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1 Charophytes in warm springs on Svalbard  
2 (Spitsbergen): DNA barcoding identifies *Chara aspera*  
3 and *Chara canescens* with unusual morphological traits  
4  
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15 **Running head:** Charophytes in warm springs on Svalbard (Spitsbergen)  
16

17 **Keywords:** Svalbard, Spitsbergen, Chara, matK, barcoding  
18  
19

20 **Abstract**

21 The Troll springs are warm springs on Svalbard (Spitsbergen). Charophytes were collected in  
22 the years 1910, 1912, 1958, 1992/1993, and 2018. However, since the *Chara* samples showed  
23 unusual morphological traits, there were doubts with respect to species identity. We here use  
24 DNA barcoding to show that there occur two *Chara* species in the Troll springs: *Chara*  
25 *aspera* and *C. canescens*.  
26  
27

28 **Biographical notes**

29 Anders Langangen is cand.real. from the University of Oslo with a thesis on Norwegian  
30 charophytes, and a retired lecturer from Oslo Cathedral school. He has worked with  
31 freshwater algae and specially with charophytes since 1968. Contribution: study design,  
32 manuscript writing.

33 Andreas Ballot is a senior scientist at the Norwegian Institute for Water Research. He is  
34 mainly interested in the ecology and phylogeny of phytoplankton and macrophytes.  
35 Contribution: genetic and phylogenetic analyses, manuscript writing.

36 Petra Nowak is a marine biologist at the University of Rostock specializing in the biodiversity  
37 and evolution of macroalgae. Besides the taxonomical aspect, she is interested in molecular  
38 and morphological approaches towards understanding the acclimation and adaptation  
39 processes of macroalgae. Contribution: sequencing, manuscript writing.

40 Susanne C. Schneider is a senior scientist at the Norwegian Institute for Water Research, and  
41 adjunct professor at the Norwegian University of Life Sciences. She is mainly interested in the  
42 ecology of macrophytes and benthic algae. Contribution: manuscript writing.

43

44

## 45 **1. Introduction**

46 In 1910 and 1912, the Norwegian scientist Adolf Hoel collected a number of algae in the  
47 Troll-springs, a group of warm springs in Bockfjorden (79°25'N, 13°17'E) on Spitsbergen,  
48 Svalbard (Hoel and Høltedahl 1911, Strøm 1921) (Fig.1). One *Chara* taxon was found in the  
49 springs, and sent to professor Otto Nordstedt in Lund, Sweden, one of the most renowned  
50 charophyte experts at that time. He determined the species as *Chara aspera* and described it  
51 (invalidly) as *C. aspera* f. *spitsbergensis* (Hoel and Høltedahl 1911). The valid description of  
52 *Chara aspera* Willd. f. *spitsbergensis* Nordstedt, based on material collected in 1912, was  
53 given in Strøm (1921).

54 In 1958, Niels Foged collected specimens of *Chara* in the same springs (Foged 1964). They  
55 were determined by the Swedish charologist Henning Horn af Rantzien, who “considered it  
56 as a rather peculiar form of *C. canescens* Lois., but at the same time he said he felt uncertain  
57 about the definition” (Foged 1964). This finding is described in Langangen (1979), where the  
58 taxon is stated to be *Chara canescens*.

59 In 1992 and 1993 charophytes were again collected from the springs by Sissel Aarvik from  
60 the Governor of Svalbard. In this sampling, two *Chara* taxa were found, which both were  
61 suggested to be different forms of *Chara canescens*. They were described and discussed in  
62 Langangen (2000). In this work the taxon *C. aspera* f. *spitsbergensis* Nordstedt was given the  
63 new combination *C. canescens* f. *spitsbergensis* (Nordstedt) Langangen (nom. inval.). The  
64 other taxon was given the name *C. canescens* subsp. *hoelii* Langangen, and described as a  
65 subspecies. We here describe and interpret the results of a new collection of *Chara* material  
66 from the Troll springs in 2018. Our aim was to confirm and if necessary, correct the species  
67 identity using DNA barcoding.

68

## 69 **2. Material and Methods**

### 70 **2.1 Study site and sampling of *Chara* material**

71 According to Hoel (1914) the Troll-springs have 14 individual ground-water sources and  
72 consist of large sinter terraces. Charophytes were found and collected in two springs named  
73 Spring 4 and Spring 6 (Hoel 1914). Spring 4 is the largest of the springs, approximately 11 m  
74 long, 7 m wide and 2 m deep. Two photos from spring 4, taken in 1912 and 2018,  
75 respectively, illustrate that the Troll springs have changed very little in more than hundred  
76 years (Fig 2). Spring 6 is approximately 3 m long, 2 m wide and 1 m deep. There is a  
77 considerable flow of water from this spring, estimated by Hoel (1914) to be at least 100 litres  
78 per minute. The outlet of spring 6 was overgrown by filamentous algae in 2018, and the  
79 spring itself was dominated by reproductive *Chara canescens*.

80

### 81 **2.2 Physical and chemical characteristics of the Troll springs**

82 Water temperature, conductivity and Ca-content were measured in the springs, using the  
83 following methods; specific conductivity was measured in 1992-93 with a Hach conductivity  
84 meter (Model 44600/CND/TDS) and in 2018 with a Milwaukee SM 301 ECmeter (range 0-  
85 1990  $\mu\text{S}/\text{cm}$ ). Calcium was measured in 1992-93 with Aquamerck 11110 Calcium test and in  
86 2018 with the Calcium MColortest from Merck. Temperature was measured in 1912 and  
87 1992-93 with unknown types of thermometers. In 2018 we used a Ziel Mercury L0110/10 305  
88 mm yellow back thermometer.

89

### 90 **2.3 DNA barcoding**

91 Two different methods were used to isolate genomic DNA from *Chara* samples investigated  
92 in this study. An overview over which samples were analyzed with which method is given in  
93 Table 1. The sequence data were deposited in the European Nucleotide Archive (ENA) under  
94 the accession numbers given in Table 1.

95 Method A: Genomic DNA from *Chara* material was isolated after Schneider et al. (2016).

96 PCR for the matK gene was performed on a Bio-Rad CFX96 Real-Time PCR Detection  
97 System (Bio-Rad Laboratories, Oslo, Norway) using the iProof High-Fidelity PCR Kit (Bio-  
98 Rad Laboratories, Oslo, Norway). Amplification of the matK gene region was conducted

99 using the primers F-Chara (agaatgagcttaaacaaggat) and R-Chara (acgatttgaacatccactataata).  
100 The following cycling protocol was used: one cycle of 5 min at 94 °C, and then 35 cycles  
101 each consisting of 10 s at 94 °C, 20 s at 62 °C, and 20 s at 72 °C, followed by a final  
102 elongation step of 72 °C for 5 min. PCR products were visualized by 1.5% agarose gel  
103 electrophoresis with GelRed staining and UV illumination. For sequencing the same primers  
104 and the intermediate primers charaintF (gatggctattcaagcagga), charaintR (ctaccgataagttcgtct),  
105 charaBt2f (datatggcaacaycaaaaagac) and charaBT2R (atacagacatgcagcytt) were used.  
106 Sequences were analysed and aligned using Seqassem (version 04/2008) and Align (version  
107 03/2007) MS Windows-based manual sequence alignment editor (SequentiX - DigitalDNA  
108 Processing, Klein Raden Germany) to obtain DNA sequence alignments, which were then  
109 corrected manually. For each PCR product, both strands were sequenced on an ABI 3730  
110 Avant genetic analyser using the BigDye terminator V.3.1 cycle sequencing kit (Applied  
111 Biosystems, (Applied Biosystems, Thermo Fisher Scientific Oslo, Norway) according to the  
112 manufacturer's instructions.

113 Method B: Preparation of total DNA was performed using the DNeasy Plant Mini Kit  
114 (Qiagen, Hilden, Germany), following the manufacturer's protocol. Amplification of the *matK*  
115 gene region was performed with a Taq PCR Master Mix (Qiagen, Hilden, Germany), using  
116 the primers matK-F2 (aatgagcttaaacaaggattc) and matK-R1b (gcagccttatgaattggatagc). The  
117 following PCR protocol was used: 10 cycles of one minute each at 94° C, 55° C, and 72° C,  
118 followed by one minute each at 94° C, 52° C, and 72° C for 25 cycles. The amplified DNA  
119 was purified with the Biometra-innuPrep Gel ExtractionKit (Analytik Jena, Jena, Germany)  
120 according to the manufacturer's instructions and was sequenced directly on a 3130×L  
121 GeneticAnalyzer (Applied Biosystems, NY, USA) using the BigDye terminator V.1.1 cycle  
122 sequencing kit (Applied Biosystems, Thermo Fisher Scientific, Darmstadt, Germany).  
123 Sequencing primers were identical to the primers that were used for the PCR reactions.  
124 Achieved sequences were proofed and manually edited using the BioEdit programme (Hall  
125 1999).

126

## 127 **2.4 Phylogenetic analysis**

128 Segments with highly variable and ambiguous regions and gaps, making proper alignment  
129 impossible, were excluded from the analyses. In addition to two samples collected in the Troll  
130 springs in 2018, a matK set containing 38 other *Chara* sequences (Table 1), including a  
131 sample of the 1992 sampling in the Troll-springs, and 1023 nucleotide positions was used for

132 phylogenetic analysis. *Nitellopsis obtusa* (AY170447) was used as an outgroup taxon in the  
133 matK tree. The dataset was analyzed using the maximum likelihood (ML) algorithm in  
134 MEGA version 7 (Kumar, Stecher, and Tamura 2016). The method selected GTR+G as the  
135 best-fitting evolutionary model for the matK gene region. ML analyses were performed with  
136 1000 bootstrap replicates in MEGA version 7 (Kumar, Stecher, and Tamura 2016).

137

### 138 **3. Results**

#### 139 **3.1 Physical and chemical characteristics of the Troll springs**

140 Water temperature was around 20 °C in spring 4, and around 26 °C in spring 6 in all years  
141 (Table 2). Conductivity varied slightly but was above 1300 µS/cm in all years. This indicates  
142 slightly brackish water. Calcium concentrations of 100 mg/L and above indicate hard water in  
143 both springs. All water samples are from August in the respective years, and we do not have  
144 any information on water chemistry from other months.

145

#### 146 **3.1 Barcoding results**

147 *Chara aspera* and *Chara canescens* were separated into two monophyletic groups supported  
148 by bootstrap values  $\geq 99$  (Fig. 3). All other taxa used in the present study were clearly  
149 separated from these two large groups. Sample S117 from spring 4 clustered with *C. aspera*,  
150 while sample S118 from spring 6, and sample AL02 which was sampled in spring 6 in 1992,  
151 clustered with *C. canescens*. Despite the well supported clusters, both *C. aspera* and *C.*  
152 *canescens* exhibited some degree of variability in the matK sequences (Fig. 3). However,  
153 sample S117 collected in the Troll springs had identical sequences to samples of *C. aspera*  
154 from Sweden, Norway, the UK, Germany and France, while samples S118 and AL02 were  
155 identical to samples of *C. canescens* collected in Sweden, Germany, Spain and Italy  
156 (Sardinia).

157

### 158 **4. Discussion**

#### 159 **4.1 Implications for taxonomy**

160 The barcoding results clearly indicate that there are two *Chara* species in the Troll-springs on  
161 Svalbard: *Chara aspera* and *Chara canescens*. The samples collected from Svalbard were

162 genetically identical to other samples of the same species from several countries in Europe.  
163 This has the following implications for taxonomy:

164 **A. Not accepted taxa:**

165 a. *Chara aspera* Willd. f. *spitsbergensis* Nordstedt in Strøm 1921.

166 Nordstedt (in Strøm 1921) gives a latin diagnosis of the new forma. The description is a  
167 combination of characters from what we now know are two species, *C. aspera* and *C.*  
168 *canescens*. This also agrees with the fact that the original material, found in 1912 is a mixture  
169 of both taxa. We designate a lectotype as the part of the original collection which matches  
170 with what we now know is *Chara aspera* and which consists of only sterile specimens:  
171 Svalbard, Bockfjorden, the Troll springs, 1912-08-03, A. Hoel, coll. (O, p.p.). This taxon is  
172 regarded as a synonym to *Chara aspera*.

173 b. *Chara canescens* subsp. *hoelii* Langangen.

174 This taxon is described and discussed in Langangen (2000). According to our barcoding  
175 results this is *Chara aspera*. We regard the taxon as an aberrant form of *Chara aspera*, due to  
176 both the morphology and the missing support for a subspecies in the genetic analysis (Fig. 3).  
177 This taxon is regarded as a synonym to *Chara aspera*.

178

179 **B. Accepted taxa:**

180 a. *Chara canescens* Desv. & Loisel. (Fig. 4)

181 *Chara canescens* is an exceptionally variable species, and many forms have been described  
182 (Schubert and Blindow 2004). The specimen we found in the Troll springs in 2018 looked the  
183 same as those found in 1992/1993 (Langangen 2000). Therefore, the morphology of this  
184 species in the Troll springs seems to be stable. For the sake of completeness, we here repeat  
185 (in condensed form) the description given in Langangen (2000).

186 Plants were unbranched to strongly branched, only slightly encrusted in part of the whorls.  
187 The axes were 400 to 750 µm in diameter, and the internodes 2 to 15 mm long, 1 to 4 times  
188 the length of the branchlets. The stem cortex was regularly haplostichous in younger  
189 internodes, and irregular or absent from older internodes. Spine cells were acute, often short  
190 but in some cases up to 1.5 times the diameter of the axes. Stipulodes were in 1-2 tiers, 2 per  
191 branchlet. Branchlets were 7-9 in a whorl, up to 4 mm long, slightly connivent, with 3 - 4  
192 segments, and with end segments of up to 3 ecorticate cells. The end segments were up to 2

193 mm long, and longer than the corticate segments. The branchlet cortex was more or less  
194 regular. Bract cells were verticillate and ca. 500  $\mu\text{m}$  long. Bracteoles were up to 1 mm long.  
195 The whorls were often “nestlike” (Fig. 4), consisting of relatively short branchlets filled with  
196 oogonia and black ripe oospores. These whorls were 3.2- 5.0 mm wide. The plants were  
197 dioecious, and only oogonia were found. Oogonia (675-825  $\mu\text{m}$  long, 275-450  $\mu\text{m}$  wide, with  
198 9-10 convolutions, coronula 50  $\mu\text{m}$  long and 125  $\mu\text{m}$  wide) were found adjacent to both  
199 corticate and ecorticate internodes, but were most common on the two lowest branchlet nodes.  
200 Oospores were black, ovoid to elliptical, (475-600  $\mu\text{m}$  long to 325-400  $\mu\text{m}$  wide, and with 9  
201 ridges). The oospores were extremely abundant.

202

203 b. *Chara aspera* Willd. (Fig. 5)

204 *Chara aspera* from the Troll springs is difficult to determine morphologically, because the  
205 plants generally are ecorticate and sterile. Plants were up to 14 cm long, mildly to strongly  
206 encrusted with calcium carbonate. Axes to 350  $\mu\text{m}$  in diameter, with internodes up to 2 cm  
207 long, ecorticate. Stipulodes were not observed. Branchlets 7-9, up to 15 mm long, with 5-10  
208 segments. Cortex of branchlets mostly rudimentary or missing, often with cortical cells  
209 standing out from the branchlet internodes. Branchlets were tipped with 2-3 ecorticate cells.  
210 In some whorls accessory branchlets are found in rows above and/or below the primary  
211 branchlets. Gametangia have not been found. Bulbils were one-celled, acute, ovoid, 750-1500  
212  $\mu\text{m}$  long and up to 600  $\mu\text{m}$  wide. They occurred as solitary bulbils, in pairs or in groups of  
213 three or four.

214 The specimens collected in the Troll springs were genetically identical to specimens of *C.*  
215 *aspera* collected in several countries in Europe (Fig. 3). We therefore regard the specimens  
216 from the Troll springs as aberrant forms of *Chara aspera*, likely caused by the “extreme”  
217 environment (warm springs in a polar environment).

218

219 **4.2 Species distribution and possible survival in an extreme Northern environment**

220 Both taxa, *C. aspera* and *C. canescens* are mesohaline species (Krause 1997), and this  
221 matches well with the conductivity measured in the Troll springs (Table 2). The occurrence of  
222 two *Chara* species so far north is, however, remarkable. The closest known locality of *Chara*  
223 *aspera* is at the Norwegian coast approximately 900 km south of the Troll springs, while the  
224 closest known locality of *Chara canescens* is approximately 1000 km south of the Troll



225 springs (Langangen, 2007). The closest currently known localities where both species occur  
226 are Alstahaug municipality in Nordland county (Gaarder et al. 2012), at a distance of  
227 approximately 1300 km from the Troll springs on Svalbard, and Eide municipality in Møre  
228 and Romsdal county (Langangen, Gaarder, and Jordal 2001), at a distance of approximately  
229 1800 km from the Troll springs (Fig. 6). Although this must remain speculative, both species  
230 may have arrived in the Troll springs via long distance dispersal by birds. Several species of  
231 geese, e.g. barnacle goose (*Branta leucopsis*), have migration routes from Scotland and South  
232 Norway, with resting places in Nordland county before breeding on Svalbard, including the  
233 area around the Troll springs (Griffin, Rees, and Hughes 2011). Fig. 6 shows that Eide and  
234 Alstahaug, where both *C. aspera* and *C. canescens* occur, are on the migration route of  
235 barnacle goose to breeding places on Svalbard.

236 It also is remarkable that the *Chara* species survive in the harsh Northern environment. In this  
237 area, the polar night (i.e. the time of darkness during which the sun never is above the  
238 horizon) lasts from October 26 to February 15 (data for Longyearbyen). The polar night is  
239 black in cloudy weather but can be surprisingly bright in clear weather and when there is  
240 moonlight. Several studies analysed light dependency of *Chara* photosynthesis, growth, or  
241 reproduction (e.g. Blindow and Schütte 2007, Schaible and Schubert 2008, Schneider et al.  
242 2015), but to our knowledge, no studies on how charophytes may survive several months of  
243 almost complete darkness exist. To our knowledge, nobody has ever collected samples in the  
244 Troll springs during winter. Consequently, it is unknown if the charophytes survive winter as  
245 green plants. Whether or not moonlight may be sufficient to sustain *Chara* photosynthesis is,  
246 to our knowledge, unknown. According to local, unpublished observations, water temperature  
247 in the Troll springs does not fall below zero, i.e. the springs stay, at least in parts, ice-free. On  
248 the one hand, the lack of any sunlight over a period of about four months should make a  
249 vegetative survival difficult. On the other hand, however, *Chara* species can in Nordic lakes  
250 survive vegetatively for four months underneath snow-covered ice (which almost completely  
251 blocks sunlight; own observations). However, charophytes are well known pioneer plants,  
252 which may persist during unfavourable conditions, e.g. the desiccation of temporal lakes, in  
253 form of resting stages. Charophytes are known to be able to quickly regrow from oospores,  
254 bulbils, or starch reserves in axial nodes (Krause 1997).

255 *Chara canescens*, as the only parthenogenetic charophyte taxon (Schaible et al. 2008),  
256 generally produces a large number of oospores, and this was also the case in the samples from  
257 the Troll springs. After the oospores ripen, *C. canescens* plants usually degenerate (Schubert

258 et al. 2016), and it therefore is assumed that this species generally regrows each year from  
259 oospores. It is unknown if this also occurs in the Troll springs on Svalbard, but the large  
260 number of oospores which occurred on the *C. canescens* samples indicate that this may be the  
261 case. However, no oospores were observed on *C. aspera* from the Troll springs, neither in  
262 2018, nor in 1992/1993 or in 1912 (Langangen 2000). *C. aspera* therefore seems to either  
263 regrow from bulbils or axial nodes in spring, or survives four months of almost complete  
264 darkness during the polar night in vegetative form.

265

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270 Oslo) for help with nomenclature, and Thomas Gregor for helpful comments on an earlier  
271 version of the manuscript. Barcoding of some other *Chara* taxa was funded by the Norwegian  
272 Biodiversity Information Centre.

273

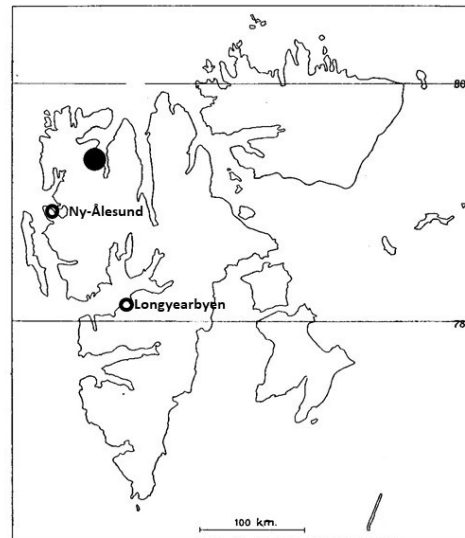
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316 21.
- 317
- 318

319 **Figures**

320



321

322 **Fig. 1.** Location of the Troll-springs on Svalbard

323

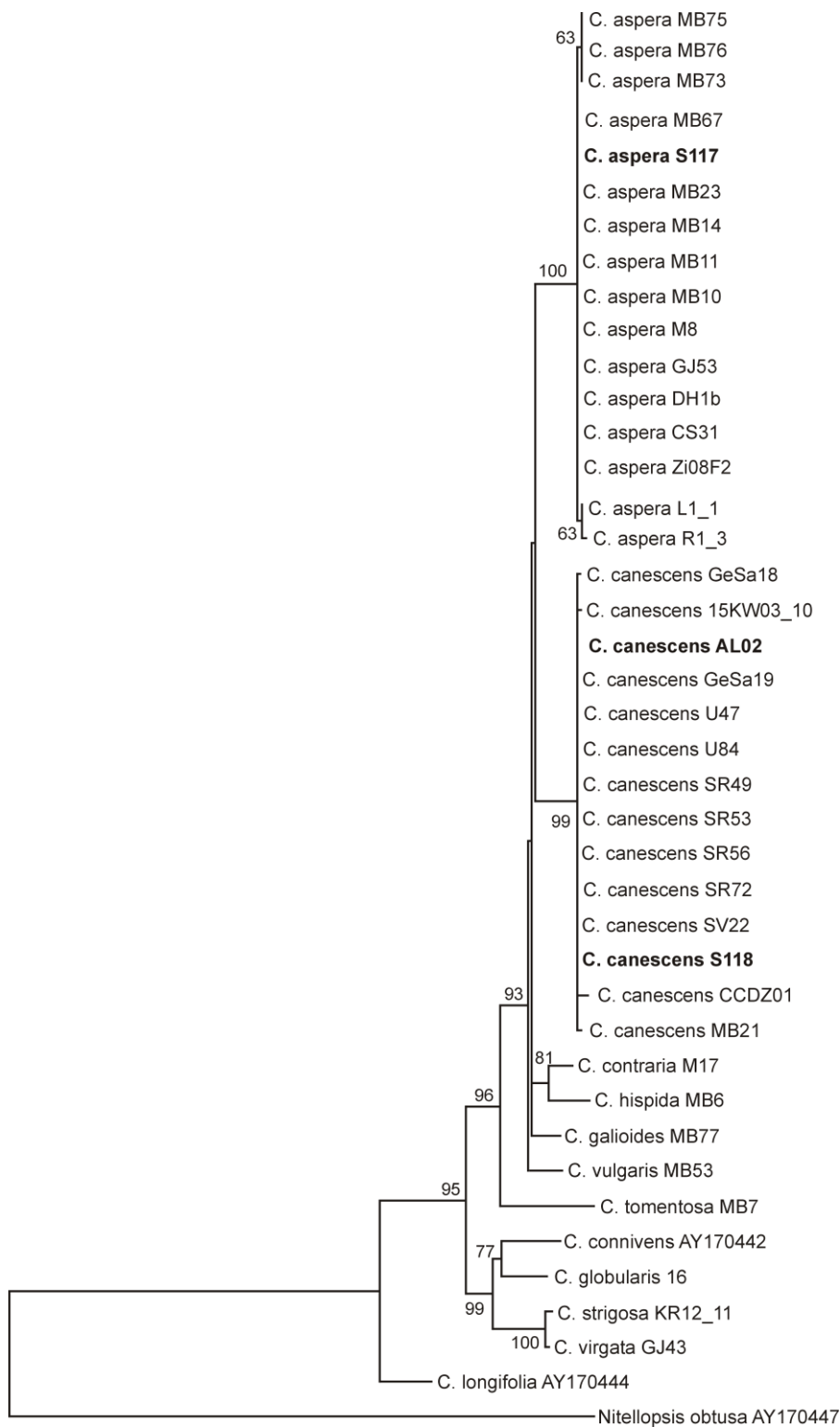


324

325 **Fig. 2.** Troll-spring 4 in 1912, August 3 (top), and 2018, August 15 (bottom). The picture  
326 from 1912 was taken during a second visit, two years after the first collection of algae. Picture  
327 taken by Adolf Hoel, Norsk Polarinstitut (top) and Gunhild Lutnæs (bottom).

328

329



330

0.020

331 **Fig. 3.** Maximum Likelihood tree of the matK gene of *Chara* spp. Bootstrap values above 50  
 332 are included. The scale bar indicates 2% sequence divergence. Sample S117 is from spring 4,  
 333 and sample S118 is from spring 6. Sample AL02 was sampled from spring 6 in 1992, and  
 334 sequences were obtained from herbarium material.

335



336

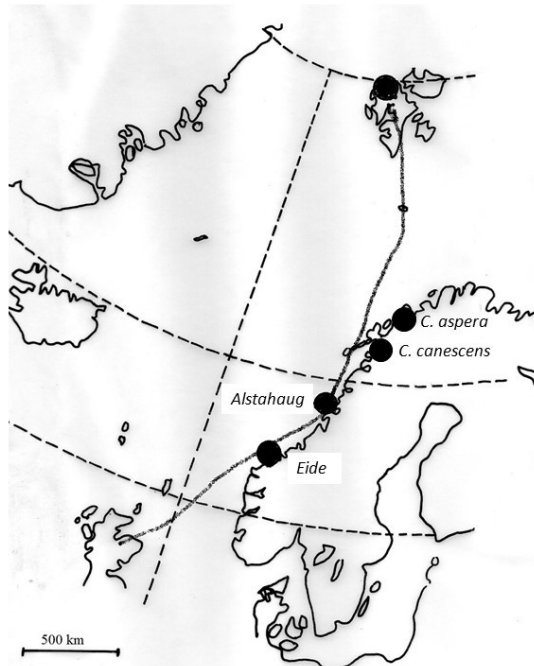
337 **Fig. 4.** *Chara canescens*. Specimens of different length from spring 6. The left specimen is  
338 covered by a brown clayish coating. The picture is from 1992, but the specimen from the  
339 2018 collection looked the same. Picture taken by A. Langangen.

340



341

342 **Fig. 5.** *Chara aspera*. Habitus of three specimens. The picture is from 1992, but the specimen  
343 from the 2018 collection looked the same. Picture taken by A. Langangen.



344

345 **Fig. 6.** Migration route for barnacle goose (*Branta leucopsis*) and the three localities where  
 346 both, *Chara aspera* and *C. canescens* occur (Eide, Alstahaug and the Troll springs).

347

348 **Tables**

349

350 **Table 1.** List of 40 *Chara* individuals (and one *Nitellopsis obtusa*) used in the present study.  
 351 “Method A and B” refers to the method used for DNA-sequencing described in 2.3. Samples  
 352 from the Troll springs are shaded.

Identification	Field ID	Genbank			country	author
		access number	method	coll. year		
<i>C. aspera</i>	MB67	LR134032	A	2005	UK	Willd. 1809
<i>C. aspera</i>	S117	LR134033	A	2018	Norway (Svalbard)	
<i>C. aspera</i>	MB23	LR134034	A	2005	Sweden	
<i>C. aspera</i>	MB14	LR134035	A	2000	Germany	
<i>C. aspera</i>	MB11	LR134036	A	2001	Germany	
<i>C. aspera</i>	M8	LR134037	A	2008	Norway	
<i>C. aspera</i>	GJ53	LR134038	B	2009	Sweden	
<i>C. aspera</i>	CS31	LR134039	B	2012	France	
<i>C. aspera</i>	DH1b	LR134040	B	2011	UK	
<i>C. aspera</i>	Zi08_F2	LR134041	B	2010	Germany	
<i>C. aspera</i>	L1_1	LR134042	B	2015	Germany	
<i>C. aspera</i>	R1_3	LR134043	B	2014	Germany	
<i>C. aspera</i>	MB10	LR134044	A	2000	Germany	
<i>C. aspera</i>	MB73	LR134045	A	2001	France	
<i>C. aspera</i>	MB75	LR134046	A	2001	France	
<i>C. aspera</i>	MB76	LR134047	A	2001	France	
<i>C. canescens</i>	SR49	LR134049	B	2010	Spain	Desv. et Loisel. 1810
<i>C. canescens</i>	SR53	LR134050	B	2010	Spain	
<i>C. canescens</i>	SR56	LR134051	B	2010	Spain	
<i>C. canescens</i>	SR72	LR134052	B	2010	Spain	
<i>C. canescens</i>	GeSa18	LR134053	B	2008	Italy (Sardinia)	
<i>C. canescens</i>	15KW03_10	LR134054	B	2015	Italy (Sardinia)	
<i>C. canescens</i>	AL02	LR134055	A	1992	Norway (Svalbard)	
<i>C. canescens</i>	GeSa19	LR134056	B	2015	Italy (Sardinia)	
<i>C. canescens</i>	U47	LR134057	B	2014	Germany	
<i>C. canescens</i>	U84	LR134058	B	2014	Germany	
<i>C. canescens</i>	CCDZ01	LR134059	B	2011	Germany	
<i>C. canescens</i>	SV22	LR134060	B	2003	Sweden	
<i>C. canescens</i>	MB21	LR134061	A	2005	Greece	
<i>C. canescens</i>	S118	LR134062	A	2018	Norway (Svalbard)	
<i>C. connivens</i>		AY170442				Salzm. ex A. Braun 1835
<i>C. contraria</i>	M17	LR134063	A	2008	Norway	A. Br. ex Kütz. 1845 s. str.
<i>C. galioides</i>	MB77	LR134048	A	2001	France	De Candolle 1813
<i>C. globularis</i>	16	LR134067	A	2009	Macedonia	Thuillier 1799
<i>C. hispida</i>	MB6	LR134064	A	2004	Germany	(L.) Hartm. 1820
<i>C. longifolia</i>		AY170444				(Rob.) R.D.Wood 1965
<i>C. strigosa</i>	KR12_11	LR134068	B	2011	Germany	A. Braun 1847
<i>C. tomentosa</i>	MB7	LR134066	A	2004	Germany	L. 1753
<i>C. virgata</i>	GJ43	LR134069	B	2009	Sweden	Kütz. 1834
<i>C. vulgaris</i>	MB53	LR134065	A	2001	France	L. 1753
<i>Nitellopsis obtusa</i>		AY170447				(Desvaux) J. Groves 1919

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356 **Table 2.** Physical and chemical characteristics of the Troll-springs 4 and 6, where  
 357 charophytes were found. The water sample taken from spring 4 in 1993 was damaged during  
 358 transport from Svalbard. The data from 1912 were taken from Hoel (1914).

	<b>3.8.1912</b>	<b>28.8.1992</b>	<b>16.8.1993</b>	<b>15.8.2018</b>
<b>spring 4</b>				
conductivity [ $\mu\text{S}/\text{cm}$ ]		1620		1380
$\text{Ca}^{2+}$ [mg/L]		125		100
Temp. [ $^{\circ}\text{C}$ ]	21		19	20
<b>spring 6</b>				
conductivity [ $\mu\text{S}/\text{cm}$ ]		1600	1470	1480
$\text{Ca}^{2+}$ [mg/L]		122	130	110
Temp. [ $^{\circ}\text{C}$ ]	26		25	27

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