STABILITY OF SOIL ORGANIC MATTER IN CRYOSOLS OF MARITIME ANTARCTIC: INSIGHTS FROM 13-C NMR AND ELECTRON SPIN RESONANCE SPECTROSCOPY

4 Evgeny Abakumov¹, Ivan Alekseev^{1,2}

¹ Department of Applied Ecology, Saint-Petersburg State University, 199178, 16-line 2, Vasilyevskiy
 Island, Russia

² Otto Schmidt Laboratory for Polar and Marine, Arctic and Antarctic Research Institute, 199397,
 ⁹ Beringa str. 38, Russia

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11 e-mail: e.abakumov@spbu.ru, e_abakumov@mail.ru, st014661@student.spbu.ru

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- 13 Key words
- 14 Antarctica, soil organic matter, stabilization, humic acids
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16 Key points

- 17 Investigation of Antarctic soil organic matter stability
- 18 Humic acids of superficial horizons contain more aromatic carbon
- 19 Humic acids of isolated layers contain more free radicals

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2122 Abstract

23

Previously, the structure and molecular composition of the Antarctic SOM has been investigated 24 25 using ¹³C-NMR methods, which showed that in typical organo-mineral soils the aliphatic carbon prevails over the aromatic one, owing to the non-ligniferous nature of its precursor material. In 26 this study, the soil organic matter (SOM) was analyzed from different sample areas (surface level 27 and partially isolated supra-permafrost layer) of the tundra-barren landscape of the Fildes 28 Peninsula, King George Island, Western Antarctica. We found that the humic acids (HAs) of the 29 30 cryoturbated, buried areas had lower amounts of alkylaromatic and protonized aromatic 31 compounds. In contrast, the HAs from the surface layers contain less alkyl carbon components. The free radical content was higher in the surface layers than in the buried layers due to the 32 33 presence of fresh organic remnants in superficial soil samples. New data on SOM quality from these two representative Cryosols will enable more precise assessment of SOM stabilization rate 34 in sub-Antarctic tundras. Comparison of the ¹³C-NMR spectra of the HAs and the bulk SOM 35 revealed that humification occurs in the Antarctic and results in accumulation of aromatic and 36 carboxylic compounds and reductions in alkylic ones. This indicates that humification is one of 37 38 the ways of soil organic matter stabilization.

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- 40

41 Highlights

42 Stabilization of soil organic matter studied

- 43 Humic acids of superficial horizons contain more aromatic carbon
- 44 Humification is one of the ways of soil organic carbon stabilization
- 46 **1. Introduction**

Polar soils play a key role in global carbon circulation and stabilization as they contain 47 maximum stocks of soil organic matter (SOM) within the whole pedosphere (Schuur et al, 2015). 48 49 Cold climate and active layer dynamics result in the stabilization of essential amounts of organic 50 matter in soils, biosediments, and grounds of the polar biome (Zubrzycki et al, 2014). Global 51 climate changes and permafrost degradation have led to the exposure of huge pools of organic 52 matter to microbial degradation (Schuur et al, 2015) and other environmental risks. Polar SOM 53 represents a vulnerable carbon source, susceptible to remobilization under increasing 54 temperatures (Schuur et al, 2015, Ejarque, Abakumov, 2016). In order to better understand the 55 implications of permafrost SOM for greenhouse gas emissions, accurate knowledge of its spatial 56 distribution, both in terms of quantity and quality (e.g. biodegradability, chemical composition, and humification stage) is needed in addition to effective evaluation of SOM's temporal 57 58 dynamics (Fritz et al, 2014, Vasilevitch et al, 2018).

59 Current estimations of soil organic carbon (SOC) stocks are around 1307 Pg throughout 60 the northern circumpolar region (Hugelius et al, 2014). These amounts surpass previous 61 estimates (Tarnocai et al, 2009) and grossly exceed the total carbon contained in the world's 62 vegetation biomass (460 - 650 Pg) or in the atmosphere (589 Pg) (Tarnocai et al, 2009). 63 However, the aforementioned SOM/SOC stock estimations are still poorly constrained (Hugelius et al, 2014). This uncertainty is largely caused by the estimates having been calculated from 64 observations that are highly spatially clustered (Hugelius et al. 2014) while extensive land areas 65 66 remain uncharacterized due to the logistic difficulties of reaching these sites. Additionally, the calculation of these stocks are based on estimated data on soil bulk density and carbon values 67 derived from dichromate oxidation methods (Abakumov, Popov, 2005, Polyakov et al, 2017). 68

69 The stocks of SOM in the Antarctic are underestimated compared to the Arctic because 70 of the lack of the data for many parts of this continent, due to the high content of stones in the soils and the high variability in the carbon content of the fine earth. Stocks of organic carbon in 71 the Antarctic soil have been reported as 0.5 kg/m^2 in its polar deserts, about 1.0 kg/m^2 in its 72 barrens, up to 3 - 5 kg/m² in the sub-Antarctic tundra, and up to 30 kg/m² in the penguin 73 74 rockeries of the maritime islands (Abakumov, 2010, Abakumov, Mukhametova, 2014, 75 Abakumov et al, 2016). To date, investigation on structural composition of the SOM from both 76 superficial and partially isolated areas has only been performed on Cryosols of the Kolyma 77 lowland (Lupachev et al, 2017), where the organic matter of modern and buried soils vary 78 greatly in terms of their molecular composition and quality.

79 Stability and biodegradability are the key features of SOM that should be taken into 80 account when estimating current and future carbon stocks and organic matter quality and dynamics. Stability is related to humification degree, as more advanced stages in the 81 humification process involve depletion of the labile molecules, as well as an increase in the bulk 82 83 aromaticity, which confers higher stability to the SOM. A number of proxies have been used to trace humification rate and SOM stability, including aromaticity level (Vasilevitch et al, 2018, 84 Kniker, 2007). Also the ratio of C-Alkyl : C-Aryl and C-Alkyl : O-N-alkyl have been 85 successfully used to assess humification degree (Kinker, 2007). C/H ratio from humic acids 86 (HAs) has been used as an index of molecular complexity, as more degrees of conjugation imply 87 less hydrogenation of the carbon chains (Zaccone et al, 2007) and C/N has been used as a 88 measure of histic material degradation (Lodygin et al, 2014). ¹³C-NMR spectrometry provides 89

90 information on the diversity in carbon functional structures (carbon species) and has been used to 91 evaluate changes in SOM during decomposition and humification. More specifically, high phenolic (150 ppm), carboxyl-C (175 ppm) and alkyl-C (30 ppm) groups, combined with low O-92 alkyl carbons, have been associated with advanced humification stages (Zech et al, 1997). So far, 93 94 studies of SOM quality from polar environments have revealed generally lowly-decomposed 95 organic molecules (Dziadowiec, 1994, Lupachev et al, 2017), which preserve much of the 96 chemical character of their precursor material due to slow progress of humification (Davidson 97 and Jansens, 2006). This is very important because polar soils are characterized by the specific 98 composition of the humification precursors.

99 The structure and molecular composition of the Antarctic SOM has been investigated using ¹³C-NMR methods (Beyer et al, 1997, Abakumov, 2017) and it was shown that in typical 100 101 organo-mineral soils the aliphatic carbon prevails over the aromatic one, owing to the non-102 ligniferous nature of its precursor material (Calace et al, 1995). Also, analyses of cryptogam extracts were conducted towards identification of individual organic precursors (Chapman et al, 103 104 1994). This feature was then shown to be typical for soils from different regions of the Antarctic 105 (Abakumov, 2010), including soil formed on the penguin rockeries (Abakumov, Fattakhova, 2015). The northern most soil of Arctic polar biome shows the same trend in organic molecules 106 organization: higher prevalence of aliphatic structures over aromatic ones. The diversity of the 107 108 individual components in aromatic and aliphatic areas is usually higher in Arctic soil because of 109 the increased diversity of humification precursors (Ejarque, Abakumov, 2016, Abakumov, 2010). A selective preservation of the alkyl moieties in the deeper soil layers has been suggested, 110 111 and little transformation processes of the SOM are detectable because soil temperatures are not high enough to stimulate further microbial break-down, even in the summer (Beyer et al, 1997). 112 It has been shown that ornitochoria play an essential role in redistribution of plant remnants in 113 114 the Antarctic (Parnikoza et al, 2016) as birds transport considerable amounts of variably 115 composed organic material within its inland landscapes. However, published data on SOM composition for the Antarctic are rare, and further studies that detail its structural compounds 116 and their distribution are needed. Recently, ¹³C-NMR was successfully used to detail the soils 117 found in endolitic communities in Eastern Antarctica and revealed that endolitic organic matter 118 119 is characterized by a low prevalence of alkyl aromatic compounds (Mergelov et al, 2018).

This study aimed to compare the structural composition of the SOM from both superficial and partially isolated (i.e. buried spots on the border with permafrost) areas and to evaluated the stabilization rate of Antarctic Cryosols. The objectives of the study were: (1) to evaluate the alterations in the elemental compositions of the HAs under partial isolation (2) to assess the ratios of aromatic and aliphatic carbon species in the topsoil and isolated areas; (3) to characterize the biochemical activity of the HAs (e.g. free radical concentration).

126 **2. Materials and Methods**

127 **2.1.** Study sites

King George Island is the largest in the South Shetland archipelago and only around 5% 128 of its 1400 km² area is free of ice (Fig. 1) (Rakusa-Suszczewski, 2002). The Fildes Peninsula and 129 Ardley Island, together around 33 km², comprise the largest ice-free area on King George Island 130 and the second largest of the South Shetland Islands. It has a gentle landscape consisting of old 131 132 coastal landforms with numerous rocky ridges (Michel et al, 2014). According to Smellie et al. 133 (2014), this area mainly consists of lava with small exposures of tuffs, volcanic sandstones, and agglomerates. The climate is cold and humid with a mean annual air temperature of -2.2°C and 134 135 mean summer air temperatures above 0°C for only up to four months (Wen et al, 1994). The 136 mean annual precipitation is 350 - 500 mm/year. The Fildes Peninsula and Ardley Island are

137 among the first areas in maritime Antarctica to become ice-free after the Last Glacial Maximum

138 (Birkenmajer, 1989). The onset of deglaciation in Fildes peninsula started as in the SSI by 8000-

139 9000 ka and

spread during the mid Holocene (Oliva et al., 2016). The patterned ground in this region dates 140 141 from 720 to 2640 BP. In the South Shetland Islands, permafrost is sporadic or non-existent at 142 altitudes below 20 m AMSL and occurs discontinuously in altitudes from 30 to 150 m AMSL 143 (Bockheim et al, 2013). Mosses, lichens, and algae are common to this area along with two vascular plants (Deschampsia antarctica and Colobanthus quitensis). Penguins, seals, and 144 145 seabirds inhabit the coastal areas and greatly impact the soil development. Major cryogenic 146 surface-forming processes in this region include frost creep, cryoturbation, frost heaving and 147 sorting, gravity, and gelifluction (Michel et al, 2014). Eight separate sites on the Fildes Peninsula have been collectively designated an Antarctic Specially Protected Area (ASPA 125) largely due 148 149 to their paleontological properties (Management plan, 2009). The average thickness of the soil is about 15 - 25 cm. Soils from King George Island have been divided into six groups (WRB, 150 151 2014): Leptosols, Cryosols, Fluvisols, Regosols, Histosols, and Technosols; this corresponds 152 well with previously published data (Navas et al, 2008, Abakumov, 2017).

153 Three soils were selected for humic substance isolation and further investigation in this study. All soils have top humus layers with a high carbon content and distinguishable layers of 154 suprapermafrost accumulation of organic matter. All three soils are classified as Turbic Cryosols 155 (Histic, Stagnic) (WRB. 2014). Soil profiles 1, 2, and 3 (SP1, SP2, SP3) were collected from 156 locations described by the following coordinates: 62,14,391 S, 58,58,549 W; 62,13,140 S, 157 58,46,067 W; and 62,10,578 S, 58, 51,446 W respectively. Sampling depth was 0 - 10 cm for the 158 superficial layers and 50 - 55, 15 - 20, 20 - 25 for SP1, SP2, and SP3 respectively. Images of the 159 soil profiles are presented in Fig. 2. SP1 is from under the mixed lichen-bryophyta cover, SP2 160 161 and SP3 are formed under species of *Bryophyta* and *Deshampsia antarctica* correspondingly.

162 **2.2.** Sampling and laboratory analysis

Soil samples were air-dried (24 hours, 20°C), ground, and passed through 2-mm sieve. Routine chemical analyses were performed using classical methods: C and N content were determined using an element analyzer (Euro EA3028-HT Analyser) and pH in water and in salt suspensions using a pH-meter (pH-150 M).

167 Humic acids (Has) were extracted from each sample according to a published protocol (Shnitzer, 1982), http://humic-substances.org/isolation-of-ihss-samples/). Briefly, the soil 168 samples were treated with 0.1 M NaOH (soil/solution mass ratio of 1:10) under nitrogen gas. 169 170 After 24 hours of shaking, the alkaline supernatant was separated from the soil residue by 171 centrifugation at $1,516 \times g$ for 20 minutes and then acidified to pH 1 with 6 M HCl to precipitate 172 the HAs. The supernatant, which contained fulvic acids, was separated from the precipitate by centrifugation at $1,516 \times g$ for 15 minutes. The HAs were then dissolved in 0.1 M NaOH and 173 174 shaken for four hours under nitrogen gas before the suspended solids were removed by 175 centrifugation. The resulting supernatant was acidified again with 6 M HCl to pH 1 and the HAs 176 were again isolated by centrifugation and demineralized by shaking overnight in 0.1 M HCl/0.3 177 M HF (soil/solution ratio of 1:1). Next, the samples were repeatedly washed with deionized 178 water until pH 3 was reached and then finally freeze-dried. HA extraction yields were calculated 179 as the percentage of carbon recovered from the original soil sample (Vasilevitch et al. 2018, 180 Abakumov et al, 2018).

Isolated HAs were characterized for their elemental composition (C, N, H, and S) using the
 Euro EA3028-HT analyzer. Data were corrected for water and ash content. Oxygen content was
 calculated by difference. The elemental ratios reported in this paper are based on weight. Solid-

state ¹³C-NMR spectra of HAs were measured with a Bruker Avance 500 NMR spectrometer in 184 185 a 3,2-mm ZrO2 rotor. The magic angle spinning speed was 20 kHz in all cases and the nutation frequency for cross polarization was u1/2p 1/4 62.5 kHz. Repetition delay and number of scans 186 were 3 seconds. Groups of structural compounds were identified by their chemical shifts values: 187 alkyl C (-10 to 45 ppm), O/N-alkyl C (45 to 110 ppm), aryl/olefine C (110 to 160 ppm), and 188 189 carbonyl/carboxyl/amide C (160 to 220 ppm) (Kniker, 2007). The ¹³C-NMR study was also 190 conducted in bulk soil samples towards characterizing changes in the initial soil material during 191 humification.

The ESR spectra (only for HAs due to low ash content) were recorded on a JES FA 300 spectrometer (JEOL, Japan) in X-diapason with a free-radical modulation amplitude of 0.06 mT and a microwave power in the cavity of 1 mW. Magnesium powder with fixed radical concentration was used as an external standard. The concentration of the paramagnetic centers in powdered samples was determined by comparison to relative signal intensities of the external standard using the program JES-FA swESR v. 3.0.0.1 (JEOL, Japan). (Chukov et al, 2017).

198 2.3. Statistical analysis

Statistical data analysis was performed using the STATISTICA 10.0 software (TX, USA). 199 One-way analysis of variance (ANOVA) was applied to test the statistical significance of the 200 differences between the data, based on estimation of the significance of the average differences 201 202 between three or more independent groups of data combined by one feature (factor). Fisher's 203 Least Significance Test (LST) was used for post-hoc analysis to provide a detailed evaluation of the average differences between groups. A feature of this post-hoc test is inclusion of intra-group 204 mean squares when assessing any pair of averages. Differences were considered significant at 205 the 95% confidence level. Concentrations of organic and inorganic contaminants were 206 207 determined with at least three replicates. The calculated average concentrations are provided as 208 mean \pm standard deviation (SD).

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210 **3. Results and Discussion**

211 It was previously suggested that temperature and humidity are the most important factors determining the most of soil-forming processes in cold climate and humid environments 212 (Campbell and Claridge, 1982, Matsuoka et al., 1990). However, Maritime Antarctica is differed 213 from the other regions of Earth by the high influence of sea birds and seals on soil-forming 214 215 processes as they provide additional source of biogenic elements and significantly change the 216 chemistry of soils. Seabird and seal colonies significantly changes biotic activity in marine terraces of Maritime Antarctica (Gonzalez-Guzman et al, 2017). Periglacial features are 217 218 dominant on Fildes peninsula (King George Island) (Lopez-Martinez et al, 2016). Total organic 219 carbon (TOC) content was high in both the superficial and buried soil layers. This is indicative of the low degree of decomposition and transformation of the precursor material and is comparable 220 221 to the data on soils from the Yamal tundra (Ejarque, Abakumov, 2016) and the Argentinian islands (Parnikoza et al, 2016). High TOC content is typical for the Antarctic Peninsula 222 223 compared to soils of the Eastern Antarctic (Beyer et al, 1997, Mergelov et al, 2017). While both 224 were elevated, the TOC was higher in the superficial levels relative to the lower ones. Previous studies describe high variability in the TOC content from the soils of King George and Galindez 225 226 Islands, mainly depending on the diversity of the ecotopes and the sources of organic matter (Abakumov, 2010, Gonzalez-Guzman et al., 2017, Moura et al., 2012, Parnikoza et al, 2016). 227 TOC was found higher previously in ornithogenic soils of rocky platforms compared to non-228

ornithogenic soils (Moura et al., 2012). Isolated (buried) soil spots are not connected with fresh
 sources of organic matter, explaining why the TOC content in these layers is lower.

231 The carbon to nitrogen ratio was narrowest in SP1, which was affected by the scuas' 232 activity (evidenced by remnants of nests). This is in line with previous studies that documented the well-pronounced ornithogenic effects on soil's nitrogen content (Otero et al., 2013, Parnikoza 233 234 et al., 2016, Simas et al., 2007). Organic matter is one of the main soil components that 235 contributes to the development of many of the physical, chemical and biological properties and is of particular importance in Antarctic soils (Beyer et al., 2000). Fine earth of soils investigated 236 237 characterized by acid reaction, which is expected for soils of this region. Values of pHH2O varied from 4.70 to 6.35. These values coincides well with previously obtained for Maritime 238 Antarctica (Moura et al., 2012, Navas et al, 2017). 239

In terms of elemental composition, soil HAs are comparable with those previously reported for the Arctic and Antarctic soil. Current exposed organic layers contain HAs with higher carbon and nitrogen and lower oxygen content. Conversely, the HAs of isolated soil patches show increased levels of oxidation. In comparison to soils of the tundra in the Komi Republic (Vasilevitch et al, 2018), HAs found in this study were more oxidized, comparable to those of the Kolyma Lowland (Lupachev et al, 2017) and previously published data from the Fildes Peninsula (Abakumov, 2017).

247 Data on the distribution of carbon species in HAs (fig. 3) and in bulk soil (fig. 4) samples indicated that aromatic compound content is generally lower than the alkyl components. This is a 248 well-known peculiarity of the soils of the polar biome (McKnihct et al, 1994, Beyeret al, 1997). 249 250 At the same time, the degree of aromaticity of the isolated HAs is three fold higher than in the bulk organic matter. This suggests the presences of the humification process in the soils of 251 Antarctica since humification involves increasing the aromatic compound content in 252 253 macromolecules. This supports the classical humification hypothesis instead of new arguments, 254 which are critical for this approach (Lehman, Kleber, 2015). Our data shows that SOM is on a continuum and HAs are the main acting constituent of this continuum; thereby confirming that 255 this model is applicable even in Antarctica. The degree of aromaticity was higher in both isolated 256 HAs and bulk soil samples from superficial levels compared to samples from isolated patches. 257 258 Carbonyl/carboxyl/amide area (160 - 220 ppm) was more prevalent in the HAs of topsoils and less abundant in the organic matter of bulk samples (this region was presented mainly by 259 carboxylic and amid carbon in the interval between 160 - 185 ppm) (Kniker, 2007). HAs 260 extracted form SP1, located under the Deshampsia antarctica, exhibited wide peaks around 110 -261 140 ppm (H-aryl, C-aryl, olefinic-C) and at 140 - 160 ppm (O-aryl and N-aryl-C), while 262 263 aromatic components of SP2 and SP3 were mainly represented by peaks between 110 - 140 ppm. 264 This difference can be explained by the organic remnants of *Deshampsia antarctica* serving as the precursor for humification. All HA samples showed intensive areas of alkylic carbon (0 - 45 265 266 ppm), aliphatic C and N, and methoxyl C (45 - 110 ppm), O-alkyl of carbohydrates and alcohols 267 (60 - 95 ppm), and acetal and ketal carbon of carbohydrates (95 - 110 ppm). Carbon composition of the bulk samples was different from isolated HAs as evidenced mainly by the presence of 268 269 alkyl carbon (0 - 45 ppm) and O- and N-alkyl carbon (45 - 110 ppm). Characteristic features of 270 the bulk organic matter include carboxylic carbon and aryl compound content was low relative to isolated HAs. Only soils with prior ornithogenic interactions showed increases in carboxylic
peaks, which corresponds well to data on relic ornithogenic soil (Beyer et al, 1997).

The C-alkyl : O-N-alkyl ratio used to indicate the degree of organic matter transformation 273 was quite variable in all samples investigated. This can be caused by diversity in the origin and 274 275 composition of the humification precursors. In case of comparisons with humic and fulvic acids of tundra soils (Vasilevitch et al, 2018), HAs of soils investigates are intermediated between 276 277 HAs and fulvic acids of tundra Histosols with partially decomposed organic matter. These data 278 are in line with a previous report (Hopkins et al, 2006) that showed soils of the Antarctic Dry Valleys have low alkyl-C : O-alkyl-C ratio using solid-state ¹³C-NMR spectroscopy) and, 279 therefore can serve as a labile, high-quality resource for micro-organisms. Beyer et al (1997) 280 showed that both the CPMAS ¹³C-NMR and the Py-FIMS spectra of the Terri-Gelic Histosol 281 were dominated by signals from carbohydrates and alkylic compounds, which is corroborated by 282 our findings. They also suggest that the ¹³C-NMR data reflected decomposition of carbohydrates 283 and enrichment of alkyl-C in deeper soil layers. In regards to the bulk SOM, this was true for 284 SP2 and SP3 but not for SP1 that formed under the vascular plant Deshampsia Antarctica. 285

286 A representative electrone spin resonanse ESR spectrum of HAs is presented in fig 5 and the 287 ESR parameters are similar to HAs and FAs of temperate soils (Senesi, 1990, Senesi et al, 2003). 288 The spectra show a single, wide line with a g-factor ranging from 1,98890 to 1,99999, attributable to the presence of stable semiquinone free radicals in the HA-containing 289 290 macromolecules (Table 5). The free radical content was higher in the superficial levels than in 291 the isolated ones. This corresponds well with previous reports (Chukov et al, 2017, Abakumov et 292 al, 2015) that connect the isolation of buried organic matter in the supra-permafrost with declining free radical content. This reveals the increased biochemical activity of HAs in topsoil. 293 294 Compared to data from Lupachev (2017), the differences between exposed and isolated areas are 295 less pronounced but, in general, the HAs of the Antarctic soils contain more unstable free 296 radicals on average than the tundra soils of the Kolyma Lowland (Lupachev et al, 2017) and are comparable to the soils from the Yamal tundra (Chukov et al, 2017). Taken together, the free 297 298 radical content found in our study was lower than in anthropogenically affected boreal and forest 299 steppe soils of the East-European plains (Abakumov et al, 2018).

4. Conclusions

301 High TOC content was fixed for the three studies representatives of Turbic Cryosols on 302 King George Island, Northwest of the Antarctic Peninsula, Western Antarctic. High amounts of TOC are characteristic for both superficial and partially isolated soil materials. HAs contained 303 304 three fold more aromatic carbon than bulk SOM, which indicates that humification appears and 305 is active in soils of the Antarctic. Moreover, the amounts of aromatic carbon and carboxyl groups 306 were higher in the HAs of the superficial layer, which is likely caused by the greater diversity of their organic precursors and more active humification than in sub-aerial conditions. The HAs of 307 308 the superficial sample layers contained lower concentrations of free radicals, an indicator of 309 active transformation in the topsoil. In general, the organic matter from partially isolated areas is 310 less stable in terms of carbon species and free radical content. This likely results from the relative lack of fresh organic precursors and the different aeration and hydration conditions of 311 stagnification bordering the permafrost table. 312

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477 Table. 1. Basic characteristics of soils

Sample	TOC, %	N, %	C/N	pH _{H2O}	pH _{CaCL2}	Color
1.0	27.63±0.23	5.18±0.42	5.33	6.35	5.30	10 YR 4/7
1 0	19.05±0.15	2.20±0.05	8.66	5.67	4.89	2.5 YR 4/4
2 [CRH]	19.05±0.15	2.20-0.03	0.00	5.07	4.09	2.5 11 4/4
	20.04±0.17	1.16±0.09	17.13	4.80	4.80	10 YR 4/4
3 0						
	12.33±0.24	0.78 ± 0.09	15.80	4.70	4.50	2.5 YR 4/3
5 [CRH]						
	10.16±0.09	0.84 ± 0.07	11.98	4.90	4.21	10 YR 5/3
4 O						
	6.66±0.07	0.81±0.09	8.20	4.70	4.35	2.5 YR 5/3
6 [CRH]						

Sample №	С	Ν	Н	0	C/N	H/C	O/C
1	49.53±0.56	5.55±0.07	6.90±0.11	38.02±0.64	8.92	0.13	0.76
2	47.14±0.45	4.30±0.06	6.79±0.09	41.77±0.21	10.96	0.14	0.88
3	45.55±0.32	5.14±0.09	5.80±0.09	43.51±0.35	8.86	0.12	0.95
4	43.77±0.24	4.72±0.11	6.90±0.08	44.61±0.21	9.27	0.15	1.01
5	49.99±0.41	4.78±0.08	6.56±0.08	38.67±0.34	10.45	0.13	0.77
6	44.45±0.034	3.99±0.07	6.77±0.10	44.79±0.25	11.14	0.15	1.01
P, One way Anova,	0.14		0.00				
superficial/buried	0.14	0.05	0.29	0.05	n.d.	n.d.	n.d.

481 Table 2. Elemental composition (%) and atomic ratios in HAs. Data presented in atomic values.

	Carbonyl/ carboxyl/	Aryl-	O-N		Calkyl/O-N	Caryl/Calkyl
Sample №	amide	olefine	alkyl	Calkyl	alkyl	, ,
Chemical shift,						
ppm	220-160	160-110	110-45	45-0		
1	11,38	33,59	39,86	14,18	0.35	2.36
2	10,75	30,45	31,86	26,05	0.81	1.16
3	19,24	23,34	29,54	27,85	0.94	0.83
4	16,48	21,42	34,23	27,87	0.81	0.77
5	16,75	33,40	29,12	20,71	0.71	1.61
6	14.39	26.86	40.07	18.68	0.46	1.43
P, One way						
Anova,						
superficial/buried						
	0.02	0.03	0.02	0.73	n.d.	n.d.

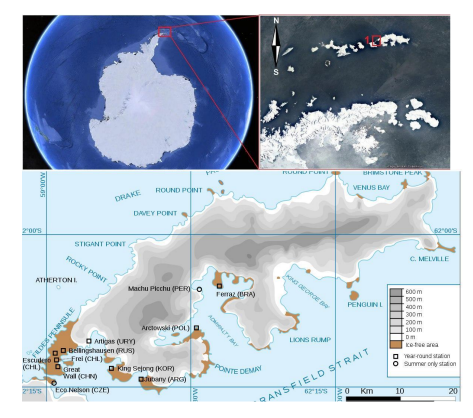
485 Table 3. Carbon species integration in molecules of the HAs, %

	Carbonyl/					Caryl/Calkyl
	carboxyl/	Aryl-	O-N		Calkyl/O	
Sample №	amide	olefine	alkyl	Calkyl	-N alkyl	
Chemical shift, ppm	220-160	160-110	110-45	45-0		
1 =113=O	7.24	11.37	46.20	35.19	0.76	0.32
2 113-Ch	18.23	10.29	40.59	30.89	0.76	0.33
3 123 O	7.34	20.48	55.12	17.06	0.31	1.20
4 123 Ch	9.34	11.27	49.50	29.90	0.60	0.37
6 149 O	5.72	13.84	62.22	18.22	0.29	0.75
6 149 Ch	22.95	9.89	46.92	20.24	0.43	0.48
						n.d.
P, One way Anova, superficial/buried	0.53	0.01	0.05	0.56	n.d.	

489 Table 4. Carbon species integration in molecules of the bulk organic matter, %

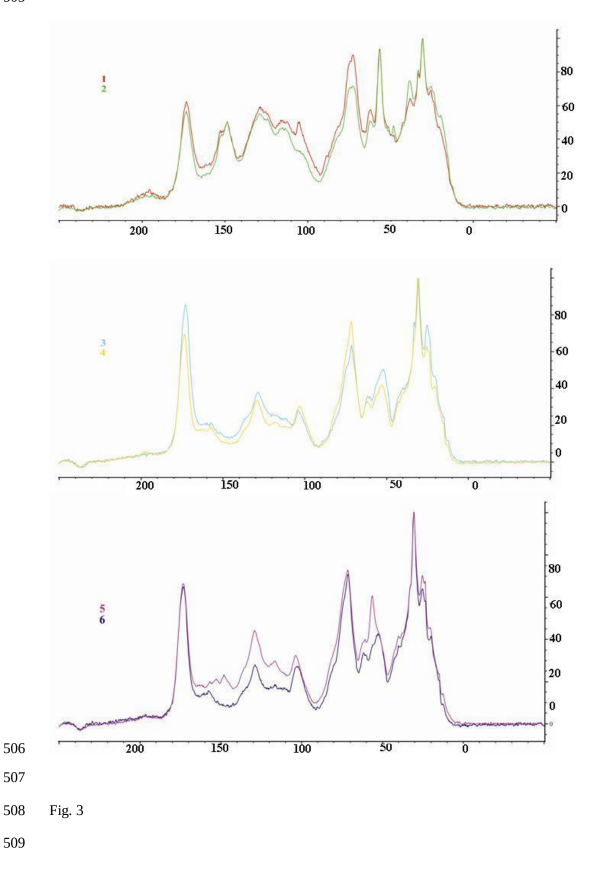
Soil horizon	Mass concentration of free radical, 10 ¹⁵ spin*g ⁻¹	g-factor
1	3.67	2.0314
2	3.04	2.3150
3	3.51	2.0314
4	2.13	2.0303
5	6.10	2.0310
6	5.86	2.0314

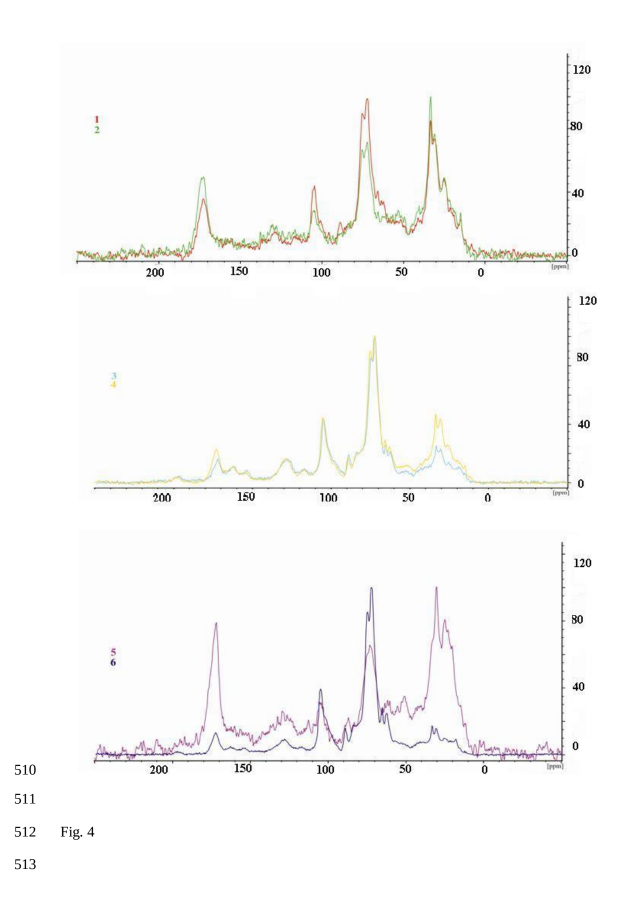
493 Table. 5. Mass concentration of free radical in humic acids











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515	Fig. 5.
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518	Fig. 1. Location of the Fildes peninsula
519	Fig.2. Soil morphology
520	Figure 3. 13-C NMR spectras of the HAs, isolated from soils (1-6 – according table 1)
521	Fugure 4. 13-C NMR spectras of bulk organic matter of soils ((1-6 – according table 1)
522	Figure 5. Typical ESR spectrum of humic substances investigated
523	Table. 1. Basic characteristics of soils
524	Table 2. Elemental composition (%) and atomic ratios in HAs
525	Table 3. Carbon species integration in molecules of the HAs, %
526	Table 4. Carbon species integration in molecules of the bulk organic matter, %
527	Table. 5. Mass concentration of free radical in humic acids
528	