**Chloromonas nivalis** subsp. *tatrae*, subsp. nov. (Chlamydomonadales, Chlorophyta): re–examination of a snow alga from the High Tatra Mountains (Slovakia)

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Abstract: Melting snow fields populated by aplanozygotes of the genus *Chloromonas* (Chlamydomonadales, Chlorophyta) are found in polar and alpine habitats. In the High Tatra Mountains (Slovakia), cells causing blooms of brownish–red snow designated as *Scotiella tatrae* Kol turned out to be genetically (18S, ITS1 and ITS2 rDNA, *rbc*L) very closely related to *Chloromonas nivalis* (Chodat) Hoham et Mullet from the Austrian Alps. Therefore, *Sc. tatrae* is transferred into the latter taxon and reduced to a subspecies as *Cr. nivalis* subsp. *tatrae*. Both exhibit a similar photosynthetic performance, thrive in similar habitats at open sites above timberline, but differ in astaxanthin accumulation and number of aplanozygote cell wall flanges. In a field sample of *Cr. nivalis* subsp. *tatrae*, polyunsaturated fatty acids formed nearly 50 % of total lipids, dominating in phospholipids and glycolipids. *Cr. nivalis* subsp. *tatrae* represents likely a variation of a common cryoflora species with distinct morphology.

Key words: aplanozygote, astaxanthin, *Chloromonas nivalis*, fatty acids, field sample, High Tatra Mountains, photosynthesis, snow algae, *Scotiella*, ultrastructure

**INTRODUCTION**

Snow inhabiting algae are polyextremophiles in terms like habitat temperature, incident irradiation or availability of liquid water and nutrients. They have been reported from many mountainous and polar regions, where melting snow packs persist until summer (Hoham & Duval 2001). Cryoflora draws attention not only because of greenish to reddish snow discolorations, but also due to striking morphologic and physiologic strategies in course of its adaptation (Komárek & Nědbalová 2007), especially in comparison with their ‘conventional’ mesophilic relatives. Cvetkovska et al. (2016) summarized the evolutionary origin and diversity of psychrophilic algae, showing that Chlorophyta harbour more than a third of all known photo–psychrophiles (plastid–bearing protists metabolically active and able to reproduce at temperatures permanently close to the freezing point of water, which do not tolerate temperatures > 20 °C). Generally, psychrophiles occur in a great variety of habitats and exhibit strategies like anti–freeze proteins or accumulation of compatible solutes like glycine (De Maayer et al. 2014).

The Tatra Mountains, which are located mainly in northern Slovakia and to a smaller part in southern Poland, are the topmost part of the Carpathian Arc. The relief of this mountain range is very complex due to glacial events (Zasadni & Kłapyta 2014). The presence of red snow in the Tatra Mountains was already reported several centuries ago (Buchholz 1783). The high cryoflora biodiversity of this massif was summarized by Kol (1975a, b).

The genus *Scotiella* (*Sc*.) was established by Fritsch (1912) with the snow alga *Sc. antarctica* Fritsch as the type. It is characterized by striking cell wall surface structures like flanges, and daughter cells were said to possess the same ribbed morphology already before being released from the mother cell. Many fusiform snow algae inhabiting polar areas and high alpine zones were initially believed to be a member of this genus (Kol 1968). However, the true nature of *Scotiella*–stages as aplanozygotes (‘aplano–’ = immotile according to Ettl & Gärtner 2014) in the life cycle of a *Chloromonas* (Cr.)–like flagellate was revealed decades after the initial description by Hoham & Mullet (1977), who showed that vegetative daughter
cells lack any cell wall ornamentation. Thus, several Scotiella like species inhabiting snow were transferred to the genus Chloromonas (Hoham & Mullet 1978), e.g. Sc. antarctica was considered to be a zygotic stage of Cr. nivalis (Chodat) Hoham & Mullet, however the synonymy has been later questioned due to different cell flange morphology between these two species (Novák 2002). The genus Scotiella is now regarded as an artificial group of species retained for those taxaon's which's reproduction is unknown (HANAGATA 1998).

In this work, we studied populations collected close to the type locality of Sc. tatrae in the High Tatras, more than 50 years after the initial description by Kol (1965). All investigations were undertaken with field material, since cultures are not available yet, most likely because continuous growth under laboratory conditions was, to our knowledge, so far unsuccessful. The overall aim of this study was to evaluate abiotic habitat parameters, to check the phylogenetic position of Sc. tatrae using molecular and morphological traits, to measure the photosynthetic activity under varying light conditions, to describe the pigments which cause the secondary reddish colouration and finally, taking a putative adaptation to low temperatures into account, analysing the repertoire of fatty acids (FAs) occurring in membranes and cellular lipid depots.

The results were compared with the closest relatives, which is especially Cr. nivalis, from snow fields in the Austrian Alps.

**MATERIAL AND METHODS**

**Sampling and snow characteristics.** The cryoflora of brownish–red snow fields in the High Tatras (Slovakia) was investigated in June 2016 (Fig. 1). Exact geographic position, sampling date, elevation and corresponding sample code are summarized in Table 1. Surface snow samples were harvested either with a sterile plastic shovel and a 10 l bucket (sample 'LP01') or directly transferred into 50 ml sterile plastic tubes (other two samples). Samples were transported the same day to a field laboratory in close proximity. Prior to physiological measurements, the snow was gently melted over night at 4 °C to 5 °C and kept in its own melt water in the dark.

Electrical conductivity (EC) and pH of melt water samples were obtained with HANNA (Combo EC, Romania) and WTW Instruments (Inolab pH, Germany), respectively. Snow density was measured by coring snow with a cylindrical polyvinylchloride corer (length 72 cm, diameter 3.7 cm, volume 0.774 l). All cores were taken in the layer visually containing a bloom of snow algae and were orientated parallel to the surface of the snowbank according to HOHAM (1975). Cores were then transferred into plastic bags and the weight of each core was immediately measured by a spring balance Medio Line (precision 2 g, Pesola, Switzerland; similarly to KRŠTEK et al. 2011). Three independent random cores were averaged. The snow water content (SWC) was expressed in percent and calculated as (weight of a snow core divided by the weight of water of the same volume as the core) × 100.

**Light and electron microscopy.** Light microscopy (LM) was conducted using an Olympus BX43 (Olympus corporation, Japan) equipped with a digital camera D-1200F (Nikon, USA). Microphotographs were processed using the QuickPHOTO Camera 3.0 software (Promicra, Czech Republic). The same software was used to measure cell sizes. Preparation of samples for scanning and transmission electron microscopy (SEM and TEM, respectively) included joint steps: samples were fixed in 1.25% glutaraldehyde in 50 mM cacodylate buffer (pH 7.0) for 2 hours at 4 °C, two times washed in cacodylate buffer at 4 °C for 15 minutes, postfixed in 1% OsO₄ overnight at 4 °C, three times washed in distilled water, gradually dehydrated through an ascending ethanol series (until 70% ethanol in 4 °C, then switched to room temperature). For SEM, fixed cells were transferred into a critical point drying chamber (Bal–Tec CPD 030). After gold–coating, specimens were examined with a JEOL 6380 LV SEM at 25 kV. For TEM, cells were embedded in Araldine and PolyBed® 812 mixture (SPI–CHEM, USA). Ultrathin sections were cut with a diamond knife, stained using uranyl acetate and lead citrate. TEM grids were examined with a JEOL 1011 TEM (JEOL Ltd., Japan) at 80 kV. Microphotographs were taken with a Veleta CCD camera equipped with Olympus image analysis software (Olympus Soft Imaging Solution, Germany) and later modified by iTEM 5.1 (Germany Soft Imaging Solution, Germany).

**Cell counting and cultivation.** A 10 ml subsample was taken at Capie Lake shore snow field (location LP01) in order to determine the cell concentration. After snow melt and homogenisation by shaking, a subsample of 0.01 ml was put into a Cyrus I chamber (Meopta, Czech Republic) for counting under a light microscope. Two replicates were analysed, each with a grid area of 1 cm². More than 200 cells were counted for one run. Attempts were made to culture the dominating snow algal species. For this, cells were placed a day after sampling on petri dishes containing BBM medium 1.5% agar (BISCHOFF & BOLD 1963) and kept at 3 °C under continuous illumination of 20 µmol photons.m⁻².s⁻¹. Light
microscopy of field samples kept in their original meltwater was performed ~100 days after sampling to verify the development of daughter cells.

Isolation of DNA, sequencing, ITS2 rRNA secondary structure prediction and phylogenetic analysis. DNA was isolated from the sample LP01 with a DNeasy Plant Mini Kit (Qiagen). Cells were mechanically disrupted by shaking for 15 min (30 Hz) in the presence of glass beads (3 mm diameter, Sigma–Aldrich) in a Mixer Mill MM 400 (Retsch, Germany). Then, samples were put in freezer at −20 °C for half an hour. Subsequently, DNA was isolated in accordance with the manufacturer’s procedure. Quality and concentration of DNA was measured on the NanoDrop® ND–1000 Spectrophotometer (NanoDrop Technologies, Inc.).

The 18S small subunit ribosomal RNA gene (18S rDNA), internal transcribed spacer regions 1 and 2 (ITS1, ITS2 rDNA) and ribulose–1,5-bisphosphate carboxylase/oxygenase large subunit (rbcL) gene regions were amplified from DNA isolates by PCR using existing primers (Table 2). Amplification reactions for 18S rDNA were performed using cycle parameters according to Katana et al. (2001) with minor modification that duration of initial denaturation was prolonged for 10 min and annealing temperatures were 50 °C, 58 °C and 61 °C. Amplification reactions for rbcL gene were performed using cycle parameters according to Hojam et al. (2002) with minor change that three different annealing temperatures were applied (53 °C, 55 °C, 59 °C). Each 20 µl of PCR reaction for 18S and rbcL amplification contained 5 µl of DNA isolates (diluted to concentration of 5 ng µl⁻¹), 0.8 µl of each 10 µM primer, 1.6 µl of 25 mM MgCl₂, 1.5 µl of 2 mM dNTPs, 2 µl of 10× Taq buffer + KCl–MgCl₂, 7.8 µl sterile Milli–Q water, and 0.5 µl of 1U.µl⁻¹ Taq DNA polymerase (Fermentas, USA). Amplification reactions for ITS2 rDNA region were performed using cycle parameters according to Goff & Moon (1993) with minor modification that the gradient of annealing temperature was included (56 °C, 58 °C, 61 °C). Each 35 µl PCR reaction contained 1 µl of DNA isolates (diluted to concentration of 5 ng µl⁻¹), 1.4 µl of each 10 µM primer, 2.8 µl of 25 mM MgCl₂, 2.6 µl of 2 mM dNTPs, 3.5 µl of 10× Taq buffer + KCl–MgCl₂, 21.8 µl sterile Milli–Q water, and 0.5 µl of 1U.µl⁻¹ Taq DNA polymerase (Fermentas). The PCR products were stained with bromophenol loading dye, quantified on 1.5% agarose gel, stained with GelRed. The amplification products were purified and sequenced using an Applied Biosystems automated sequencer (ABI 3730xl) at Macrogen (Korea). The newly obtained sequences were entered in GenBank (USA).

<table>
<thead>
<tr>
<th>Sample reference</th>
<th>Date</th>
<th>Location</th>
<th>Altitude</th>
<th>GPS</th>
</tr>
</thead>
<tbody>
<tr>
<td>LP01 (this study)</td>
<td>18 June 2016</td>
<td>shore of Capie Lake</td>
<td>2073</td>
<td>N49°10.081 E20°02.279</td>
</tr>
<tr>
<td>Kol (1965)</td>
<td>22 June 1932</td>
<td>shore and ice cover of Okrúhle Lake</td>
<td>2000</td>
<td>N49°10.242 E20°02.178</td>
</tr>
</tbody>
</table>

Table 1. Samples of Chloromonas nivalis subsp. tatrae from the High Tatra Mountains with sample codes, collection date, sampling site, altitude (m a.s.l.) and geographic position (GPS).

Fig. 2−4. Overview of sampling sites of Chloromonas nivalis subsp. tatrae: (2) sampling location of this study next to Capie Lake (code LP01, red arrowhead) and the first observation in 1932 in Okrúhle Lake (asterisk); (3) coloured snowfields at LP01 (4) detail view of brownish–red snow after harvest.
submitted to NCBI Nucleotide sequence database (accession numbers for Cr. nivalis subsp. tatrae LP01: 18S, ITS1, ITS2 – KY499614, rbcL – KY499615; Cr. nivalis P24/DR4: rbcL – KY499616).

The nuclear rDNA ITS2 region was identified using a web interface at the ITS2 database showing position of 5.8S and 26S motives (http://its2.bioapps.biozentrum.uni-wuerzburg.de/cgi-bin/index.pl?amnator; KOETSCHEM et al. 2010). The sequence was then folded with 5.8S–LSU stem regions using the Mfold server accessible at http://mfold.rna.albany.edu/?q5mfold (Zuker 2003). A model of the secondary structure consistent with the specific features of nuclear rDNA ITS2 was selected: four helixes, U–U mismatch in helix II, and the UGGU motif near the 5´–end site apex of helix III (COLEMAN 2007).

Photosynthesis. In vivo chlorophyll fluorescence parameters were obtained in a chamber for liquid samples at approximately 0 °C with a pulse–amplitude modulated fluorometer (PAM 2000, Heinz Walz GmbH, Germany). To gain sufficient biomass, cells were concentrated by passive sedimentation of melt water in a 1 l plastic cylinder overnight and the snow meltwater was then used for measurements. Prior measurement, approximately 0.5 g of cells was kept in the snow meltwater in the dark for 30 minutes. Then, cells were exposed to photon flux densities (PFDs) of 5, 34, 66, 104, 201, 366, 622, 984, 1389, 1666 and 2018 µmol photons m⁻² s⁻¹ for 30 seconds each. Four replicate measurements were carried out. After each light exposure, a saturating pulse was given to detect effective photosynthetic quantum yield of photosystem II (PS II):

\[ Y(II) = \frac{(F'_m - F)}{F'_m} \] (GENTY et al., 1989)

where \( F'_m \) = maximal fluorescence yield emitted by chlorophyll–a in PS II when PSII reaction centres are closed by a saturation pulse and \( F \) = minimum fluorescence yield at steady state.

Afterwards, the relative electron transport rate of PSII (rETR) was calculated as follows:

\[ rETR = Y(II) \times \frac{PFD}{PPS} \times \frac{P_{PSII}}{P_{PSII}} \times ETR–Factor \]

where \( Y(II) = \) the effective PSII quantum efficiency, \( PFD = \) photon flux density, \( P_{PSII} = \) photons absorbed by PSII relative to photons absorbed by all photosynthetic pigments (default value 0.5), \( ETR–Factor = \) an empirical corrective factor (default value 0.84), which considers fraction of incident photons absorbed by the chlorophyll a molecules in PSII.

A curve of rETR upon PFD was calculated and fitted by the mathematical model according to WALSBY (1997) where photoinhibition is assumed. Presence of photoinduction was indicated by β:\n
\[ rETR = ETR_{max} \times \left(1 - \exp(-\alpha \times \frac{PFD}{ETR_{max}})\right) + \beta \times PFD \]

where \( rETR = \) relative electron transport rate of PSII, \( ETR_{max} = \) maximum relative electron transport rate of PSII, \( \alpha = \) initial slope alpha and \( \beta = \) slope of the curve at high irradiances, \( PFD = \) photon flux density. The parameters \( ETR_{max} \), \( \alpha \) and \( \beta \) were found using the Solver software in Microsoft Excel.

If \( \beta > 0 \) (no photoinduction), curve of rETR upon PFD was calculated and fitted according to model WEBB et al. (1974) where photoinhibition is not assumed:

Table 2. List of primers used for amplification of 18S rDNA, ITS1 rDNA, ITS2 rDNA (ITS) and rbcL markers; (F) forward; (R) reverse.

<table>
<thead>
<tr>
<th>Primer</th>
<th>Marker</th>
<th>Direction</th>
<th>Sequence</th>
<th>Reference</th>
</tr>
</thead>
<tbody>
<tr>
<td>SSU</td>
<td>ITS</td>
<td>F</td>
<td>CTGCCGGAAGGATCATGATT</td>
<td>PIERCEY–NORMORE &amp; DEPRIEST (2001)</td>
</tr>
<tr>
<td>LSU</td>
<td>ITS</td>
<td>R</td>
<td>AGTTCACTGCGGTTTGGTCTG</td>
<td>PIERCEY–NORMORE &amp; DEPRIEST (2001)</td>
</tr>
<tr>
<td>Al1500af</td>
<td>ITS</td>
<td>F</td>
<td>GCCGGCTCAGACTGATGC</td>
<td>HELMS et al. (2001)</td>
</tr>
<tr>
<td>LR3</td>
<td>ITS</td>
<td>R</td>
<td>GGGCCGTGTGGTCCAGACGG</td>
<td>VILGALYS &amp; HESTER (1990)</td>
</tr>
<tr>
<td>rbcL1F</td>
<td>rbcL</td>
<td>F</td>
<td>CTGCTTTTATCTGCGAACTGC</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>rbcL7R</td>
<td>rbcL</td>
<td>R</td>
<td>AAATAATACCAAGCCTACG</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>rbcL10F</td>
<td>rbcL</td>
<td>F</td>
<td>GTTCCGCCCTCAGGTTTACC</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>rbcL14R</td>
<td>rbcL</td>
<td>R</td>
<td>CACGGCTGAGGGTCCGTTG</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>rbcL19F</td>
<td>rbcL</td>
<td>F</td>
<td>CAAATCAGTTCACTGCGTTG</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>rbcL4R</td>
<td>rbcL</td>
<td>R</td>
<td>GAAATGAAAGCCTGCCGCTCC</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>18S1F</td>
<td>18S</td>
<td>F</td>
<td>CTGCTTTTATCTGCGAACTGC</td>
<td>HOHAM et al. (2002)</td>
</tr>
<tr>
<td>34F</td>
<td>18S</td>
<td>F</td>
<td>GCTCTAAAGTATGAGCCATGC</td>
<td>MIKHAILYUK et al. (2008)</td>
</tr>
<tr>
<td>895R</td>
<td>18S</td>
<td>R</td>
<td>AAATCCAAAGAATTTACCTC</td>
<td>MIKHAILYUK et al. (2008)</td>
</tr>
<tr>
<td>18F</td>
<td>18S</td>
<td>F</td>
<td>AACCGGTGGATCCTGCGACT</td>
<td>KATANA et al. (2001)</td>
</tr>
<tr>
<td>18R</td>
<td>18S</td>
<td>R</td>
<td>TGAATCCTTGCAGGGTCCACTACG</td>
<td>KATANA et al. (2001)</td>
</tr>
<tr>
<td>1122R</td>
<td>18S</td>
<td>R</td>
<td>CAATTCTTTAAGGGTCCAG</td>
<td>MIKHAILYUK et al. (2008)</td>
</tr>
</tbody>
</table>
rETR = ETR_max(1−exp(−α × PFD/ETR_max))

The fitting model with a lower sum of squared deviations was chosen.

**Pigment analysis.** Chlorophylls and carotenoids were isolated and determined by a protocol optimized for snow algae described by Remias & Lütz (2007) and Remias et al. (2013). Briefly, cells were lyophilized on glass fibre filters and broken in a shaking mill and using quartz balls and mill jars pre-cooled with liquid nitrogen. Extraction was performed with dimethylformamide (Sigma–Aldrich) and compounds were identified and quantified by high performance liquid chromatography with diode array detector (Agilent ChemStation 1100).

**Lipid extraction.** The extraction procedure was based on the method of Bligh & Dyer (1959), except that 2–propanol was substituted for methanol, since 2–propanol does not serve as a substrate for phospholipases (Kates & Volcani 1996). The alcohol–water mixture was cooled, one part chloroform was added and the lipids from the lyophilized cells were extracted for 30 min. Insoluble material was sedimented by centrifugation and the supernatant was separated into two phases. The aqueous phase was aspirated off and the chloroform phase was washed three times with two parts 1 M KCl each. The resulting chloroform phase was evaporated to dryness under reduced pressure. 10 mg of total lipid extract in 1 ml of chloroform was applied to Sep–Pak Vac Silica cartridge 35cc (Waters; 10 g normal–phase silica), and then eluted with 30 ml each of the following solvents: chloroform (neutral lipids), acetone (glycolipids), and methanol (phospholipids) at 4 ml min⁻¹ (Saunders & Horrocks 1984). The eluates of each lipid class were then evaporated prior to further analysis.

**Fatty acid methyl esters analysis (FAMEs).** The lipids (neutral lipids, glycolipids and phospholipids) (~5 mg of total dry weight) were saponified overnight in 10% KOH–MeOH at room temperature. A FA fraction obtained from saponification was partitioned between alkali solution (pH 9) and diethylether to remove basic and neutral components. The aqueous phase, containing FAs, was acidified to pH 2 and extracted with hexane. The FA fraction was methylated using CH₃N₁₃. Gas chromatography coupled with mass spectrometry (GC–MS) of FAMEs mixture was done on a Finnigan 1020 B with electron ionization (EI) mode. Splitless injection was at 100 °C for 1 min, subsequently increasing at 20 °C min⁻¹ to 180 °C and at 2 °C min⁻¹ to 250 °C, which was maintained for 1 min.

The carrier gas was helium at a linear velocity of 60 cm.s⁻¹. All spectra were scanned within the range of m/z 50–500. The structures of FAMEs were confirmed by comparison of retention times and fragmentation patterns with those of standard FAMEs (Supelco, Prague) (Rézanka 1990; Dembitsky et al. 1991).

**Statistical analysis.** Cell sizes (length, width, length to width ratio) of Sc. tatrae (LP01) were compared with those from Cr. nivalis at the Austrian Alps, (P12, P24/DR4, DR43, GK05, GK09; collection data of these field samples in Table S1 and in Remias et al. 2010). Kruskal–Wallis test was used for testing the hypothesis that cell size median of all Cr. nivalis samples from the Austrian Alps is not different. Moreover, Mann–Whitney test was used for testing the hypothesis that the median of two groups is identical (Sc. tatrae vs. Cr. nivalis).

**RESULTS**

**Collection sites and habitat conditions**

In the High Tatra Mountains (Slovakia), brownish–red snow fields were found during June 2016 at elevations of approximately 2000 m a. s. l. (Figs 2–4). A site in the vicinity to the locality of the first observation of the snow alga Sc. tatrae was sampled: the coloured snow was harvested at the shore of Capie Lake in Mlynická Valley (sample LP01) (Table 1). The population contained virtually only cells with a morphology according to Sc. tatrae. Scattered cells of the fungi Chionaster nivalis Bohlin (Wille) occurred. The habitat conditions of this locality are shown in Table 3. Moreover, snow–fields dominated mainly by morphotype Cr. nivalis and rarely of Sc. tatrae were found in other sampling spots slightly distant from the Okrúhle Lake (at the shore of Vysné Kozie Lake in Mlynická Valley – 0.5 km far away; at the shore of Upper Sesterske Lake in Velká Studená Valley – about 10 km far away).

**Morphology and ultrastructure**

Cell wall surface and intercellular organisation of field samples was described by light and electron microscopy (Figs 5–27). The fusiform cells of Sc. tatrae were taxonomically determined by their characteristic surface structures, and the harvested cell stage was tentatively designated as an aplanozygote on the basis of the life cycle proposed by Hoham & Mullet (1977). The

<table>
<thead>
<tr>
<th>Sample/reference</th>
<th>EC</th>
<th>pH</th>
<th>SWC</th>
<th>cells per ml meltwater</th>
<th>cell length</th>
<th>cell width</th>
<th>L:W ratio</th>
<th>flanges</th>
</tr>
</thead>
<tbody>
<tr>
<td>LP01 (this study)</td>
<td>15</td>
<td>5.7</td>
<td>52.6±3.8</td>
<td>19950±1995</td>
<td>19.8±1.3</td>
<td>12.8±0.9</td>
<td>1.55 ± 0.12</td>
<td>(9)10–12</td>
</tr>
<tr>
<td>KOL (1965)</td>
<td>—</td>
<td>5.2</td>
<td>—</td>
<td>—</td>
<td>18–24</td>
<td>12–15</td>
<td>—</td>
<td>10–12</td>
</tr>
</tbody>
</table>

Table 3. Abiotic habitat parameters and cell sizes of Chlamydomonas nivalis subsp. tatrae field samples from the High Tatra Mountains. Electrical conductivity (EC; µS cm⁻¹), pH of meltwater and snow water content (SWC; %), population density ± SD (standard deviation), average sizes of cells in µm ± SD, length to width ratio (L:W ratio) ± SD and number of cell wall flanges counted at equatorial region of cell in cross–section view are shown.
ornamented cell wall structures were made of straight to sometimes slightly undulated flanges (Figs 5−6, 13−14), and LM showed that (9)10−12(14) flanges per cell were present in equatorial region, which became evident at cross–section view (Fig. 7). The average cell sizes are summarised in Table 3. The cytoplasm was almost entirely occupied by lipid bodies containing a secondary pigmentation, thus causing an overall orange impression of the cells (Fig. 8). Rarely, young aplanozygotes surrounded by a smooth (primary) cell wall were observed (Fig. 9). SEM confirmed the presence of 10 to 14 flanges (Fig. 18), and further details of their arrangement could be observed: most common were flanges reaching from pole to pole (Fig. 15) and from one pole nearly to the antapex (Fig. 16). Rarely, flanges reached from one pole only to the equatorial position of the cell (Fig. 16), bifurcations (Fig. 17) or isolated shorter flanges (Fig. 15) also occurred. In a few cases all these secondary surface structures were found on a single aplanozygote. The ultrastructure observed by TEM is presented in Figs 19−27. Three different life cycle stages were recognized. Firstly, ovoid young aplanozygotes still without cell wall flanges, usually with one compact chloroplast containing starch grains (Fig. 19). In a few cases, an undulated cell membrane was found (Fig. 19). Moreover, mitochondria were observed (Fig. 20). Well–developed Golgi bodies and a cytoplasm rich in vesicles and ribosomes indicated an active metabolism (Fig. 21). Vacuoles full of electron dense, sometimes crystalline content were also present (Fig. 21). Secondly, intermediate aplanozygotes with flanges at the secondary cell wall surface, but still surrounded by a primary cell wall were observed (Figs 22−25). Thirdly, and prevailing in the sample, ‘mature’ aplanozygotes with fully developed flanges. The majority of the cytoplasmatic volume was occupied by large lipid bodies (containing secondary pigments) (Figs. 26−27). Instead of a compact chloroplast, several smaller spherical plastids located around the nucleus were present. Starch grains were very small or absent. Other life cycle stages like vegetative flagellates or any cleaving activities (formation of daughter cells) were not observed directly after sampling. 

In order to discriminate Sc. tatrae from its’ closest relative, Cr. nivalis from the Austrian Alps, cell sizes of aplanozygotes of both snow algae were compared. Sc. tatrae aplanozygotes were significantly smaller in term of cell length (17.6−23.1, median 19.5 vs. 18.35−27 µm, median 22.4), width (10.9−16.2, median 12.9 vs. 10.6–16 µm, median 13.2) and length to width ratio (1.33−1.8, median 1.55 vs. 1.38–2.04, median 1.7) (Figs S1−S4). However, significant differences in cell sizes were found also among samples of Cr. nivalis (Table. 4).

**Population density and cultivation assay**

The population density of Sc. tatrae (LP01) was 19 950±1995 cells.ml⁻¹ meltwater (Table 3). Cultivation assays were unsuccessful, and no cell cleavages have been observed up to three months after inoculation on
solid medium. However, in a field sample kept in its original meltwater for approximately 100 days, many cells started to contract the protoplast (Fig. 10) and to generate four elongate daughter cells (4.5−6.7 µm wide × 15.4−20.4 µm long), still surrounded by the mother cell wall (Figs 11−12). The constant number of four daughter cells points to foregoing meiotic divisions. The daughter cell pigmentation was similarly orange–greenish like that of the mother cell before cleavage.

**Phylogenetic analysis**

The phylogenetic affiliation of *Sc. tatrae* was elucidated based on molecular analysis of three nuclear (18S rDNA, ITS1 rDNA, ITS2 rDNA) and one plastid marker (*rbc*L). The 18S rDNA gene sequence of *Sc. tatrae* was identical with *Cr. nivalis* from the Austrian Alps (sample P24/DR4, accession number GU117576.1, Remias et al. 2010) except for two ambiguous bases at the end of the P24/DR4 sequence resulting in identity 99.9% (identical 1697 from 1699 bases). This affiliation was supported by comparing the secondary structures of the ITS2 region between the Slovak and the Austrian populations (letters outside the structure indicate differences; Fig. 28): Twenty nucleotide differences were found, one in the helices I and II, both of them were located in single stranded regions, ten differences in the most conserved helix III and eight changes in expansion segments between the III. helix and LSU stem. There was no compensatory base change (CBC; changes on both sides of structures which maintain the pairing between nucleotides). One hemi–CBC (hCBC; change only on one side maintaining the pairing) was found in helix III (A–U in *Cr. nivalis* vs. A–G in *Sc. tatrae*, marked by asterisk in Fig. 28). ITS2 rDNA identity was 94.5% (identical 342 out of 362 bp). A close affiliation between both taxa was deduced also from a pairwise comparison of the variable nuclear marker ITS1 rDNA, revealing 98.4% identity (identical 308 of 313 bp; three substitution detected and two ambiguous bases in the P24/DR4 sequence). There was no other more similar ITS2 sequence in public databases which can be used for secondary structure comparison with *Sc. tatrae* since other hits in BLAST search showed identity less than 76% and query cover less than 57%. The plastid marker (*rbc*L) of *Sc. tatrae* was 99.6% identical with the corresponding of *Cr. nivalis* P24/
DR4 (560 out of 562 bp, two ambiguous bases in P24/DR4 sequence were present in one codon, thus one amino acid in the product of translation was labelled as 'X', sequenced in this study) and showed a sister relationship to Cr. sp. ‘Gassan B’ from Japan (identity 96.5%, identical 1004 out of 1039 bp, sequence LC012743.1, see the phylogenetic tree – Fig. S14 in Matsuzaki et al. 2015).

Photosynthesis

In order to test the photosynthetic activity of this snow alga, the performance of field samples was measured under different light levels at the temperature of the melting snow habitat (photosynthesis/irradiance-curves). In general, aplanozygotes of Sc. tatrae were physiologically active and exhibited an α value of 0.19 ± 0.01, a relative ETR_{max} of 21.8 ± 1.0 and an
Fig. 28. Comparison of secondary structure of ITS2 rDNA transcripts between *Chloromonas nivalis* subsp. *tatrae* (accession number: KY499614, this study) and *Chloromonas nivalis* from the Austrian Alps (accession number GU117576.1, Remias et al. 2010). Note the U–U mismatch in the helix II (arrowheads). Differences characteristic for the latter are shown by nucleotides outside of the structure. The asterisk indicates hemi–compensatory base change.

$I_k$ value of $149 \pm 18$ µmol.photons.m$^{-2}$.s$^{-1}$ (Fig. 29). Photoinhibition was noticed from 600 µmol.m$^{-2}$.s$^{-1}$ on. For comparison, values of *Cr. nivalis* from the Austrian Alps (from Remias et al. 2010) were integrated in Fig. 29 and showed a likewise photosynthetic performance. The only difference was a lower parameter $I_k$ about one third for *Cr. nivalis* ($I_k = 106$ µmol.photons.m$^{-2}$.s$^{-1}$). The irradiance level from which on *Cr. nivalis* was subject to photoinhibition is not known, because application of higher irradiance levels in Remias et al. (2010) were not possible due to technical reasons.
Table 4. Statistical significance in differences of aplanozygotes length, width and length to width ratio (L:W) between Cr. nivalis subsp. tatrae from the High Tatra Mts. and Cr. nivalis from the Austrian Alps, and among latter group; n.s. not significant.

<table>
<thead>
<tr>
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<th>Cr. nivalis subsp. tatrae vs. Cr. nivalis</th>
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<tbody>
<tr>
<td>Length</td>
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<tr>
<td>Width</td>
<td>&lt;0.05</td>
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<tr>
<td>L:W ratio</td>
<td>&lt;0.0001</td>
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</table>

Pigments
In mature aplanozygotes, chlorophyll–a and –b accounted for 46.6%, primary (plastidal) carotenoids 12.0% and secondary (non-plastidal) carotenoids 41.4%. The latter were identified as derivatives of the keto-carotenoid astaxanthin by their spectral absorbance and occurred as 3 peaks. Their late retention times the non-polar region (compared to non-derivatives astaxanthin standard) of the chromatogram point to the presence of astaxanthin esterified with a variety of different FAs. The ratio astaxanthin to chlorophyll–a was 1.465. Other secondary carotenoids were not detected.

Fatty acid composition
The relative content of FAs (as % of total lipids and as % in the three major lipid groups) in Sc. tatrae is given in Table 5. FAs with 14 to 18 carbon atoms were detected. The cells had a high level of PUFAs (49 % in total lipids), whereas the content of saturated acids did not exceed 34 % (formed mainly by palmitic acid, 16:0). The main monounsaturated fatty acid (MUFA) was oleic acid (18:1n9, 14 %). The major PUFA were α-linolenic acid (18:3n3, 19%), followed by linoleic acid (18:2n6, 10 %), hexadecatetraenoic acid (16:4n3, 7 %) and stearidonic acid (18:4n3, 6 %). There were significant differences in FA composition among the three major lipid classes. The level of saturated lipids was the highest in neutral lipids (48 %), whereas PUFAs were dominant in phospholipids (67.1 %) and glycolipids (69 %) (Table 5). The total lipid content in the biomass was 17.6 % (of dry weight).

Taxonomic treatment
Chloromonas nivalis subsp. tatrae (Kol) Procházková, Remias, Řezanka et Nedbalová subsp. nov.
Synonym: Scotiella tatrae Kol 1965: 147, figs 1–5, 8–11.
Basionym: Pteromonas nivalis Chodat 1902: 145, fig. 70.

Description: mature zygote (causing the coloured snow phenomenon) elongate to somewhat fusiform, 16.4–24.0 µm long, 10.0–16.2 µm wide, length to width ratio 1.55–1.64, one nucleus, single axial chloroplast or several smaller spherical ones, abundant peripheral lipid globules containing orange carotenoids (astaxanthin to chlorophyll ratio ~ 1.5), two layered cell wall, most frequently 10–12 flanges, rarely less (9) or more (up to 14) in the equatorial region of the cell surface; aplanozygote division into four daughter cells, which are smooth–walled elongate 15.4–20.4 µm long, 4.5–6.7 µm wide; occurring in open sites above timberline; flagellates unknown; no pyrenoid or stigma visible; young zygote smooth–walled still without flanges, one compact chloroplast.

Holotype: specimen LP01 deposited at Herbarium of CAUP (Charles University in Prague), material consists of gold coated aplanozygotes.
Iconotype: Figs 5–27.

Etymology: The species epithet tatrae is based on the name of the geographic region, the Tatra Mountains, where it was found. It is genitive of the substantive ‘Tatra’ and this form corresponds with former description of ‘Sc. tatrae’ done by Kol.

Type locality: Snow fields close to the rocky shore of Capie Lake (Mlynická Valley), High Tatra Mountains, Poprad District, Slovakia.

Distribution: so far known from snowfields on shores of four alpine lakes: Okrúhle Lake, Capie Lake, Vyšné Kozie Lake (Mlynická Valley) and Upper Sesterske Lake (Velká Studená Valley) in the High Tatra Mountains, Slovakia.

Remarks: Any deposited material from the first observation of ‘Sc. tatrae’ is unknown. Kol (1965) reported 10–12 cell wall flanges on basis of L.M. Cr. nivalis subsp. tatrae differs from Cr. nivalis sensu Chodat (1902) in having mostly 10 or more flanges on cross–section instead of 8–9. Besides, Cr. nivalis in the Austrian Alps (Remias et al. 2010) differs from subsp. tatrae physiologically in the accumulation of...
secondary carotenoids (about three times lower astaxanthin to chl–a ratio in mature cells for the former) and slightly larger cell sizes of aplanozygotes: (18.4–27 × 10.6–16 µm). Aplanozygotes of *Cr. nivalis* from North America (Hoham et Mullet 1977) possess a lower number of flanges at cross–section (six to eight), are larger (16–37 × 10−27 µm) and lack fusing/diverting flanges apart from the apex (the latter are typical for *Cr. nivalis* from the Austrian Alps and for *Cr. nivalis* subsp. *tatrae*).

**DISCUSSION**

*Scotiella tatrae* was initially described from red and pink snow in the High Tatra Mountains at Okrúhle Lake, 2000 m a.s.l. (formerly Döller Lake) where it developed an almost monospecific bloom (Kol 1965). In a much lower extend, *Cr. nivalis* was also present, but no exact proportion to the former was given (‘in einer kleineren Menge’ according to Kol 1965). In that report, scattered cells of *Chlamydomonas (Cd.) nivalis* (Bauer) Wille, fungi *Chionaster nivalis* and *Chionaster bicornis* Kol were found as well. We collected our sample from a snow field close to the shore of Capie Lake, which is situated about 150 m south from the place of the initial observation (Fig. 2). Generally, the subspecies *tatrae* seems to be restricted to the High Tatra Mountains and can thus be likely regarded as endemic. A single report of scattered cells from elsewhere (Kühhtai region, 2500 m a. s. l., Tyrolean Alps, Austria) by Kol (1970) should be questioned, because such observation of *Sc. tatrae* in Austrian Alps has never been repeated in spite of intensive cryobiological research of this region (Remias et al. 2010).

For the population harvested in course of this study, the new name combination *Cr. nivalis* subsp. *tatrae* is proposed, since it shows a very close molecular affiliation to *Cr. nivalis* from the Austrian Alps based on the investigated molecular markers (18S rDNA, ITS1 rDNA, ITS2 rDNA, *rbcL*). CBCs in ITS2 secondary structures correlate with Ernst Mayr’s biological species concept (Coleman 2009). This hypothesis is also referred to as the CBC species concept. In our study, no CBCs in conserved parts (helices II and III) were detected, however, nucleotide difference in this

<table>
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<th>Fatty acid</th>
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variable marker between *Cr. nivalis* subsp. *tatrae* and *Cr. nivalis* of about 5.5 % suggests that a molecular divergence between them is ongoing. In comparison, it was nearly two fold higher than in two strains of *Cd. reinhardtii* P. A. DANGEARD, NIES–2463 and SAG 11–32a, which can cross and produce zygotes (3.3 %, NAKADA et al. 2010). On the other hand, it was slightly less than that between *Microcystis* *basinucleata* DEMCHENKO, MIKHAIYUK & PRÖSCHOLD and *M. monadina* (EIHENBERG) STEIN (5.9 %), recently separated taxa based on polyphasic approach including several CBCs in helix III (DEMCHENKO et al. 2012). On the other hand, even in absence of any CBC, algae may belong to the different species with a probability of 0.24 (WOLF et al. 2013) because total loss of gamete compatibility may be observed sooner than any CBC appear in the most conserved region of helix III (COLEMAN 2009). Between *Cr. nivalis* and subsp. *tatrae*, there were found similarities in the intracellular organization of aplanozygotes, cell sizes, photosynthetic performance and type of preferred habitat (see REMIAS et al. 2010). However, the significant differences in cell wall surface organization, a feature of taxonomic relevance, that seems to be stable considering no detectable changes since first descriptions of this snow alga several decades ago, justified the setup of a new subspecies. It remains open, if hybrids between subsp. *tatrae* and the main form of *Cr. nivalis* occur.

**Ecology**

Low electrical conductivities and slightly acidic pH were characteristic for the habitat of *Cr. nivalis* subsp. *tatrae* (Table 3). These findings are in good agreement with environmental conditions recorded for *Cr. nivalis* in the Austrian Alps (pH = 4–6.2, EC = 2.8–7.2 µS.m⁻².s⁻¹; REMIAS et al. 2010). pH values of snow meltwater with algae present were summarised in HOHAM et al. (2007). They ranged from 3.8 to 8.1, with *Cr. pichinchae* WILLE in the most acidic environment close to coniferous trees (HOHAM 1975) and *Cr. polyptera* (FRITSCH) HOHAM, MULLET & ROEMER in the most alkaline meltwater influenced by a guano input from close penguin colonies (LING & SEPPLET 1998). Similarly high pH of meltwater (7.7–7.9) was caused by input of volcanic ash (LUTZ et al. 2015). Meltwater conductivities with snow algae present varied from <1 to 950 µS.m⁻².s⁻¹. For instance, the very low values were found in arctic habitats of red orange cells affiliated to clade of *Cd. nivalis* (EC = 0.83) and *Cr. nivalis* (EC = 7.4) in Svalbard (MÜLLER et al. 1998). High EC was reported from the environment of *Cr. polyptera* in Antarctica in close proximity to bird colonies (LING & SEPPLET 1998). The snow water contents in snow surface samples containing *Cr. nivalis* subsp. *tatrae* are in good agreement with *Cr. nivalis* (45–52 %; HOHAM & MULLET 1977). This parameter is influenced by air temperatures during day and depends also to the depth of sampling (HOHAM & MULLET 1977). Generally, the availability of liquid water in the snow influences the life cycle development of *Chloromonas* snow algae: sexual stages occur if SWC is lower (47–54 %), whereas the asexual phase dominate during a higher SWC values (57–63 %, HOHAM & DUVAL 2001). The population density of *Cr. nivalis* subsp. *tatrae* was twice times higher than the detection limit from which on snow discoulouration can be recognized by bare eyes (>10⁵ cells per ml⁻¹ of meltwater; HOHAM & DUVAL 2001). Generally, the intensity of colouration may change in dependence on the cellular pigment composition, prevailing life cycle stage and the seasonality. Due to melt water flush out processes, cryoflora can have an important impact on biota feeding on algae in adjacent lakes. The period of serious acidification of the High Tatra lakes in the 20th century was accompanied with a decrease of phytoplankton biomass to nearly zero values. The inflow of cryoflora (probably mainly *Cr. nivalis* subsp. *tatrae*) into Ohrůžle Lake was thus believed to be the key food supply for planktonic calanoid copepod *Arctodiapomus alpinus* IMHOF that enabled its survival despite of the acidification-induced processes of oligotrophication (HOŘÍČKA et al. 2006).

**Morphology and ultrastructure**

The aplanozygote sizes of ‘rediscovered’ populations of *Sc. tatrae* (16.4–24 × 10–16.2 µm) fit to the original description (18–24 × 12–15 µm; KOI 1965). The morphological similarity of ’*Sc. tatrae*’ and *Cr. nivalis* (formerly *Sc. nivalis*) was confirmed earlier by a cluster analysis where all species with *Scotiella*–like morphology were included (KOMÁROVY 1982). The striking difference is a morphologic variation: most commonly 10 to 12 flanges per cell in cross–section typical are a constant feature for *Cr. nivalis* subsp. *tatrae*. On the contrary, *Cr. nivalis* aplanozygotes from the Austrian Alps have usually 8 to 9 flanges only (REMIAS et al. 2010). In both taxa, the organization of flanges is consistent, except for the rare case of two flanges reaching from opposite cell poles and finally touching at the equatorial region of the cell (Fig. 16), which has not been observed in *Cr nivalis* from Alps. The aplanozygote sizes of the new subspecies were slightly smaller than those of the relative from the Alps. However, a large variability of aplanozygote lengths among samples of *Cr. nivalis* (Fig. S1, Table 4) is probably reflecting individual growth conditions depending on the date of sampling in course of the season, and furthermore the abiotic microhabitat and weather conditions before sampling. The known life cycle of *Cr. nivalis* comprises vegetative flagellates which cleave into two, four, eight or sixteen daughter cells. Daughter cells behave as zoospores or gametes. Generative fusion of two gametes leads to development of a quadriflagellate motile zygote. After a while, loss of flagella takes place (transformation to an aplanozygote), the primary cell wall is shed and exposing the developing secondary wall structures of the zygote (HOHAM & MULLET 2001).
intracellular reorganisation may be following: this raise in plastid membrane surface to volume ratio provides probably advantages due to better metabolite exchange for survival in drastically changing cold habitats at the end of growing season. Also the development secondary cell wall flanges during aplanozygote maturation can be generally regarded as a mechanism of mechanical protection, which becomes effective after complete snowmelt and cells are subject to draught when exposed at bare rock surfaces (Remias 2012).

Other genetically and morphologically related snow algal species

Zygotes of *Cr. cf. nivalis* have been recorded from all continents except Australia (Kol 1968; Duval et al. 1999). However, according to 18S rDNA and rbcL phylogeny, *Cr. nivalis*-like aplanozygotes were found in at least four independent lineages, and each lineage is located in a different geographic region (Japan, Svalbard, North America and Europe; see the phylogenetic tree in Fig. S13 in Matsuzaki et al. 2015), suggesting a geographical barrier of dispersion or isolation. Another explanation could be an independent snow colonization several times in the evolution history of *Cr. cf. nivalis*.

Zygotes of the snow alga *Cr. pichinchae* from the North America were reported to resemble *Sc. tatrae* in number of flanges per cell ([6]–11(15)) and cell size (17–20 × 11–14 µm) (Hoham 1975). Accordingly, Hoham & Mullet (1978) suggested in their taxonomic notice that *Sc. tatrae* is a zygotic stage of *Cr. pichinchae*. However, strain *Cr. pichinchae* UTEX SNO 33 is well separated from the *Cr. nivalis* from the Austrian Alps according to 18S phylogeny (see Fig. S13 in Matsuzaki et al. 2015). Independent evolution can be inferred also from ecology, because *Cr. pichinchae* prefers shaded habitat under heavy forest canopy (Hoham 1975) whereas *Cr. nivalis* subsp. *tatrae* causes blooms at exposed sites above timberline. *Cr. hohamii* Ling & Seppelt is another morphologically related snow alga, however with a higher numbers of usually undulating flanges per cell [12–16(20)] (Hoham et al. 1983; Ling & Seppelt 1998). Additionally, it has, like in the case of *Cr. pichinchae*, a different molecular affiliation than *Cr. nivalis* subsp. *tatrae*. The 18S rDNA sequence of *Cr. hohamii* UTEX SNO 67 represents an independent lineage (see Fig. S13 in Matsuzaki et al. 2015) which is also supported by a different geographical distribution (Hoham et al. 1983, Ling & Seppelt 1998). The sister lineage to *Cr. nivalis* from the Austrian Alps (and thus to *Cr. nivalis* subsp. *tatrae*) is represented by *Cr. polypetra* (based on 18S phylogeny in Matsuzaki et al. 2015), a snow alga described from Maritime Antarctica (Fritsch 1912). The putative vegetative hyponblasts of this species occur in coastal, penguin-influenced places and differ morphologically in possessing a very high number of flanges (18–23 per cell; Akiyama 1979), in preference of neutral to alkaline pH
(6.7–8.1; Ling & Seppelt 1998) and in elevated supply of nutrients from bird colonies (electrical conductivity 56–950 μS cm⁻¹ in Ling & Seppelt 1998; 42–95 μS cm⁻¹ in Remias et al. 2013). Contrary, habitat conditions Cr. nivalis subsp. tatrae are characterised by low conductivities and slightly acidic pH (Table 3).

Recent approaches use effective high-throughput sequencing protocols to elucidate OTUs (operational taxonomic units) and their abundances in whole cryoflora communities, thus describing the biodiversity of snow microorganisms (Lutz et al. 2016). Apparently, the use of a single gene for eukaryotes, especially when only sections of the conservative 18S rDNA are available by this method in the context of Chlorophyta, does not provide sufficient resolution to reflect the actual taxonomic richness. Cr. nivalis subsp. tatrae is an example that a ‘rare’ or regionally distributed taxon could be overseen depending on the used gene, because clear morphologic differences to the standard form of Cr. nivalis cannot be evaluated by using 18S rDNA.

Pigment composition
The survival of microalgae at exposed, nutrient-poor snow surfaces frequently goes along with the accumulation of secondary carotenoids (Remias 2012). One the one hand, these pigments effectively absorb excessive visible and near ultraviolet irradiation, thus protecting the photosystems against photoinhibition (Bidigare et al. 1993). On the other hand, carotenoids are nitrogen-free compounds which can be abundantly synthesized also in nitrogen-poor habitats. So far, Chlorophycean snow algae have been reported to accumulate exclusively astaxanthin and derivatives, which are stored in cytoplasmic lipid globules (Remias et al. 2016). It is the same with Cr. nivalis subsp. tatrae, which’s pigments have, to our knowledge, been analysed for the first time. Generally, the cellular amount of astaxanthin in microalgae is illustrated as a relative ratio to chlorophyll (Bidigare et al. 1993; Remias & Lutz 2007). This ratio reflects either the level of aplanozygotes maturity during life cycle (the higher ratio, the older the cells; Remias et al. 2010), or in a tentative chemotaxonomical approach, it shows a distinct range of ratios found for certain clades or taxa. For example, the astaxanthin to chlorophyll ratio was three times higher for mature aplanozygotes of Cr. nivalis subsp. tatrae than that of comparable cells of Cr. nivalis from the Austrian Alps (Remias et al. 2010). This suggests a higher tolerance to excessive irradiation for the former population, but still five times lower values in comparison with dark red cells of the common snow alga Cd. nivalis, which is usually found at the most exposed high alpine and polar sites (Remias et al. 2005). The relative proportions of pigment classes in aplanozygotes of Cr. nivalis subsp. tatrae and Cr. nivalis (Remias et al. 2013) slightly differ, containing twice as many secondary carotenoids, but one third less primary carotenoids and one quarter less chlorophylls for the former alga. In both cases, astaxanthin was the only secondary carotenoid. Accordingly, the cellular astaxanthin content reflects the specific snow coloration such as brownish–red for subsp. tatrae or blood–to crimson red snow for Cd. nivalis (Remias et al. 2005).

Photosynthesis
The adaptation of Cr. nivalis subsp. tatrae to rather high irradiation conditions can be deduced not only from abundant levels of astaxanthin, but also from its photosynthetic performance, such as high light saturation point (Iₚ = 149), light saturation rate (ETRₜₐₛₙₜ = 21.8) and the irradiance level from which on, photoinhibition takes place (600 μmol photons m⁻²·s⁻¹). Moreover, photosynthesis showed positive rETR values even at irradiance levels (< 2100 μmol photons m⁻²·s⁻¹) close to full sunlight (up to 2500 μmol.m⁻²·s⁻¹ at Ladővė Lake with an elevation of 2057 m, the High Tatra Mountains; unpublished results from the automatic meteorological station). Cells may be subject to such irradiances and have to cope with photoinhibition when they are accumulated on snow surface due to proceeding snow melt. Additionally, this species is also able to perform well under low light conditions: Since penetration of irradiation decreases logarithmically with depth of snow column (Gorton & Vogelman 2003), a high alpha value of 0.19 supports cells located a few centimetres below the snow surface to perform positively. In comparison, mature aplanozygotes of Cr. nivalis from the Austrian Alps seems to be saturated by light at slightly lower irradiance levels (Iₚ = 106) what is probably related to their lower astaxanthin content. Flagellate stages of Cr. nivalis are lacking secondary pigments (Hoham & Mullet 1977). One of their protective mechanisms against high irradiance is believed to be the active migration to deeper parts of the snow pack (Kvíderová 2010). Diurnal decrease of rETR and alpha value during noon and successful recovery a few hours later in response to irradiance at open sites was documented for vegetative flagellates tentatively designed as Cr. cf. nivalis (Kvíderová 2010). The preference of low irradiances (about 100 μmol photons.m⁻²·s⁻¹) for flagellate stages of Cr. nivalis originating from shady sites was demonstrated (Stibal 2003). Similar irradiance levels below tree canopy were found in the upper 5 cm of snowpacks containing populations of Cr. hohamii and Cr. nivalis in the North America (Hoham et al. 1998) and were proven to be optimal for mating of Cr. tughilensis and Cr. chenangoensis at lab conditions (Hoham et al. 2006). In our study, we have shown physiological borders for photosynthesis of aplanozygotes of Cr. nivalis subsp. tatrae, whereas growing optima of vegetative stages remain unknown due to a lack of cultures.

Fatty acid composition
The desaturation of FAs in biomembranes is considered as one of the main mechanisms of adaptation and acclimation of photosynthetic microorganism to cold
environments (Morgan–Kiss et al. 2006). The number and position of double bonds together with FA chain length markedly influence melting point (MP) of FA and thus fluidity, permeability and stability of biological membranes (e.g. MP = −11.58 °C for α–linolenic acid; MP = 65.45 °C for palmitic acid, 16:0; Knotte & Dunn 2009). For example, the levels of oleic acid (18:1n9) and α–linolenic acid (18:3n3) decreased in thylakoid membranes upon increase of exposure temperatures from low to mesophilic (Lukéš et al. 2014). The α–linolenic acid as the dominant unsaturated FA in cells of Cr. nivalis subsp. tatrae (19.3 % in total lipids) is consistent with profiles of other Chloromonas species harvested from snow (Cr. pichinchae, 22.9–25.7%; Řezanka et al. 2014) or of a strain kept in nitrogen deficient medium (Cr. nivalis CCCryo 005–99, ~35%; Spijkerman et al. 2012). A high contribution of this FA seems to be typical for all 22 Chloromonas species screened in the study of Lang et al. (2011) (average 27 %). On the contrary, snow algae from a different clade related to Cd. nivalis, show a dominance of oleic acid (~59%; Bidgeare 1993; ~45%; Spijkerman et al. 2012), suggesting that phylogenetic aspects can play a role in native FA patterns within the Chlamydomonadales. This study showed that FAs containing C16 and C18 chains with 0 to 4 double bonds were dominating in aplanozygotes of Cr. nivalis subsp. tatrae. A different profile with dominating medium to long chain FAs of C11 to C18 chains with 0 to 5 double bonds were found in snow dwelling flagellates of Cr. brevispina (Fritsch) Hoham, Roemer & Mullet (Řezanka et al. 2008). The high level of PUFAs of Cr nivalis subsp. tatrae (48%) is comparable with field samples of other Chloromonas species from snow (Cr. brevispina, 74%, Řezanka et al. 2008; Cr. pichinchae, 59–70%, Řezanka et al. 2014) and reflects a suitable strategy for life at temperatures close freezing point (Kvíčerová 2010). The proportion of PUFAs in Cr. nivalis subsp. tatrae was nearly 70% in lipid groups associated with membranes, and it was twice lower in neutral lipids which are likely deposited in cytosolic lipid bodies as storage products (Thompson 1996). Generally, this is in agreement with data published on other microalgae, where PUFA content of the glycolipid and phospholipid fractions was higher than that of the neutral lipid fraction (e.g. Henderson et al. 1998). However, this pattern is extremely pronounced in Cr. nivalis subsp. tatrae suggesting that the high accumulation of PUFAs in membranes is important for its survival in the harsh snow environment.

CONCLUSION
In summary, Cr. nivalis subsp. tatrae can be presumably regarded as an endemic variation of a common cryoflora species. It has been found so far only in the High Tatra Mountains (Slovakia). According to the analysis of several molecular markers, it is closely related to Cr. nivalis from the Austrian Alps. Despite having similar physiological performance and ecological requirements, they differ in the abundance of secondary carotenoids and certain morphological traits. Aplanozygotes (Cr. hohamii and Cr. pichinchae) and putative vegetative hypnoblasts (Cr. polyptera) of further morphologically similar species living in snow can be distinguished by different habitat ecology, molecular traits and morphological details like number of flanges. High levels of polyunsaturated fatty acids were detected especially in membrane lipids. The taxonomic affiliation of Cr. cf. nivalis–like species inhabiting further localities in the High Tatra Mountains listed in (Kol 1975a, b) and Lukáský (1994) is still unclear. A phylogenetic position of other Scotiella species associated with snow (e.g. Sc. norvegica Kol) is lacking. To our experience, establishing cultures from aplanozygotes remains difficult. A polyphasic approach including molecular analysis of multiple DNA regions, description of ecology, cytology and physiology might help to reveal species identity of snow algae and evaluate biodiversity of polar and alpine regions.

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Supplementary material

the following supplementary material is available for this article:

Table S1. Collection data and sample codes of mature aplanozygotes of Chloromonas nivalis from exposed snow fields in the Austrian Alps.

Fig S1. Comparison of lengths of mature aplanozygotes between Chloromonas nivalis subsp. tatrae (LP01) and Chloromonas nivalis from the Austrian Alps (P12, P24/DR4, DR43, GK05, GK09).

Fig S2–S4. Comparison of lengths, widths and length to width ratio (L:W) of mature aplanozygotes between Chloromonas nivalis subsp. tatrae and Chloromonas nivalis from the Austrian Alps (five samples).

This material is available as part of the online article (http://fottea.czechphycology.cz/contents)