Taxonomic study of three new Antarctic *Asterochloris* (Trebouxiophyceae) based on morphological and molecular data

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INTRODUCTION

Lichens are among the best-known examples of symbiotic organisms and consist of fungal partners (mycobionts) and photosynthetic eukaryotic algal (phycobionts) or cyanobacterial (cyanobionts) partners. Lichens are biologically and ecologically important in terrestrial environments due to their influence on global carbon and nitrogen circulation, playing an essential role during early ecological succession (Fernández-Mendoza et al. 2011, Elbert et al. 2012, Pérez-Ortega et al. 2012, Borchhardt et al. 2017). Within lichen associations, the lichen-forming phycobionts transfer carbohydrates to the heterotrophic host in the form of sugar alcohol or glucose (Richardson et al. 1967, Richardson and Smith 1968), while mycobionts protect their partners from solar radiation and harsh environmental conditions such as extreme temperature fluctuations and desiccation (Honegger 2012). These partners have species selectivity, meaning that they correspond to each other, but not always in a one-to-one manner (Chodat 1913, Jaag 1929, Raths 1938, Kroken and Taylor 2000, Piercey-Normore and DePriest 2001, Muggia et al. 2018). To date, approximately 19,400 species of lichens have been described,
distributed in diverse climates from tropical to polar regions (Kirk et al. 2008, Garrido-Benavent and Pérez-Ortega 2017). Many studies of lichen taxonomy have focused on the mycobiont, but relatively less attention has been paid proportionally to the lichen-forming phycobiont. Although approximately 40 genera and 100 species of photobionts are known globally (Tschermak-Woess 1988, Friedl and Büdel 2008, Muggia et al. 2018), taxonomic studies of eukaryotic algae acting as phycobionts are not sufficient to understand their species diversity, especially in the Antarctic region (Engelen et al. 2016, Garrido-Benavent et al. 2017, Beck et al. 2019).

*Asterochloris* is closely related to the genus *Treboxxia* in having similar morphological characters of vegetative cells. In 1867, Schwendener recognized that a lichen is an association of green bodies and fungal hyphae. Subsequently, Treboux (1912) included the green bodies as members of the formerly described genus *Cystococcus*, following the classification system of Nägeli (1849). Warén (1920) made an attempt to divide *Cystococcus* into two subgenera (*Eucystococcus* and *Eleuterococcus*) based primarily on the number of autosporules and added eight new species to the two subgenera. Later, Puymaly (1924) recognized that *Cystococcus humicola*, the type species described by Nägeli (1849), was a different species from the lichen-forming green coccoid alga and suggested a new genus name, *Treboxxia*, based on observations of the gonidia in the thallus of the lichen species *Xanthoria parietina*. Thereafter, Ahmadjian (1960, 1967) classified *Treboxxia* into two groups using chloroplast features and vegetative cell shape. In addition, Archibald (1975) established a new genus, *Pseudotreboxxia*, based on differences in cell-wall characters during asexual reproduction. However, morphological and molecular data did not support Archibald’s classification system (Gärtner 1985, Friedl and Zeltner 1994, Friedl and Rokitta 1997). In 1980, Tschermak-Woess established a new genus, *Asterochloris*, for microalgae devoid of autosporules.

Recently, many molecular studies have supported *Treboxxia* and *Asterochloris* as distinct monophyletic clades. Early molecular studies performed based on nuclear small subunit and large subunit (LSU) rDNA sequence data suggested that *Treboxxia magna* and *T. erici* were more closely related to *Myrmecia biatoraeae* than other members of *Treboxxia* s. str. (Friedl and Zeltner 1994, Friedl 1995, Friedl and Rokitta 1997). Piercey-Normore and DePriest (2001) showed the monophyly of *Asterochloris* using nuclear internal transcribed spacer (ITS) 1, 5.8S and ITS2 sequence data. Later, an 18S rDNA-based phylogenetic tree of lichen photobionts revealed that *Asterochloris* was a monophyletic genus distinct from the closely related genera *Myrmecia* and *Treboxxia* (Friedl and Büdel 2008). Škaloud and Peksa (2010) emended the diagnosis of the genus *Asterochloris* and revealed 16 lineages in the genus, including nine new species-level lineages (Škaloud et al. 2015). More recently, eight new species of *Asterochloris* have been reported based on chloroplast morphology and molecular data (Moya et al. 2015, Škaloud et al. 2015, Kim et al. 2017).

To understand the phylogenetic relationships and taxonomy of *Asterochloris* isolated from lichen thalli in the vicinity of King Sejong Station, King George Island (Antarctica) and Morro Chico (Chile), the morphological characters of lichen photobionts were investigated by using light microscopy (LM), confocal laser microscopy (CM), and transmission electron microscopy (TEM). Molecular data analyses were also performed based on three genetic markers (nuclear-encoded internal transcribed spacer [ITS]), actin, and plastid-encoded ribulose bisphosphate carboxylase large chain (rbcL) to understand the phylogenetic relationships among the newly isolated *Asterochloris* and known members of this genus.

**MATERIALS AND METHODS**

**Taxon sampling**


**Phycobiont isolation and cultivation**

Initially, a small piece of rehydrated lichen thallus was fragmented by chopping with a sharp razor in a Petri dish or macerated by using a glass slide. Each free-laying
algal cell surrounded by broken pieces of the fungal hyphae was isolated by a micropipette. Then, each cell was washed five times with 3N Bold’s Basal medium (3NBBM) modified according to Thomas and Montes (1978) and transferred into the same medium in a well of a 96-well plate. The cells within the plate were incubated at 17°C under a 14 : 10 h light : dark cycle with 30 µmol photons m⁻² s⁻¹ using cool white fluorescent bulbs.

**Light microscopy**

Each culture strain was observed and identified under an Axio Imager.A2 (Carl Zeiss Inc., Hallbergmoos, Germany) equipped with differential interference contrast optics. Cell images were obtained with an AxioCam HRc (Carl Zeiss Inc.) photomicrographic system. Numerical values of morphological characters were determined by measuring 25-30 cells of each taxon from photographic images. We followed the terminology from Škaloud et al. (2015) and Gärtner (1985) for descriptions of detailed chloroplast morphology.

**Confocal microscopy**

CM was conducted with an LSM5 laser scanning confocal microscope equipped with an argon-krypton laser (Carl Zeiss Inc.). A 488 nm excitation line and an AOBS filter-free system were employed to collect emitted light between 498 and 700 nm. The autofluorescence of chlorophyll was used to visualize chloroplast structure. Collections of optical sections through chloroplasts were used to obtain a three-dimensional reconstruction of chloroplast morphology. The reconstructions of chloroplasts were made by LSM 5 software version 3.5 (Carl Zeiss Inc.).

**Transmission electron microscopy**

For TEM, aliquots of the culture were pelleted by centrifugation for 2 min at 2,300 xg (5,000 rpm) in an Eppendorf 5415D centrifuge (Eppendorf, Hamburg, Germany). After removing the supernatant, the pelleted cells were fixed in 2.5% glutaraldehyde mixed with 3NBBM culture medium for 1 h at 4°C. The glutaraldehyde-fixed cell pellets were washed 3 times in 3NBBM culture medium and postfixed in 1% OsO₄ for 1 h at 4°C. Dehydration, embedding, and polymerization were performed following the methods of Kim et al. (2017). The polymerized blocks were thinly sectioned to a thickness of 70 nm. Sections were collected on formvar-coated slot copper grids, stained with 3% uranyl acetate and Reynold’s lead citrate (Reynolds 1963) and examined and photographed with a JEM-1010 transmission electron microscope operating at 80 kV (JEOL, Tokyo, Japan). Images of the sections were recorded on Kodak EM Film 4489 (Eastman Kodak Co., Rochester, NY, USA) and scanned in tagged image file (TIF) format using Epson Perfection V700 Photo (Epson Korea Co., Ltd., Seoul, Korea).

**DNA isolation, amplification, and sequencing**

Cultured cells were harvested by centrifugation at the exponential growth phase (Eppendorf 5415D). Genomic DNA was extracted from the culture strains with a Dok-Do-Prep Genomic DNA Kit (Elpis-Biotech Inc., Daejeon, Korea), employing the animal tissue protocol described by the manufacturer. The nuclear ITS rDNA, actin and plastid rbcL coding gene were amplified by using a combination of forward and reverse primers in a T100 thermal cycler (Bio–Rad, Hercules, CA, USA). Amplification of the internal ITS rDNA, actin, and rbcL coding gene was performed with the following primers: ITS5F (5’-GGA AGT AAA AGT CGT AAC AAG G-3’) (White et al. 1990), ITS4R (5’-TCC TCC GCT TAT TGATAT GC-3’) (White et al. 1990), actinF2 (5’-AGC CGG GGT ACA GCT TCA C-3’) (Škaloud et al. 2017), actinR2 (5’-CAG CAC TTC AGG GCA GCG GAA-3’) (Škaloud and Peksa 2010), PRASF5 (5’-ATG GGT CCA CAA ACA GAA AC-3’) (Škaloud et al. 2015), PRASR5R (5’-TTG TCA ATA GTA TCA AAT TC-3’) (Sherwood et al. 2000, Škaloud et al. 2015), Astero_rbcL (5’-GGC T TTG AAA GCA GGT GTA AAA G-3’) (this study), and Astero_rbcLR (5’-CAC GAC CSG CGT TTC GKG-3’) (this study). The three genes were amplified by using a 25 µL reaction mixture containing 2 µL of DNA (20 ng), 1.0 µL of each primer (10 pmol), 1 µL of Accupower PCR PreMix (Bioneer Corporation, Daejeon, Korea) and enough distilled water to reach a total reaction volume of 25 µL. The DNA amplifications were performed using the following program: 5 min of denaturation at 94-95°C; 35-40 cycles of 94-95°C for 30 s, 45-55°C for 1 min, and 72°C for 1-2 min; a final extension at 72°C for 7-10 min; and an infinite hold at 4°C. The polymerase chain reaction (PCR) products were loaded onto an agarose gel and purified using a LaboPass PCR purification kit and LaboPass gel extraction kit (Cosmo Genetech Corporation, Daejeon, Korea) according to the manufacturer’s protocols. The purified PCR products were sequenced by using the PCR primers at Macrogen Corp. (Seoul, Korea). Forty-six new sequences were generated: 14 sequences of the nuclear ITS1, 5.8S, and ITS2 rDNA regions; 14 sequences of the actin gene and 18 se-
sequences of the rbcL gene. These sequences were aligned by eye using the Genetic Data Environment (GDE 2.6) program (Smith et al. 1994). The conserved regions of the three loci sequences were aligned across taxa and used for phylogenetic analyses. We excluded the parts of the sequences that could not be aligned.

Phylogenetic analyses

The combined dataset consisted of previously published and newly generated sequences. The dataset included 110 ITS rDNA, 102 actin gene and 24 rbcL gene sequences from Asterochloris (Supplementary Table S1). The dataset consisted of 2,252 characters (nuclear ITS = 506, actin = 426, rbcL = 1,320) for phylogenetic analysis. Maximum likelihood (ML) analyses were performed using RAxML version 8.2.12 (Stamatakis 2014) with a single general time-reversible plus gamma (GTR + GAMMA) model. We performed 1,000 tree inferences by using the -# option of the program to identify the best tree. Bootstrap values (ML bootstrap support) were computed based on 1,000 pseudoreplicates with the same substitution model. Bayesian analyses were implemented using MrBayes 3.2.6 (Ronquist et al. 2012), with the starting tree constructed randomly, two simultaneous runs (nruns = 2) and four Metropolis-coupled Markov chain Monte Carlo (MC³) algorithms for 10 × 10⁶ generations, with one tree maintained every 1,000 generations. Burn-in point was identified graphically by tracking the likelihoods (Tracer v.1.6; http://tree.bio.ed.ac.uk/software/tracer/). The first 2,500 trees were discarded, and the remaining 7,501 trees were used to calculate the posterior probabilities (PP) of each clade. Additionally, the sump command in MrBayes was used to confirm convergence. This analysis was repeated twice independently, with both analyses resulting in the same tree. The trees were visualized using the FigTree v.1.4.4 program, available at http://tree.bio.ed.ac.uk/software/figtree/.

Phycobiont ITS secondary structure

Secondary structures of the nuclear ITS sequences were built using the Mfold v.2.3 program (Walter et al. 1994, Zuker 2003), and the folding temperature was 25°C. The structures were compared with the published ITS structures of Asterochloris photobionts (Kim et al. 2017), and common secondary structures were drawn with PseudoViewer3 (http://pseudoviewer.inha.ac.kr/) and used to identify compensatory base changes (CBCs), hemi-CBCs, and single-base changes.

RESULTS

Morphology

Mycobiont specimens collected from King George Island (Antarctica) and Morro Chico (Chile), included Cladonia sp., C. gracilis, Sphaerophorus globosus, and Stereocaulon sp. (Fig. 1). Asterochloris cells among isolates of lichen-forming phycobionts were mostly spherical and occasionally oval in shape (Figs 2-4). The vegetative cells were 4-22 µm in diameter and had thick and smooth cell walls. Occasionally, partially thickened cell walls could

Fig. 1. Habitus of lichens from which the new species of Asterochloris were discovered. (A) Cladonia sp. showing primary squamules and cup-shaped podetia covered with powdery soredia. (B) Cladonia gracilis showing brown-colored, slender and unbranched podetia. (C) Sphaerophorus globosus showing yellowish-brown color and dichotomous branching. (D) Stereocaulon sp. showing a pale gray and granular thallus.
be distinguished in vegetative cells and sporangia. Most of the cell volume was occupied by the chloroplast, which contained one or more pyrenoids. The chloroplast in young cells was positioned parietally and had irregular margins. In mature cells, the chloroplast moved to the center and formed various shapes with different terminals depending on the species. These cells produced either aplanospores or zoospores for reproduction (Figs 2H, 3H & 4H).

Ultrastructural study was performed using culture strains of three new species to understand pyrenoid characteristic features (Fig. 5). In the case of *Asterochloris antarctica* (2015KGS-049A), mature cells had few thyla-koid tubules in the pyrenoid matrix, which were associated with a few pyrenoglobuli, as seen in the impressa-type (Fig. 5A & B). In mature cells of *A. pseudoirregularis*...
A dataset of nuclear ITS rDNA and actin and plastid \( \text{rbcL} \) genes revealed 18 well-resolved lineages with high support values (Fig. 6). The 15 formerly described species (\textit{Asterochloris erici}, \textit{A. magna}, \textit{A. glomerata}, \textit{A. irregularis}, \textit{A. excentrica}, \textit{A. leprarii}, \textit{A. gaertneri}, \textit{A. friedlii}, \textit{A. mediterranea}, \textit{A. echinata}, \textit{A. italiana}, \textit{A. phycobiontica}, \textit{A. lobophora}, \textit{A. sejongensis}, and \textit{A. woessiae}) were well recognized and formed their own monophyletic and well-supported lineages. The phylogenetic tree was di-

**Fig. 3.** Light micrographs and confocal reconstructions of the chloroplast structures in \textit{Asterochloris pseudoirregularis} (2015KGS-064C). (A & B) Mature aplanospores showing peripheral nucleus (N), shallowly lobed chloroplast (Cp) and pyrenoid (Py). (C-F) Vegetative cells showing pyrenoid positioned in the center of the shallowly lobed chloroplast. (G) Mature vegetative cell showing a shallowly lobed chloroplast with short, pointed terminal. (H) Aplanosporangium. (I-K) Confocal reconstructions of a crenulate chloroplast (I) or shallowly lobed chloroplast with flat ends (J) and aplanospores (K). Scale bars represent: A-K, 10 μm.

(2015KGS-064C) and \textit{A. stereocaulonica} (2015KGI-037A), several thylakoid membranes penetrated into the pyrenoid matrix and connected with many pyrenoglobuli (Fig. 5C-F), consistent with the irregularis-type according to Friedl (1989).

**Phylogenetic analysis**

The Bayesian and ML analyses based on the combined
consisted of 14 lineages. These lineages formed a monophyletic clade with strong support values (PP = 1.00, ML = 99), but the relationships among them were not fully resolved. The clade included five undesignated lineages and nine species, including a new species: *A. echinata*, *A. italiana*, *A. phycobiontica*, *A. lobophora*, *A. antarctica* sp. nov., *A. sejongensis*, *A. woessiae*, *A. friedlii*, and *A. mediterranea*. The new species *A. antarctica* also exhibited a strong sister relationship with *A. sejongensis* and *A. woessiae* (PP = 1.00, ML = 100).

The first clade (A) included nine species that were further divided into two subclades (PP = 1.00, ML = 99). The first subclade consisted of five species: *A. erici*, *A. glomerata*, *A. irregularis*, *A. magna*, and *A. pseudoirregularis* sp. nov. The new species *A. pseudoirregularis* was sister to *A. irregularis* (PP = 0.99, ML = 77). The second subclade included four species: *A. excentrica*, *A. stereocaulonicola* sp. nov., *A. leprarii*, and *A. gaertneri* (PP = 0.98, ML = 68). The second new species, *A. stereocaulonicola*, was sister to *A. leprarii* (PP = 1.00, ML = 97). The second major clade (B) consisted of 14 lineages. These lineages formed a monophyletic clade with strong support values (PP = 1.00, ML = 99), but the relationships among them were not fully resolved. The clade included five undesignated lineages and nine species, including a new species: *A. echinata*, *A. italiana*, *A. phycobiontica*, *A. lobophora*, *A. antarctica* sp. nov., *A. sejongensis*, *A. woessiae*, *A. friedlii*, and *A. mediterranea*. The new species *A. antarctica* also exhibited a strong sister relationship with *A. sejongensis* and *A. woessiae* (PP = 1.00, ML = 100).
Fig. 5. Transmission electron micrographs of Antarctic photobionts. (A) Chloroplast (Cp) morphology of *Asterochloris antarctica* (2015KGS-049A). (B) Enlarged image of the pyrenoid (Py) region showing irregularly scattered pyrenoglobuli (Pg) and thylakoid invagination (arrow). (C) Chloroplast of *Asterochloris pseudointegrilis* (2015KGS-64C). (D) Enlarged image of pyrenoid (Py) in (C) showing pyrenoglobuli (Pg) on the thylakoid membrane. (E) Vegetative cell of *Asterochloris stereocaulonicola* (2015KGIC-037A). (F) Enlarged image of the pyrenoid region in (E) showing pyrenoglobuli connected with thylakoid membranes. N, nucleus. Scale bars represent: A, C & E, 1 μm; B, D & F, 0.5 μm.
Fig. 6. Bayesian tree based on concatenated nuclear-encoded ITS rDNA, actin and plastid-encoded rbcL sequences. The numbers on each node represent posterior probabilities (PP; left) and maximum likelihood bootstrap values (ML; right) The bold branches have strong support values (PP = 1.00 and ML = 100%).
ITS secondary structures

The nuclear ITS rRNA secondary structures of each new species were compared with those of closely related species to determine the occurrence of CBCs (nucleotide changes on both sides of a paired base) and hemi-CBCs (change on only one side of the nucleotide pair, with the pairing preserved) according to Coleman (2000, 2003) and Ma et al. (2015). The ITS1 secondary structure was composed of four paired regions (helices I-IV, see Fig. 7A-F). In A. antarctica, the base changes were compared with those in the ITS1 and ITS2 secondary structures of A. sejongensis (Fig. 7A & B). In ITS1 (Fig. 7A), the bases were different in helix I (UCC – UUCUCC), in helix II (C:G – U:G), and in helix IV (U:U – U:C). In ITS2 (Fig. 7B), the bases were different in helix I (UACCA – CAUG), between helix II and helix III (AU – AGU), in helix III (C:G – U:G), and in helix IV (A:U – A:C). The ITS1 and ITS2 secondary structures of A. pseudoirregularis were compared with those of A. irregularis (Fig. 7C & D). The base differences of A. pseudoirregularis in helix II (GCUUUC – UCUUGC) and in helix IV (AAUU – AUUU) were detected only in ITS1 (Fig. 7C). The ITS1 and ITS2 secondary structures of A. stereocaulonicola were compared with those of A. leprarii (Fig. 7E & F). In ITS1 (Fig. 7E), the bases were different in helix I (UUCAC – UUUAC), in helix II (G:U – A:U and CUUG – CCUG), and in helix IV (UUCAAACCAUACU). In ITS2 (Fig. 7F), the bases were different in helix I (CCC – CCA), in helix III (UUGAUAUAAU and CCUGUCUUGU) and between helix IV and the LSU rDNA (UUCAAAUJC – UCIJAAUJC). The above three species had conserved motifs (Mai and Coleman 1997) that included a UU mismatch in helix II and a UGGU sequence in helix III of ITS2 (Fig. 7B, D & F).

Taxonomy

Taxon descriptions.

*Asterochloris antarctica* Y. J. Kim, J. I. Kim et W. Shin sp. nov.

Cells are mostly spherical in shape. Vegetative cells are 6-19 µm in diameter. Cells have thick and flat cell walls (Fig. 2A-G). Occasionally, a partially thickened cell wall can be distinguished. Most of the cell volume is taken up by the chloroplast (Fig. 2A-F). Chloroplast in young cells is positioned parietally with irregular margins and has one or two pyrenoids (Fig. 2A, C, D & G). In mature cells, the chloroplast is in the center and is shallowly lobe-shaped with elongated terminals (Fig. 2D & E) or crenulate (Fig. 2I & J). This species produces 64-128 aplanospores for asexual reproduction (Fig. 2H & K). Rarely, zoospores burst from oval or spherical sporangia.

Molecular signatures. Base changes (UCC – UUCUCC) in helix I of ITS1 (Fig. 7A) and base changes (UACCA – CAUG) in helix I and (A:U – A:C) in helix IV of ITS2 (Fig. 7B). Structures were compared with those of the ITS1 and ITS2 secondary transcripts of A. sejongensis, a closely related sister taxon in the tree.

Holotype. Deposited in the herbarium of Chungnam National University, Daejeon, Korea (CNUK).

Type strain. Deposited in the culture collection of Chungnam National University (strain number: 2015KGS-049A).


Etymology. The specific epithet “antarctica” is derived from the name Antarctica, corresponding to the geographic origin of this photobiont.

Distribution. Currently, the only known distribution area is in the King George Island, Antarctica.


*Asterochloris pseudoirregularis* Y. J. Kim, J. I. Kim et W. Shin sp. nov.

Cells are mostly spherical in shape. Vegetative cells are 4-18 µm in diameter. Cells have thick and flat cell walls (Fig. 3A-G). Cells at the sporangium stage occasionally have partially thickened cell walls. Most of the cell volume is occupied by the chloroplast-bearing one central pyrenoid (Fig. 3B-E). Chloroplast in young cells is positioned parietally and deeply lobed with irregular margins (Fig. 3E-G) while in mature cells, the chloroplast is located in the center of the cell (Fig. 3D & E). Chloroplast is shallowly or deeply lobed or crenulate (Fig. 3A-G) with flat or elongated terminals (Fig. 3E, F, I & J). This species produces 64-128 aplanospores for asexual reproduction (Fig. 3H & K). Rarely, zoospores are released from oval or spherical sporangia.

Molecular signatures. Base changes (GCUUUC – UCUUGC) in helix II and base changes (AAUU – AUUU) in helix IV of ITS1 (Fig. 7C). Structures were compared with those of the ITS1 secondary transcripts of A. irregularis.

Holotype. Deposited in the herbarium of Chungnam National University, Daejeon, Korea (CNUK).

Type strain. Deposited in the culture collection of Chungnam National University (strain number: 2015KGS-049A).


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Molecular signatures. Base changes (GCUUUC – UCUUGC) in helix II and base changes (AAUU – AUUU) in helix IV of ITS1 (Fig. 7C). Structures were compared with those of the ITS1 secondary transcripts of A. irregularis.

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Etymology. The specific epithet “antarctica” is derived from the name Antarctica, corresponding to the geographic origin of this photobiont.

Distribution. Currently, the only known distribution area is in the King George Island, Antarctica.

Fig. 7. Predicted secondary structures of the internal transcribed spacer (ITS) 1 and ITS2 transcripts of Antarctic Asterochloris species. (A & B) Predicted secondary structures of the ITS1 (A) and ITS2 (B) transcripts of A. antarctica (2015KGS-049A) drawn by comparison with those of the closely related species A. sejongensis. Base changes of Asterochloris species are indicated in red. (C & D) Predicted secondary structures of the ITS1 (C) and ITS2 (D) transcripts of A. pseudoirregularis (2015KGS-064C) drawn by comparison with those of the closely related species A. irregularis. (E & F) Predicted secondary structures of the ITS1 (A) and ITS2 (B) transcripts of A. stereocaulonicola (2015KGIC-037A) drawn by comparison with those of the closely related species A. leprarii. Base changes between two Asterochloris genotypes are indicated: the black boxes indicate base changes, and the red boxes indicate base changes in each new species that can serve as diagnostic molecular signatures. ITS2 transcripts, highly conserved U-U mismatches and UGGU motifs are marked with [ ].
National University, Daejeon, Korea (CNUK).

**Type strain.** Deposited in the culture collection of Chungnam National University (strain number: 2015KGS-064C).

**Type locality.** Samples of Cladonia sp. and Cladonia gracilis were collected from rocks and soil crusts in the vicinity of King Sejong Station, King George Island, Antarctica (-62.131490 S, -58.458890 W), in February 2015 and 2016. The lichen specimens have been deposited in the herbarium of the Korea Polar Research Institute (2015KGIC-037A, 2015KGIC-037B, and 2015KGIC-037D).

**Etymology.** The specific epithet “pseudoirregularis” is derived from that of the species A. irregularis and Greek "pseudo-" (= false) indicating the high similarity with this species.

**Distribution.** Currently, the only known distribution area is in the King George Island, Antarctica.

**Ecology.** In Cladonia gracilis commonly growing on gravelly soil and mosses.

*Asterochloris stereocalonicolica* Y. J. Kim, J. I. Kim et W. Shin sp. nov.

Cells are mostly spherical and occasionally oval in shape (Fig. 4). Vegetative cells are approximately 9-22 µm in diameter. Cells have thick and flat cell walls (Fig. 4A-F). Occasionally, the cell wall is partially thickened. Most of the cell volume is taken up by the chloroplast (Fig. 4). Chloroplast has one or two pyrenoids. Chloroplast in young cells is positioned parietally, with irregular margins. In mature cells, the chloroplast is moved to the center of the cell and is echinate with pointed terminals (Fig. 4H) or shallowly or deeply lobed with elongated terminals (Fig. 4B-E). This species produces 64-128 aplanospores for asexual reproduction. Rarely, zoospores are released from oval or spherical sporangia.

**Molecular signatures.** Hemi-CBCs (G:U – A:U) and base changes (CUUG – CCUG) in helix II and base changes (UUCAACCAAUACU – UUCAACCAAUACU) in helix IV of ITS1 (Fig. 7E) and base changes (CCC – CCA) in helix 1 and base changes (UUCAAUUU – UCUAAUUU) between helix IV and the LSU rDNA region in ITS2 (Fig. 7F). Structures were compared with those of the ITS1 and ITS2 secondary transcripts of A. leprarii.

**Holotype.** Deposited in the herbarium of Chungnam National University, Daejeon, Korea (CNUK).

**Type strain.** Deposited in the culture collection of Chungnam National University (strain number: 2015KGIC-037A).

**Type locality.** Phycobiont of Stereocalon sp. collected from rocks, mosses and soil crusts in the vicinity of King Sejong Station, King George Island, Antarctica (-62.131230 S, -58.458890 W), in February 2015. The lichen specimens have been deposited in the herbarium of the Korea Polar Research Institute (2015KGIC-037A, 2015KGIC-037B, and 2015KGIC-037D).

**Etymology.** The name “stereocalonicolica” refers to the genus name of the fungal partner Stereocalon and Greek “-cola” (= dweller).

**Distribution.** Currently, the known distribution area is in the King George Island, Antarctica.

**Ecology.** In Stereocalon species growing on rocks, scrubland, and soil on Barton Peninsula, King George Island.

**DISCUSSION**

Although more than 19,400 species (ca. 27% of all known fungi) of lichen-forming fungi are known (Jaktisch et al. 2016, Lücking et al. 2017), lichen-associated photobiont diversity is still underestimated (Muggia et al. 2018). In particular, being an extremely harsh environment, Antarctica is covered mostly by mosses, lichens, green algae and only two flowering plants, which thrive in the coldest, driest and windiest climates (Laybourn-Parry and Pearce 2007, Convey et al. 2009, Chae et al. 2019). On this inhospitable continent, the diversity of phycobiont species in the genus Asterochloris has been studied only once so far (Kim et al. 2017), but few studies on Trebouxia have been already performed (Romeike et al. 2002, Fernández-Mendoza et al. 2011, Pérez-Ortega et al. 2012, Ruprecht et al. 2012). In this study, we discovered three new Asterochloris species associated with the lichenized fungal genera Cladonia, Sphaerophorus, and Stereocalon in the vicinity of King Sejong Station (King George Island, Antarctica). These new species were differentiated from other species by morphological and molecular data.

Among the common green algal genera of lichen photobionts, Asterochloris has consisted of 15 species and is more closely related to the recently established genus Vulcanochloris than to Trebouxia and Myrmecia (Vančurová et al. 2015). However, both Asterochloris and Vulcanochloris can be differentiated by morphological and molecular characteristics. The genus Vulcanochloris is characterized by the unique formation of spherical incisions within the pyrenoid and by forming a strongly supported monophyletic lineage in Bayesian trees based on rbcL and ITS data, respectively (Vančurová et al. 2015),
but the pyrenoid type may be similar to the corticolar-type according to Friedl (1989). Chloroplast and pyrenoid types are important diagnostic features for species delimitation among the species of the genus *Asterochloris* (Gärtner 1985, Friedl 1989, Škaloud et al. 2015). *Asterochloris* has multiple chloroplast types depending on developmental stage and species. Our new species also have various chloroplast morphologies and share chloroplast features with previously described *Asterochloris* species (Škaloud et al. 2015).

The new species *Asterochloris antarctica* grouped with *A. sejongensis* and *A. woessiae* and was characterized by spherical cells possessing shallowly lobed and / or crenulate chloroplasts with flat terminals. Based on comparisons with closely related species, *A. antarctica* shared the crenulate chloroplast and spherical cell shape characters with *A. sejongensis*. However, *A. antarctica* did not have deeply lobed and / or echinate chloroplasts or oval or pyriform cells. The new species was differentiated from *A. woessiae* based on its deeply lobed chloroplasts with flat terminals. Compared with the ITS secondary structures of *A. sejongensis*, those of *A. antarctica* had a base change in helix I of ITS1 and base changes in helix I and helix IV of ITS2 (see Fig. 7). Although these species were closely related, they had morphological differences and unique molecular signatures in terms of ITS rRNA secondary structures, suggesting that *A. antarctica* is differentiated from *A. sejongensis*.

*A. pseudoirregularis* had shallowly and / or deeply lobed or crenulate chloroplasts and spherical cells. Based on comparisons with closely related species, *A. pseudoirregularis* was similar to *A. irregularis* in having shallowly and / or deeply lobed chloroplasts and spherical cells, but *A. pseudoirregularis* did not have oval or pyriform cells. Compared with the ITS secondary structures of *A. irregularis*, those of *A. pseudoirregularis* had base changes in helix II, and one single-base change in helix IV of ITS1. Although these two species are closely related, they have morphological differences and unique molecular signatures, suggesting that *A. pseudoirregularis* is different from *A. irregularis*.

The new species *Asterochloris stereocaulonicola* was characterized by spherical or oval cells, with vegetative cells possessing echinate chloroplasts with pointed terminals or shallowly and / or deeply lobed chloroplasts with elongated terminals. Compared with a closely related species, *A. leprarii*, the new species has distinctive echinate chloroplasts. Compared with the ITS secondary structures of *A. leprarii*, those of *A. stereocaulonicola* had a hemi-CBC and base change in helix I of ITS1 and base changes in helix 1 and (UUCAAAUU – UCUAAUU) between helix IV and the LSU rRNA of ITS2. Although these two species are closely related, they have morphological differences and many unique molecular signatures, suggesting that *A. stereocaulonicola* is differentiated from *A. leprarii*.

In ultrastructural studies of chloroplasts, the thylakoid membranes in the pyrenoid matrix were associated with pyrenoglobuli (Fisher and Lang 1971), and the form and arrangement of chloroplast thylakoids was divided into six categories (Friedl 1989). However, these patterns were not reflected in the phylogenetic relationships among the species. According to the morphological type of pyrenoids described by Friedl (1989), two new species, namely, *A. pseudoirregularis* and *A. stereocaulonicola*, had irregularis-type pyrenoids, while *A. antarctica* had gigantea- or impressa-type pyrenoids. When the pyrenoid types were mapped onto our Bayesian tree, the ultrastructure data were not congruent with the phylogenetic tree, suggesting that diverse pyrenoid types may have arisen independently in the evolutionary history of the genus *Asterochloris*. For example, irregularis-type pyrenoids were distributed in diverse clades in our phylogenetic tree (Fig. 6): *A. excentrica*, *A. glomerata*, *A. irregularis*, *A. italiana*, *A. pseudoirregularis*, *A. sejongensis*, and *A. stereocaulonicola*. *A. antarctica* has gigantea- and impressa-type pyrenoids, but *A. sejongensis* has irregularis-type pyrenoids, even though the two species are included in the same lineage.

In summary, the phylogenetic tree based on the combined nuclear ITS, actin, and *rbcL* sequence data had a topology similar to that of recently published phylogenies (Moya et al. 2015, Škaloud et al. 2015, Kim et al. 2017), and the genus was largely divided into two clades with moderate to high support values. The phylogenetic relationships among the members of *Asterochloris* were not fully resolved in clade B of our tree, but each species formed a monophyletic clade despite the morphology being misleading for species identification. Compared with closely related species, the three new species had unique CBCs, hemi-CBCs and single-base changes in ITS secondary structures, supporting that the new species were differentiated from closely related species in the tree.

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SUPPLEMENTARY MATERIALS

Supplementary Table S1. Strains of the genus Asterochloris taxa used in this study and the GenBank accession numbers for their nuclear ITS rDNA, actin, and plastid rbcl gene sequences (https://e-algae.org).

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