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A new corticolous Megaspora (Megasporaceae) species from Armenia

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Abstract: The corticolous species *Megaspora cretacea* is described as new for science. The species is characterized by a thick, cretaceous thallus and a pale bluish, rather coarse soredia covering most of the thallus. It grows on *Juniperus* bark in open arid woodlands in Armenia. A key to the three species included in the genus *Megaspora* is presented. Phylogenetic analysis based on nrITS sequences revealed that *M. cretacea* clustered within the *Megaspora* clade as sister species to *M. rimisorediata* with high support.

Key words: lichens, *Megasporaceae*, *Megaspora*, taxonomy, new species, sorediate, *Juniperus*, South Caucasus, Armenia, Khosrov Forest State Reserve, ITS

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Introduction

According to recent phylogenetic studies, Megasporaceae Lumbsch is monophyletic (Nordin & al. 2010). They are mostly saxicolous crustose lichens (Valadbeigi & al. 2011). In Armenia, they are among the more common lichen families, in species diversity but especially in abundance, covering large parts of most siliceous rock faces and also present on limestone. The genus Megaspora (Clauzade & Cl. Roux) Hafellner & V. Wirth is closely related to the genus Circinaria Link (Nordin & al. 2010). It is an exception within the family, in that it is predominantly corticolous, with two species on trees, one of which is also occasionally terricolous. Both currently accepted species, M. rimisorediata Valadbeigi & A. Nordin and M. verrucosa (Ach.) Hafellner & V. Wirth (Valadbeigi & al. 2011), occur in Armenia (Gasparyan & Sipman 2013; Harutyunyan & al. 2011).

During a lichenological excursion to Armenia, organized by the second author, we collected a sorediate crustose lichen at the bases of trees of *Juniperus polycarpos* K. Koch in the Khosrov Forest State Reserve. The territory of the Reserve was already considered as a protected area in the fourth century C.E. by the Armenian king Khosrov Kotak (Khanjyan 2004). In 1958, the Khosrov Forest was officially declared as a state reserve (Anonymous 2008). The natural landscapes of phryganoid vegetation, open arid forests and montane steppes have high biological diversity and are recognized as a priority area for conservation. So far, 1849 species of vascular plants (including 24 endemic species) and 176 lichenized and lichenicolous fungi have been registered in the reserve (Anonymous 2008; Gasparyan & al. 2015).

While in the field it was not possible to recognize the collected specimens as representatives of *Megasporaceae*; rather they gave the impression of a species of the

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Caloplaca albolutescens (Nyl.) H. Olivier / C. teicholyta (Ach.) J. Steiner group or, less likely, a species of Lepraria Ach., but subsequent examination of the material revealed a few black apothecia immersed in the thallus, with large, thin-walled ascospores and a greenish epihymenium, suggesting Megasporaceae.

In the framework of a phylogenetic study of Asian *Megasporaceae*, the first author sequenced the material and found that it clusters inside *Megaspora* as a sister species to *M. rimisorediata*. Therefore, we describe it as a new species in this genus.

Megaspora rimisorediata has a restricted distribution. It was described from Iran (Valadbeigi & al. 2011) and later found also in S Armenia (Gasparyan & Sipman 2013; Gasparyan & al. 2015). Megaspora verrucosa has been reported from Europe, Africa, Asia, North and South America, New Zealand and Antarctica (Smith & al. 2009).

Currently, Armenia is the centre of diversity of the genus, with all three currently known species present. The new species has been reported from two localities. Further comprehensive studies are required to explore distributional and ecological patterns of the new species.

Material and methods

Identification and descriptive work was carried out in Soest and BGBM using an Olympus SZX7 stereomicroscope and an Olympus BX50 compound microscope with interference contrast, connected to a Nikon Coolpix digital camera. Sections were mounted in tap water, in which also all measurements were taken. The specimens from this study are preserved in ABL and B (herbarium codes after Thiers 2016+). The chemistry of the type specimen was investigated by thin-layer chromatography (TLC) using solvent A (Orange & al. 2001).

DNA extraction — We used nuclear ITS1-5.8S-ITS2 rDNA sequences of specimens in the molecular study because it has been shown that among the regions of the ribosomal cistron, the internal transcribed spacer (ITS) region has the highest probability of successful identification for a range of fungi (Schoch & al. 2012; Divakar & al. 2015). Total DNA was extracted from freshly collected material according to Park & al. (2014). We followed the instructions given in that paper except for the following steps: we used a 1 \times 1 mm² piece of medulla and mixed it with beadbeader without liquid nitrogen; instead of chloroform we used Roti®-C/I (chloroform/isoamyl alcohol at a ratio of 24:1); and at the end we used only 30 μ L TE buffer instead of 100 μ L because of the low quantity of DNA.

PCR amplifications and sequencing — The primer pair ITS1F (Gardes & Bruns 1993) and ITS4 (White & al. 1990) was used for the PCR amplifications. PCR amplifications were performed in a 12.5 μL volume contain-

ing 2 μL undiluted DNA, 0.5 μL of each primer (10 mM), 6.4 μL of sterile water, 1 μL dNTP (2 mM), 1 μL s-buffer, 1 μL MgCl2, 0.1 μL Taq-polymerase. Thermal cycling parameters were initial denaturation for 5 min at 95 °C, followed by 30 cycles of 30 sec at 95 °C, 30 sec at 54 °C, and 1 min at 72 °C; following the last cycle a final extensions for 3 min at 72 °C was included. Amplification product was viewed by electrophoresis on 1% agarose gels and stained with ethidiumboromide and was purified by adding 2 μL ExoSAP-ITTM (Exonuclease 1-shrimp alkaline phosphatase) to 5 µL of the PCR products, followed by a heat treatment of 15 min at 37 °C and 15 min at 80 °C. The PCR product was sequenced in both directions by Bik-F Laboratory in Frankfurt am Main. For the reconstruction of a phylogenetic tree, all ITS sequences of Megasporaceae from Valadbeigi & al. (2011) were used as well as seven accessible sequences of Megaspora from NCBI GenBank (http://www.ncbi.nlm.nih.gov/genbank/). Two sequences were obtained from the new species and submitted to the NCBI GenBank (Table 1). The sequences were aligned through the Muscle V4 program web server (Edgar 2004) with the default settings. The aligned sequences were adjusted manually in PhyDE software (Müller & al. 2010). Gblocks 0.91b (http://molevol.cmima.csic.es/castresana/ Gblocks server.html) was used to eliminate ambiguously aligned positions, applying settings allowing for smaller final blocks, gap position within the final blocks and less strict flanking position (Castresana 2000).

Phylogenetic analyses — MrModeltest (Nylander 2004) was used to determine the most appropriate model using AIC, with GTR + I + G found to be the best-fitting model of nucleotide evolution. Bayesian inference of phylogeny with Markov chain Monte Carlo sampling was performed on the Bayesian inference of phylogeny with Markov chain Monte Carlo sampling was performed on the 477 unambiguously aligned nucleotide positions. Bayesian analyses were conducted with Mr-Bayes v. 3.2.2 (Ronquist & Huelsenbeck 2003) using the GTR model of nucleotide substitution including a proportion of invariable sites and a discrete gamma distribution with six rate categories. Two independent runs, each with four Metropolis-Coupled Markov Chain Monte Carlo chains and a temperature of 0.2 were initiated and run for 1000000 generations, with tree and parameter sampling every 100 generations. Burn-in was set to discard 25 % of samples. Maximum parsimonious trees (MPs) were reconstructed in PAUP (Phylogenetic Analysis Using Parsimony) v. 4.0b10 (Swofford 2003) using the heuristic search option with 100 random sequence additions and tree bisection and reconstruction (TBR) as the branch-swapping algorithm. Alignment gaps were treated as missing and all characters were unordered and of equal weight. The robustness of the trees obtained was evaluated by 1000 bootstrap replications with ten random sequence additions. Molecular Evolutionary Genetics Analysis software (MEGA verWilldenowia 46 – 2016 247

Table 1. Voucher specimens and NCBI GenBank accession numbers of the ITS sequences used in the phylogenetic analyses.

Taxon	Locality, voucher	GenBank acc. no.	
Aspicilia cinerea	Sweden, Dalarna, <i>Hermansson 13275</i> (UPS)	EU057899	
Aspicilia indissimilis	Sweden, Nordin 5943 (UPS)	EU057909	
Aspicilia laevata	Sweden, Tibell 23659 (UPS)	EU057910	
Circinaria calcarea	Sweden, Nordin 5888 (UPS)	EU057898	
Circinaria contorta	Sweden, Nordin 5895 (UPS)	EU057900	
Circinaria leprosescens	Sweden, Nordin 5906 (UPS)	EU057911	
Lobothallia melanaspis	Norway, Own-Larsson 8943a (UPS)	JF825524	
Lobothallia radiosa	Sweden, Nordin 5889 (UPS)	JF703124	
Megaspora cretacea	Armenia, Aptroot 73835 (B)	KX253974	
Megaspora cretacea	Armenia, Gasparyan 600199170 (B)	KX253975	
Megaspora rimisorediata	Iran, Valadbeigi 2250 (TARI)	JF825525	
Megaspora rimisorediata	China, Xinjiang, XJU 20116002	KT443790	
Megaspora rimisorediata	China, Xinjiang, XJU 20136001	KT443789	
Megaspora rimisorediata	China, Xinjiang, XJU 20111617	KT443788	
Megaspora rimisorediata	China, Xinjiang, XJU 91815043	KT443787	
Megaspora verrucosa	Austria, Trinkaus (GZU)	AF332121	
Megaspora verrucosa	Austria, <i>Hafellner 48544 & Ivanova</i> (GZU)	AF332122	
Megaspora verrucosa	China, Xinjiang, XJU 200753	KT443786	
Megaspora verrucosa	China, Xinjiang, XJU 20000724	KT443785	
Megaspora verrucosa	U.S.A., Colorado, St. Clair C54042 (BRY)	KC667053	
Sagedia mastrucata	Norway, Troms, Nordin 5708 (UPS)	EU057913	
Sagedia simoensis	Sweden, Own-Larsson 9000 (UPS)	EU057926	
Sagedia zonata	Sweden, Nordin 5932 (UPS)	EU057949	
Ochrolechia parella	France, Brittany, Feige (ESS-20864)	AF329174	

sion 7.0) was used to reconstruct the Maximum Likelihood phylogenetic tree based on the GTR + I + G model (Nei & Kumar 2000; Kumar & al. 2016). Initial

tree(s) for the heuristic search were obtained automatically by applying Neighbor-Join and BioNJ algorithms to a matrix of pairwise distances estimated using the Maximum Composite Likelihood (MCL) approach, and then selecting the topology with superior log likelihood value. A GTR model of nucleotide substitution including a proportion of invariable sites and a discrete gamma distribution with five rate categories (GTR + I + G) were used in Maximum Likelihood approach.

Results

Phylogeny

The maximum parsimony analysis resulted 12 most parsimonious trees with 513 steps, consistency index (CI) = 0.591, retention index (RI) = 0.690, rescaled consistency index (RC) = 0.407 and homoplasy index (HI) = 0.409. The Maximum Likelihood analysis resulted a tree with the highest log likelihood (-2014.3274). Majority rule consensus tree for maximum parsimony analysis was congruent with the tree ob-

tained by Bayesian and maximum likelihood phylogenetic inference. The majority rule consensus tree of Bayesian analysis is shown here (Fig. 1) with posterior probabilities

Table 2. The main distinguishing characteristics of Megaspora cretacea, M. rimisorediata and M. verrucosa.

	Megaspora cretacea	Megaspora rimisorediata	Megaspora verrucosa
Thallus	whitish grey, irregularly delimited to almost lobate	ochraceous to bluish grey, dense net of elongate cracks over thallus	white to grey-white, continuous to areolate to verrucose
Soredia	pale bluish grey, covering most of thallus, c. 100 µm in diam.	dark bluish green, produced along sides of elongate cracks, 50–70 µm in diam.	absent
Hymenium	not inspersed, c. 150 μm high	not inspersed, to 150 μm high	inspersed at times, 200–250 μm high
Paraphyses	unbranched	branched and anastomosing	branched but not anastomosing
Asci	$125-140 \times 25-31 \ \mu m$	c. 145 × 46 μm	$200-230 \times 45-50 \ \mu m$
Ascospores per ascus	s 4	4–8	8
Ascospores	27–31 × 18–21 μm	$35-42 \times 23-27 \ \mu m$	$30-60 \times 21-42 \ \mu m$
Substrate	bark of <i>Juniperus</i> sp.	bark of <i>Juniperus</i> sp., <i>Quercus</i> sp.	soil, mosses, plant remains on calcareous rocks, bark

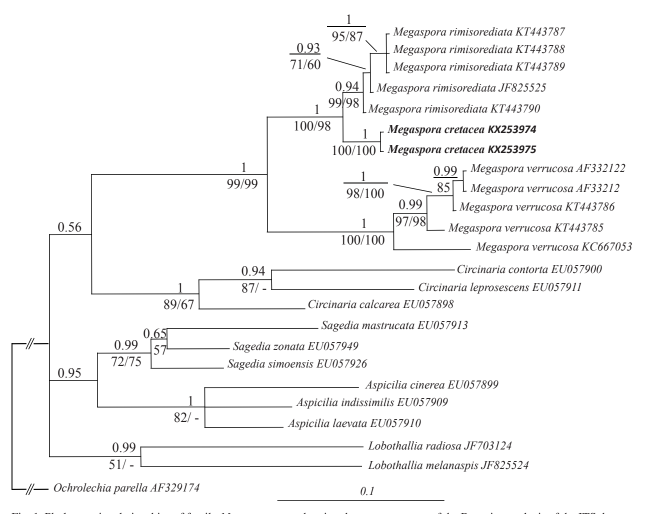


Fig. 1. Phylogenetic relationships of family *Megasporaceae* showing the consensus tree of the Bayesian analysis of the ITS dataset. Bayesian posterior probabilities are shown above the branches and MP/ML bootstrap values ≥ 50 are shown below the lines adjacent to the branches. Distance of outgroup and ingroup root is shortened three times.

of Bayesian analysis and bootstrap numbers of Maximum Parsimony and Maximum Likelihood analysis.

The molecular phylogenetic results confirmed affiliation of the new species to the genus *Megaspora*. It is clusters in a phylogenetic tree in *Megaspora*, as sister to *M. rimisorediata* (PP = 1; MP/ML BS= 100/100). The phylogenetic trees resulting from the three different analyses also confirmed *Megaspora* clade as a monophyletic group even after adding the new species samples (*M. cretacea*) with a high posterior probability and bootstrapping values (PP = 1; MP/ML BS = 99/99). Monophyly of species *M. verrucosa* and *M. rimisorediata* were confirmed with a high supporting values (PP = 1; MP/ML BS = 100/100 for *M. verrucosa* and PP = 0.94; MP/ML BS = 99/98 for *M. rimisorediata*).

Taxonomy

Megaspora cretacea Gasparayan, Zakeri & Aptroot, **sp. nov.** – MycoBank #817072 – Fig. 2A–C. Holotype: Armenia, Ararat, Vedi, Urtsadzor, Khosrov For-

est State Reserve, 40°00'42"N, 44°54'04"E, 1600 m, on

Juniperus polycarpos bark, 17 Jun 2015, *A. Aptroot 73835* (B 600200932; isotypes: ABL, GLM).

Diagnosis — Megaspora with thallus whitish grey, cretaceous, fully sorediate with soredia c. 0.1 mm in diam.; apothecia sparse, immersed; ascospores 4 per ascus, broadly ellipsoid, $27-31 \times 18-21 \mu m$, hyaline, thin-walled.

Description — Thallus whitish grey, crustose, ecorticate, to 0.2 mm thick, irregularly delimited to almost lobate, occupying areas up to 5 cm in diam. Medulla white, cretaceous. Soralia covering most of thallus surface, pale bluish grey; soredia c. 100 μm in diam. Photobiont chlorococcoid. Apothecia sparse, dispersed, immersed in thallus, round, 0.3–0.5 mm in diam.; disc black, concave; margin black, raised above disc, incurved, c. 0.1 mm wide, with some crenations. Hymenium IKI+ blue, c. 150 μm high, not inspersed with oil droplets. Subhymenium hyaline. Epihymenium greenish, colour unchanged in KOH. Hypothecium hyaline. Paraphyses 2–2.5 μm thick, not branched. Asci clavate, 125–140 × 25–31 μm. Ascospores 4 per ascus, broadly ellipsoid, 27–31 × 18–21 μm, hya-

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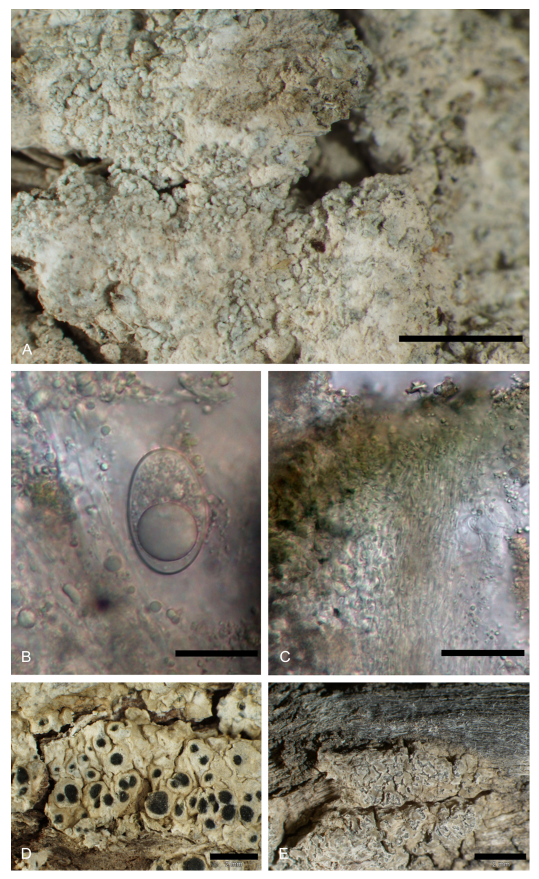


Fig. 2. A–C: *Megaspora cretacea*, holotype; A: thallus with soredia and apothecia; B: ascospore; C: hymenium (with excipulum at left). – D: *M. verrucosa* thallus with apothecia. – E: *M. rimisorediata* thallus with net of cracks and soredia. – Scale bars: A, D, E = 2 mm; B, C = $20 \mu m$.

line, thin-walled (less than 1 μm). *Pycnidia* not observed. *Conidia* not observed.

Chemistry — Thallus KOH-, C-, Pd-, UV-. TLC: No lichen substances detected.

Distribution and ecology — The species is known from two separate localities within the Khosrov Forest State Reserve, Armenia. It occurs on bases of trees of *Juniperus polycarpos* K. Koch in arid, open, montane forests. The forest ecosystems in the Khosrov Forest State Reserve, at 1400–2300 m, are generally dominated by oak trees (*Quercus macranthera* Fisch. & C. A. Mey. ex Hohen.) and sparse juniper (*J. polycarpos*) formations, accompanied by *Fraxinus excelsior* L., *Sorbus aucuparia* L., and species of *Acer* L. and *Pyrus* L. (Khanjyan 2004).

Etymology — The epithet is derived from word *cretaceus* (resembling chalk) in reference to the colour and texture of the thallus.

Additional specimen examined — Armenia: Ararat, Vedi, Urtsadzor, Khosrov Forest State Reserve, 39°59'07"N, 44°53'51"E, 1390 m, on *Juniperus polycarpos* bark, 17 Jun 2015, *A. Gasparyan* (B 600199170).

Discussion

Megaspora cretacea is a morphologically distinctive species, from which the two other species of the genus, M. verrucosa (Fig. 2D) and M. rimisorediata (Fig. 2E), can be separated as follows (Table 2): M. verrucosa has no soredia, whereas M. cretacea and M. rimisorediata are both sorediate; the closely related M. rimisorediata differs from M. cretacea by the presence of a dense net of elongate cracks over the thallus, dark bluish green soredia, branched paraphyses and larger ascospores.

Key to the species of *Megaspora*

1.	Soredia absent
_	Soredia present
2.	Thallus ochraceous to bluish grey with a dense net
	of elongate cracks; soredia produced along sides of
	elongate cracks, dark bluish green
	M. rimisorediata
_	Thallus whitish grey, irregularly delimited to almost
	lobate; soredia covering most of thallus, pale bluish
	grey

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