. Ronikier</i> , KRAM F-49440
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	–	6.5–8.5(–9.5) × (5.5–)6.0–7.5(–9.0)	7.3 ± 0.7 × 6.8 ± 0.6	Slovenia, Carniola, "Schibeneggergraben bei Ratschach", 3 Jun. 1885, <i>J.C. Eques Pittoni a Dannenfeldt</i> , KRAM 108297
<i>Microbotryum silenes-saxifragae</i>	<i>Silene saxifraga</i>	ITS: AY588101 LSU: JN000076	5.5–7.5 × (4.5–)5.0– 6.5(–7.0)	6.7 ± 0.5 × 6.2 ± 0.5	Slovenia, Bovec, Trenta, Juliana Alpine Botanical Garden, cultivated, 7 Aug. 2001, <i>D. Begerow</i> & <i>M. Lutz</i> , TUB 11790

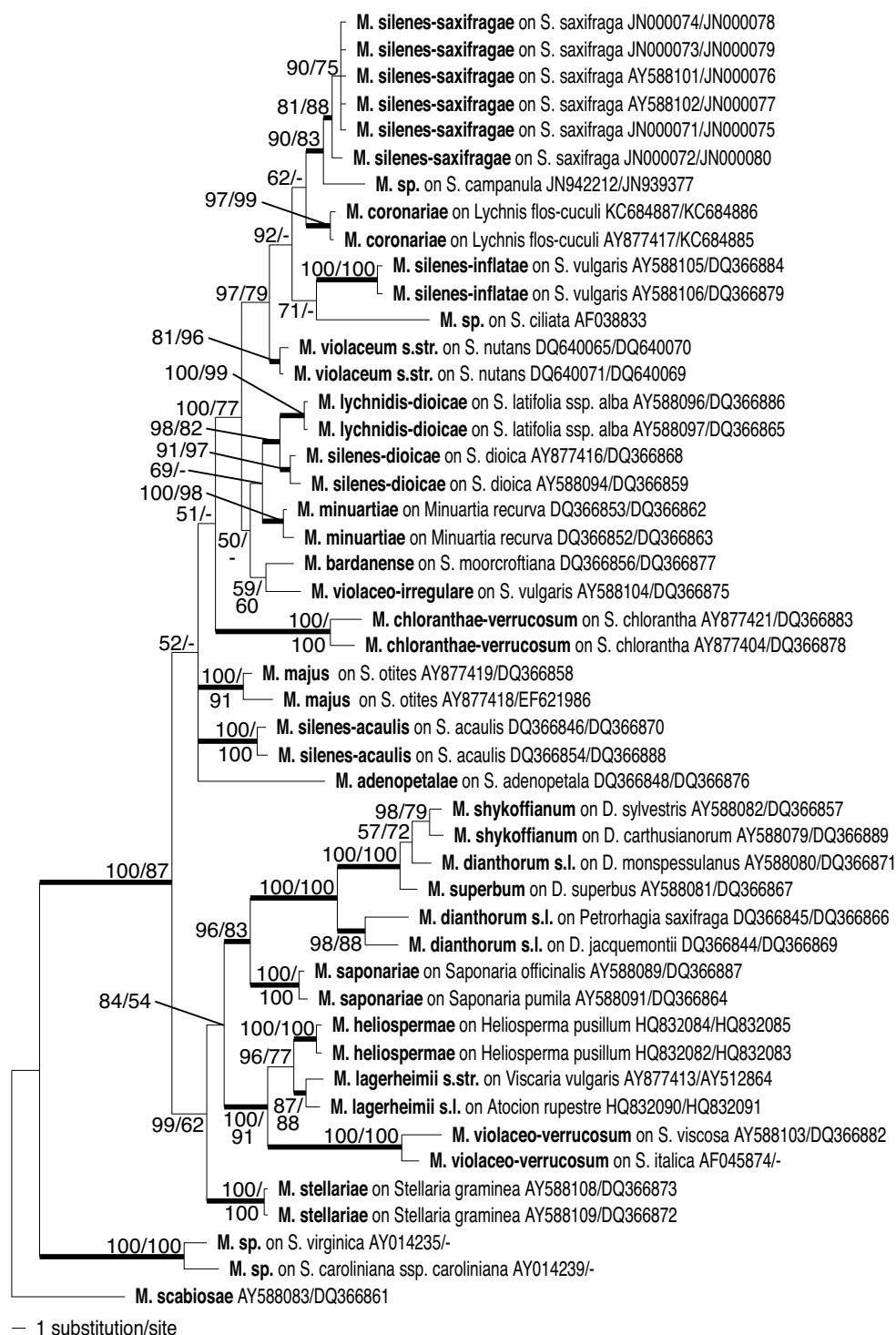


Fig. 1. Bayesian inference of phylogenetic relationships between the sampled *Microbotryum* species: Markov chain Monte Carlo analysis of an alignment of concatenated ITS + LSU base sequences using the GTR+I+G model of DNA substitution with gamma distributed substitution rates and estimation of invariant sites, random starting trees and default starting parameters of the DNA substitution model. A 50% majority-rule consensus tree is shown computed from 75 000 trees that were sampled after the process had reached stationarity. The topology was rooted with *Microbotryum scabiosae*. Bold branches indicate support values higher than 80 in all analyses. Numbers on branches before slashes are estimates for a *posteriori* probabilities; numbers on branches after slashes are ML bootstrap support values. Branch lengths were averaged over the sampled trees. They are scaled in terms of expected numbers of nucleotide substitutions per site. D. = *Dianthus*, M. = *Microbotryum*, S. = *Silene*.

¹H.U.V. – Herbarium *Ustilaginales* Vánky, Gabriel-Biel-Str. 5, D-72076 Tübingen, Germany; KR-M – Mycological Herbarium of the Staatliches Museum für Naturkunde Karlsruhe, Germany; KRAM – Phanerogamic Herbarium of the W. Szafer Institute of Botany, Polish Academy of Sciences, Kraków, Poland; KRAM F – Mycological Herbarium of the W. Szafer Institute of Botany, Polish Academy of Sciences, Kraków, Poland; TUB – Herbarium of the Eberhard-Karls-Universität Tübingen, Germany.

²Taken from Lutz *et al.* (2005).

TAXONOMY

Microbotryum silenes-saxifragae M. Lutz, M. Piątek & Kemler, **sp. nov.**
 MycoBank MB800823
 (Figs 2–4)

Etymology: The name of the species refers to the host plant species, *Silene saxifraga*.

Description: Parasitic on *Silene saxifraga*. Sori in anthers; all anthers of the inflorescence infected, and most flowers in a clump contain smut spores; spore mass powdery, dark brownish violet. Spores pale violaceous or violaceous in transmitted light, regular in shape and size, globose, subglobose, broadly ellipsoidal, or rarely ovoid, 5.0–8.5 (–10.5) × (4.5–)5.0–7.5(–9.0) μm; wall reticulate, ca. 0.5–0.7 μm high, meshes more or less polyhedral, usually irregular, rarely regular, 5–8 (usually 6–7) meshes per spore diameter, interspaces usually smooth as observed by LM (sometimes weak tubercles are visible at very high magnification using Nomarski optics), almost smooth, rough, verruculose or tuberculate as observed by SEM.

Type: **Austria:** *Carinthia:* Villach, Finkenstein, southern part of the Kanzianiberg, near the church, [630 m a.s.l.], on *Silene saxifraga*, 24 June 2006, M. Lutz (KR-M-23890 – **holotype**). The ITS/LSU hologenotype sequences are deposited in GenBank as JN000071/JN000075, respectively.

Additional specimens examined (paratypes): **Austria:** *Carinthia:* Karawanken, 7 km WSW of Bad Eisenkappel, Trögern valley, 700–800 m a.s.l., on *Silene saxifraga*, 22 July 1962, H. Teppner (KR-M-34470, Dupla Graecensia Fungorum 237); Villach, Finkenstein, Kanzianiberg, [630 m a.s.l.], on *Silene saxifraga*, 18 June 2003, M. Lutz (TUB 11791); Villach, Finkenstein, north of Kanzianiberg, [630 m a.s.l.], on *Silene saxifraga*, 7 July 2005, M. Lutz (KR-M-23889); **France:** *Central Pyrenees:* rocks between Gavarnie village and Cirque de Gavarnie, [between 1400 and 1600 m a.s.l.], on *Silene saxifraga*, 11 July 1961, S. Batko (KRAM 1762). – **Germany:** *Baden-Württemberg:* Tübingen, Botanical Garden (originating from Slovenia, Bovec, Vas na Skali, 17 July 1994), [440 m a.s.l.], on cultivated *Silene saxifraga*, 11 June 1999, C. Vánky & K. Vánky (H.U.V. 19570); Tübingen, Botanical Garden (originating from Slovenia, Bovec, Vas na Skali, 17 July 1994), [440 m a.s.l.], on cultivated *Silene saxifraga*, 24 May 2011, M. Lutz (KRAM F-49439). – **Italy:** *Tridentum:* Doss Trento, [300 m a.s.l.], on *Silene saxifraga*, 25 May 1893, Evers (KRAM 1760); *Alpi Maritime:* Valle de Gosso, [? – site not located in Google Earth], on *Silene saxifraga*, 7 June 1992, M. Schubert (KR-M-23949). – **Montenegro:** *Dinaric Alps:* Durmitor Mts, along trail Sedlo-Bobotov Kuk, Surutka valley, ca. 2090 m a.s.l., on *Silene saxifraga*, 10 August 2009, A. Ronikier & M. Ronikier (KRAM F-49440). – **Slovenia:** *Carniola:* “Schibeneggergraben bei Ratschach”, [Ratschach - 870 m a.s.l., Schibeneggergraben not located in Google Earth], on *Silene saxifraga*, 3 June 1885, J.C. Eques Pittoni a Dannenfeldt (KRAM 108297); Bovec, Trenta, Juliana Alpine Botanical Garden, [600 m a.s.l.], on cultivated *Silene saxifraga*, 7 August 2001, D. Begerow & M. Lutz (TUB 11790).

Host range and distribution: On *Silene saxifraga* (*Caryophyllaceae* subfam. *Silenoideae*); Europe (Austria,

France, Germany, Italy, Montenegro, Slovenia). The localities in Austria, Italy and Slovenia are placed in the Alps, the locality in France in the Pyrenees and the locality in Montenegro in the Dinaric Alps. The locality in Germany is artificial as plants were cultivated.

Ecology: The infected plants were found from May to August in the natural localities, in May and June cultivated in the Botanical Garden in Tübingen (Germany), and in August cultivated in the Juliana Alpine Botanical Garden in Trenta (Slovenia). At the type locality, the Kanzianiberg (Austria), large populations of infected plants were observed between 2003 and 2006 in different places on the limestone hill. In other localities for which data on the habitat are available, *Microbotryum silenes-saxifragae* occurred on plants growing on rocks, together with *Rhamnus pumila* (France: between Gavarnie and Cirque de Gavarnie), between limestone rocks (Austria: Karawanken) and on grassland (Montenegro: Surutka valley). Altitude data were available for few of the examined specimens, and the approximate altitude data (included in square brackets in the list of specimens examined) for most of the remaining specimens were obtained through Google Earth (google.earth.com). The lowest locality was recorded at 300 m a.s.l., and the highest locality at 2090 m a.s.l., which indicates that *Microbotryum silenes-saxifragae* is a submontane-montane species with wide altitudinal amplitude.

DISCUSSION

Silene saxifraga is widely distributed on rocks and screes in southern European mountains, extending northwards to West Austria (Tutin *et al.* 1993). The anther smut of *Silene saxifraga*, although reported in the literature and referred to the catch-all name *Microbotryum violaceum* (syn. *Ustilago violacea*), is poorly known and has never been critically examined using samples from different populations. In this study, molecular phylogenetic analyses and morphological data were used to resolve the systematic position of the anther smut on *Silene saxifraga*.

The phylogenetic analyses of the concatenated ITS + LSU dataset showed that the analysed specimens on *Silene saxifraga* from three geographically distinct populations (Austria, Slovenia, and Montenegro; the German specimen derived from a Slovenian population) form a well supported independent evolutionary lineage of caryophyllaceous anther smuts. The genetic divergence of the lineage is comparable to genetic distances between the remaining anther smuts on caryophyllaceous hosts (Lutz *et al.* 2005, 2008, Piątek *et al.* 2012) or between *Microbotryum* species on non-caryophyllaceous hosts (Kemler *et al.* 2006, 2009). Within this lineage, the anther smut specimen on *Silene saxifraga* from Montenegro (Dinaric Alps) revealed some sequence divergence in comparison to the specimens from populations in Austria and Slovenia (Alps). A similar phylogeographic split between specimen from the Alps and specimens from the Carpathians and the Dinaric Alps has been observed in another montane species, namely *Microbotryum heliospermae*, a parasite on *Heliosperma pusillum* (Piątek *et al.* 2012). This is congruent with a similar phylogeographical pattern observed in

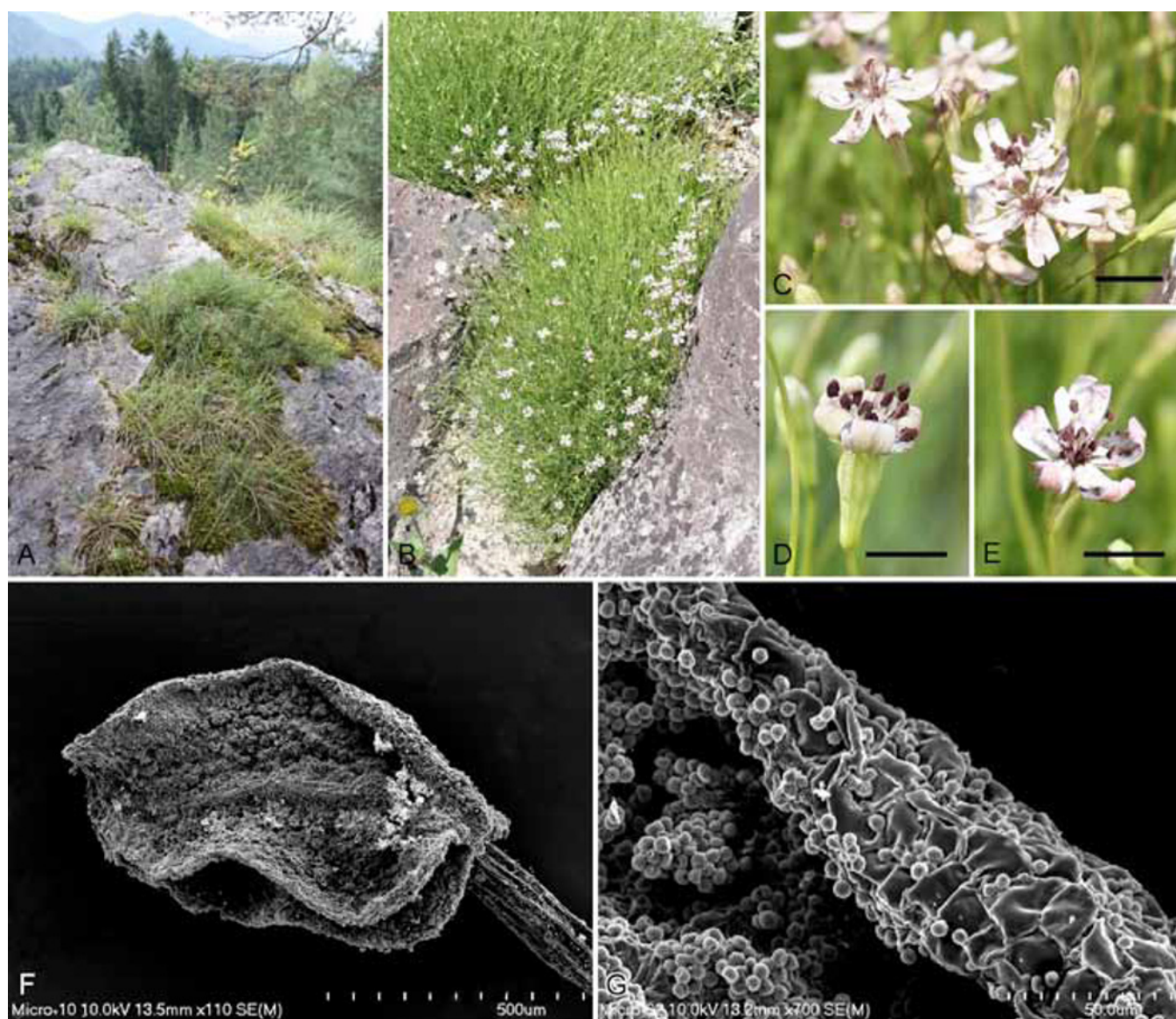


Fig. 2. *Microbotryum silenes-saxifragae* sp. nov. on *Silene saxifraga*. **A.** The type locality at the Kanzianiberg, Austria. **B.** The clump of *Silene saxifraga* with infected flowers in the Botanical Garden of Tübingen, Germany. **C–E.** Infected inflorescences, with the fungus sporulating in the anthers in the Botanical Garden of Tübingen, Germany. **F.** Infected anther: an open pollen sac filled with teliospores seen at the foreground, made in SEM (KRAM F-49440). **G.** Teliospores inside the pollen sac and the anther's epidermis, seen by SEM. (KRAM F-49440). Bars: C–E = 5 mm, F = 500 μ m, G = 50 μ m.

several vascular plant species (Ronikier 2011). It may indicate long-term separation and different evolutionary histories of populations in the Alps and the Dinaric Alps (and South-East European mountains in general).

The closest phylogenetic relative of the analysed anther smut specimens on *Silene saxifraga* may be the anther smut on *Silene campanula* assigned by Schoch *et al.* (2012) to *Microbotryum violaceum*¹. However, it does not belong to this species, which is restricted to *Silene nutans*. According to our molecular phylogenetic analyses, the anther smut on *Silene campanula* occupies a sister position to the anther smut on *Silene saxifraga*, well separated from *Microbotryum violaceum* s. str. (incl. neogenotype ITS/LSU sequences DQ640065/640070, Lutz *et al.* 2008).

The morphological examination of the anther smut specimens on *Silene saxifraga* revealed some characteristics,

such as a reticulate ornamentation of spores, spore shape and size, that are shared with other *Microbotryum* species in anthers of caryophyllaceous hosts (Lutz *et al.* 2005, 2008, Denchev *et al.* 2009, Piątek *et al.* 2012, Vánky 2012). However, in contrast to other species of the caryophyllaceous anther smuts lineage, the interspaces in spores (= bottom of the muri) were variably ornamented from almost smooth to rough, verruculose or, in some spores, distinctly tuberculate. This feature is almost invisible in LM (sometimes weak tubercles are visible at very high magnification using Nomarski optics),

¹see: <http://www.fungalbarcoding.org/BioMICS.aspx?Table=Fungal%20barcodes&Fields=All&Rec=2110> -- data retrieved on 1 March 2013; the specimen is deposited in private herbarium and it was not possible to re-examine it during the course of this study.

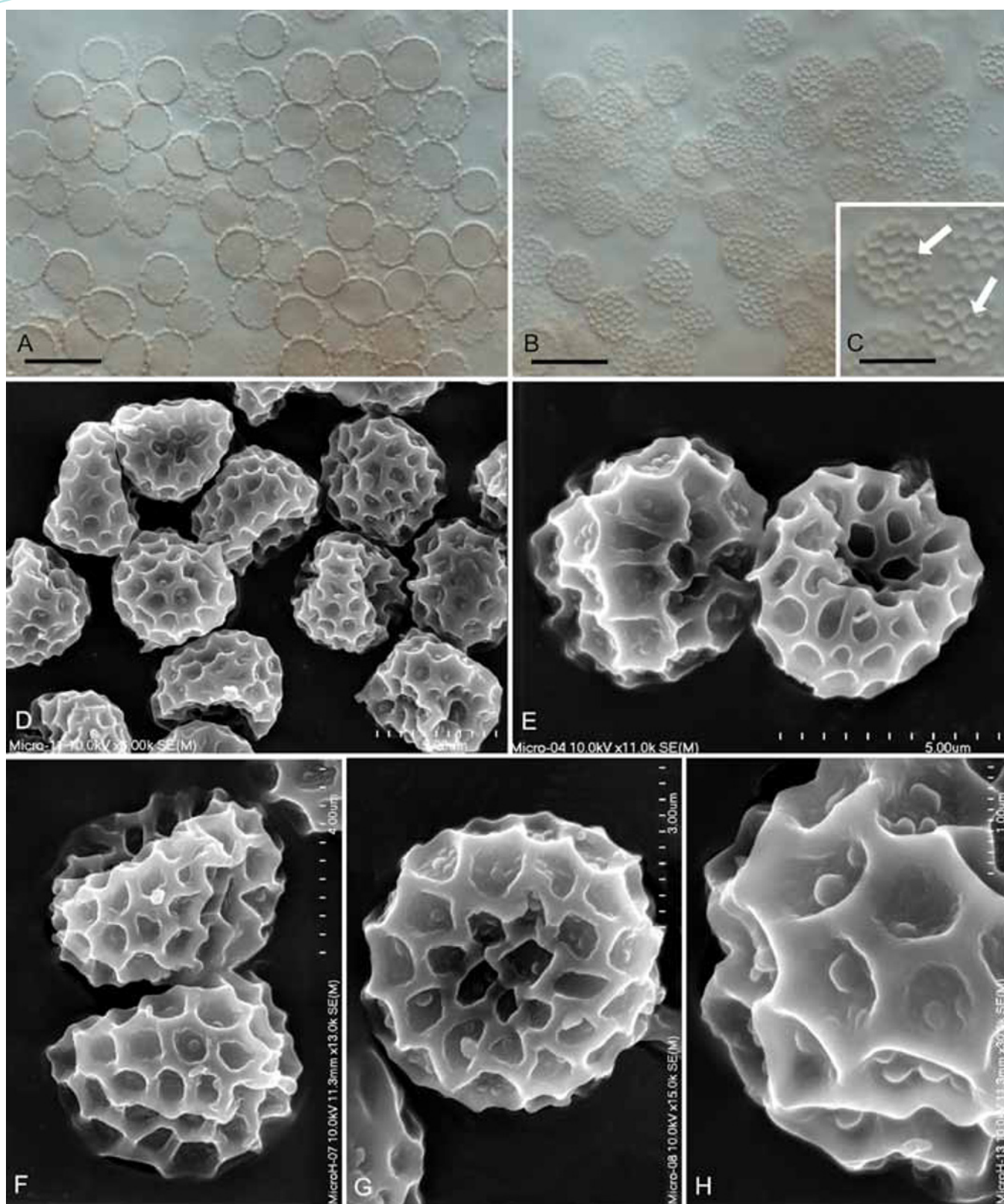


Fig. 3. *Microbotryum silenes-saxifragae* sp. nov. on *Silene saxifraga* (KR-M-23890 – holotype). **A–B.** Spores seen by LM, median and superficial views. **C.** Hardly visible tubercles in LM at very high magnification using Nomarski optics, indicated by arrows. **D–G.** Spores with tuberculate, rough and verruculose interspaces seen by SEM. **H.** Close-up of spore ornamentation seen by SEM. Bars: A–B = 10 μ m, C–E = 5 μ m, F = 4 μ m, G = 3 μ m, H = 1 μ m.

but visible in SEM. The reason for morphological variation in different specimens is uncertain, but it might be related to different developmental stages or environmental conditions.

In conclusion, the genetic divergence, the host plant and the distinct spore morphology indicate that the anther smut on *Silene saxifraga* represents a new species that is

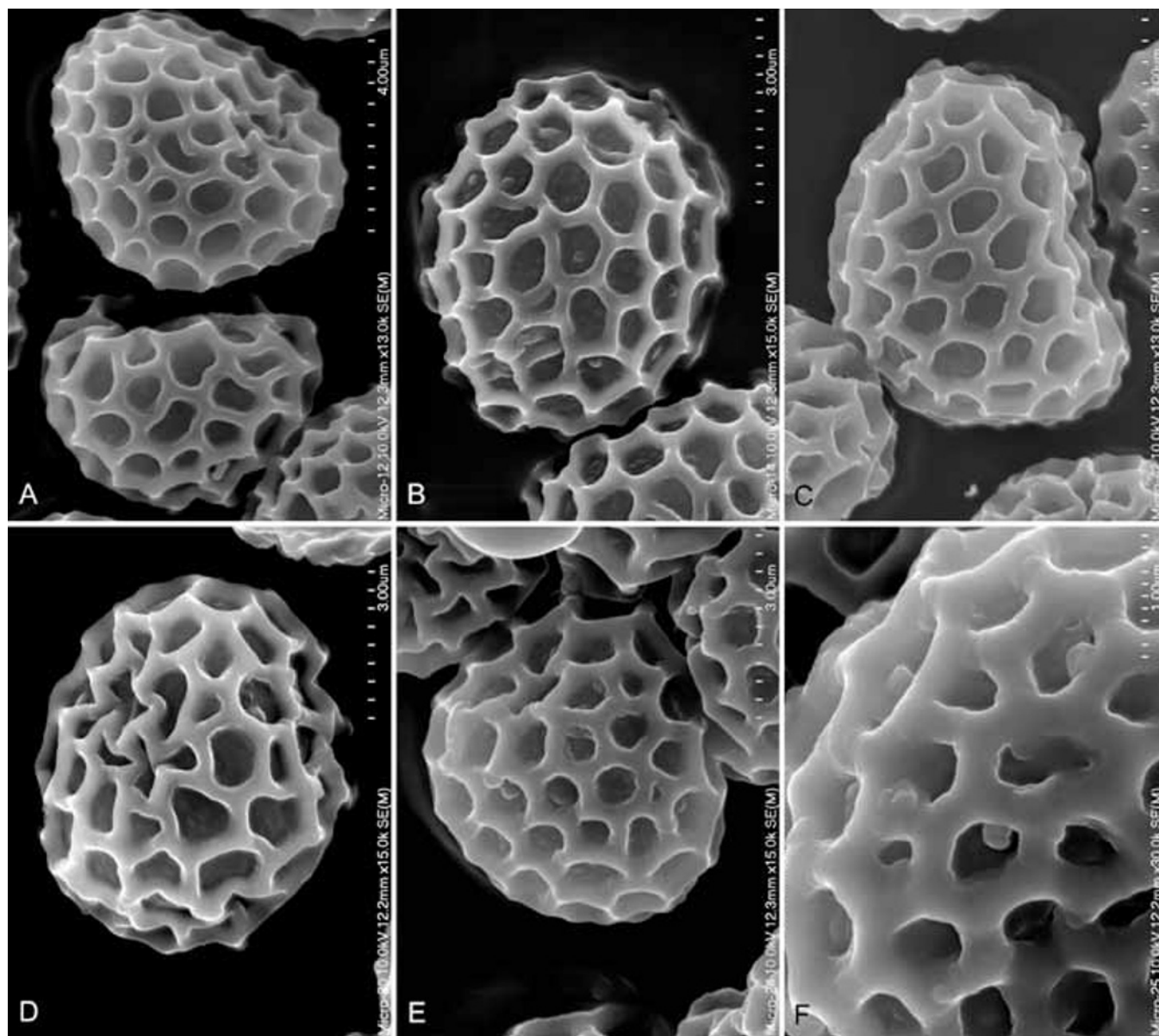


Fig. 4. Variability of interspaces ornamentation in different spores and specimens of *Microbotryum silenes-saxifragae* sp. nov. seen by SEM. **A–B.** From KRAM F-49439. **C–D.** From KRAM F-49440. **E–F.** From KRAM 1760. Bars: A, C = 4 μ m, B, D–E = 3 μ m, F = 1 μ m.

described in this work as *Microbotryum silenes-saxifragae*. The genetic divergence and only one sequenced specimen of the anther smut on *Silene campanula* do not allow the possible assignment of this specimen to *Microbotryum silenes-saxifragae*.

The distinct morphological trait of tuberculate interspaces observed in a certain portion of reticulate spores of *Microbotryum silenes-saxifragae* is unique in the lineage of caryophyllaceous anther smuts. Most species of this clade have spores with reticulate ornamentation but with smooth interspaces (Lutz *et al.* 2005, 2008, Piątek *et al.* 2012, Vánky 2012), exceptionally rough or at most verruculose interspaces (*Microbotryum adenopetalae*, Lutz *et al.* 2008, *M. carthusianorum*, *M. shykoffianum*, Denchev *et al.* 2009). There are only four evident exceptions from this general pattern, namely two species have verruculose spores (*Microbotryum chloranthae-verrucosum*, *M. violaceo-verrucosum*, Lutz *et al.* 2005, Vánky 2012) and two species have irregularly

verruculose-reticulate spores (*M. bardanense*, *M. violaceo-irregulare*, Deml & Oberwinkler 1983, Chlebicki & Suková 2005, Vánky 2012). *Microbotryum silenes-saxifragae* adds a new type of ornamentation to the lineage of caryophyllaceous anther smuts.

The development of reticulate spores having tuberculate interspaces is a common feature in other *Microbotryum* species, and it is especially widespread in many species infecting flowers of different plants from the family *Polygonaceae* that cluster in *Microbotryum* group II resolved in the phylogenetic study of Kemler *et al.* (2006). The reticulate spores with tuberculate interspaces are also produced by some anther smuts on non-caryophyllaceous hosts (Vánky 2012) and are developed in some *Microbotryum* species sporulating in the ovaries of caryophyllaceous plants (Piątek 2005, Vánky 2012). Most of these species cluster in *Microbotryum* group II (Kemler *et al.* 2006), though not for all of them sequence data are available. It seems that the feature of reticulate spores

with tuberculate interspaces is a homoplasy that evolved convergently in different *Microbotryum* species.

Microbotryum silenes-saxifragae is embedded in a well supported clade composed of three other described species, *M. coronariae* on *Lychnis flos-cuculi*, *M. silenes-inflatae* on *S. vulgaris* and *M. violaceum* s. str. on *Silene nutans*, as well as the unresolved *Microbotryum* sp. on *Silene campanula* and *Microbotryum* sp. on *S. ciliata*. In accordance with the studies of Lutz *et al.* (2005, 2008), where only two sequences of anther smut on *Silene saxifraga* were available, *Microbotryum violaceum* s. str. is revealed as sister taxon to all other clades of this group. All other phylogenetic relations received only moderate to low support with one exception, the sister relation of *Microbotryum* sp. on *Silene campanula* and *M. silenes-saxifragae*. The host plant phylogenetic relations and ecology indicate a complex evolutionary history within this clade of caryophyllaceous anther smuts. While *Silene campanula*, *S. nutans* and *S. saxifraga* are phylogenetically closely related (Greenberg & Donoghue 2011, Kemler *et al.* 2013), *S. ciliata* and especially *Lychnis flos-cuculi* and *Silene vulgaris* have distant phylogenetic relations (Greenberg & Donoghue 2011, Kemler *et al.* 2013). Moreover, the host plants occupy different ecological niches, *Silene campanula*, *S. ciliata*, *S. nutans* and *S. saxifraga* occur on calcareous rocks, mostly in mountains, *Lychnis flos-cuculi* on wet meadows, in mountains and lowlands, and *S. vulgaris* on dry meadows or disturbed places (roadsides, fields) in mountains and lowlands. It appears possible that the evolution of *Microbotryum* species within this clade was driven by host shifts combined with the changes in the ecological niches of the hosts. Host shifts, both evolutionary ancient (Refrégier *et al.* 2008) or recent (Antonovics *et al.* 2002, López-Villavicencio *et al.* 2005, Kummer 2010), were often reported in caryophyllaceous anther smuts. Furthermore, artificial cross-inoculation experiments indicate that the potential host range of some *Microbotryum* species is larger than the actual host range observed in natural populations, but the ability to infect non-host plants is higher for plants that are phylogenetically closer related to the original host (de Vienne *et al.* 2009). This phenomenon could promote higher speciation rates in this group of plant parasites. In this respect, the clade warrants further study with increased sampling of anther smut specimens, especially on other hosts, likely to reveal other closely related species.

The lineage of caryophyllaceous anther smuts is predominantly composed of cryptic species, with only a few species differing morphologically (Lutz *et al.* 2005, 2008). In some species (*Microbotryum adenopetalae*, *M. heliospermae*, *M. minuartiae*) subtle morphological features were found after their initial detection by molecular methods (Lutz *et al.* 2008, Piątek *et al.* 2012). In the absence of molecular support or cross infection experiments, these features could easily be considered as phenotypic variation within a single species. In contrast, the tuberculate interspaces observed in a certain portion of reticulate spores of *Microbotryum silenes-saxifragae* are rather evident, although not previously noticed (probably due to the absence of SEM studies of anther smut specimens on *Silene saxifraga*). Species that have only been recognized as morphologically distinct after application of methods other

than comparative morphology, usually methods of molecular biology, are called pseudo-cryptic species in different studies on systematically diverse organisms (Knowlton 1993, Amato & Montresor 2008, Luttikhuisen & Dekker 2010, Medina *et al.* 2012). The species mentioned in this paragraph, including *Microbotryum silenes-saxifragae*, match that concept very well.

The discovery of morphological species showing genetic differences that are comparable with genetic divergences between cryptic species embedded within the lineage of caryophyllaceous anther smuts supports the narrow species concept for this group of smut fungi advocated by Liro (1924) and confirmed by recent molecular studies (Lutz *et al.* 2005, 2008, Le Gac *et al.* 2007, Refrégier *et al.* 2008, Devier *et al.* 2010, Piątek *et al.* 2012). Furthermore, the strict correlation of *Microbotryum silenes-saxifragae* with its host species confirms the earlier conclusion that host-specific species delimitation might reflect the evolution of many anther smut parasites best (Lutz *et al.* 2005). It is likely that the uncharted species diversity of anther smuts is much higher and that, in addition to cryptic species, also pseudo-cryptic species still could be detected as forming part of caryophyllaceous anther smuts lineage.

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