| 1  | Contrasting potential for biological N <sub>2</sub> -fixation at three   |
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| 2  | polluted Central European Sphagnum peat bogs: Combining  |
| 3  | the <sup>15</sup> N <sub>2</sub> -tracer and natural-abundance isotope approaches  |
| 4  |  |
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| 17 | ABSTRACT   |
| 18 |  |
| 19 | Availability of reactive nitrogen $(N_r)$ is a key control of carbon $(C)$ sequestration in wetlands. To complement  |
| 20 | the metabolic demands of Sphagnum in pristine rain-fed bogs, diazotrophs supply additional Nr via biological   |
| 21 | nitrogen fixation (BNF). Since breaking the triple bond of atmospheric $N_2$ is energy-intensive, it is reasonable to                                      |
| 22 | assume that increasing inputs of pollutant $N_r$ will lead to BNF downregulation. Yet, recent studies have   |
| 23 | documented measurable BNF rates in Sphagnum-dominated bogs also in polluted regions, indicating adaptation   |
| 24 | of N2-fixers to changing N deposition. Our aim was to quantify BNF at high-elevation peatlands located in  |
| 25 | industrialized Central Europe. A $^{15}\mathrm{N}_2\text{-tracer}$ experiment was combined with a natural-abundance N-isotope study                        |
| 26 | at three Sphagnum-dominated peat bogs in the northern Czech in an attempt to assess the roles of individual  |
| 27 | BNF drivers. High short-term BNF rates $(8.2 \pm 4.6 \text{ g N m}^2 \text{ d}^{-1})$ were observed at Male Mechove Jezirko                                |
| 28 | receiving ~17 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> . The remaining two peat bogs, whose recent atmospheric $N_r$ inputs differed from                |
| 29 | Male Mechove Jezirko only by 1-2 kg ha <sup>-1</sup> yr <sup>-1</sup> (Uhlirska and Brumiste), showed zero BNF. The following                              |
| 30 | parameters were investigated to elucidate the BNF difference: $NH_4^+$ - $N/NO_3^-$ - $N$ ratio, temperature, wetness,                                     |
| 31 | Sphagnum species, organic-N availability, possible P limitation, possible Mo limitation, SO42- deposition, and   |
| 32 | pH. At Male Mechove Jezirko and Uhlirska, the same moss species (S. girgensohnii) was used for the ${}^{15}N_2$  |
| 33 | experiment, and therefore host identity could not explain the difference in BNF at these sites. Temperature and  |
| 34 | moisture were also identical in all incubations and could not explain the between-site differences in BNF. The   |
| 35 | N:P stoichiometry in peat and bog water indicated that Brumiste may have lacked BNF due to P limitation,   |
| 36 | whereas non-detectable BNF at Uhlirska may have been related to 70 times higher $SO_4^{2-}$ concentration in bog   |
| 37 | water. Across the sites, the mean natural-abundance $\delta^{15}N$ values increased in the order: atmospheric deposition (-                                |

38  $5.3 \pm 0.3 \%$ ) < Sphagnum (-4.3 ± 0.1 ‰) < bog water (-3.9 ± 0.4 ‰) < atmospheric N<sub>2</sub> (0.0 ‰). Only at 39 Brumiste, N in *Sphagnum* was significantly isotopically heavier than in atmospheric deposition, possibly 40 indicating a longer-term BNF effect. Collectively, our data highlight spatial heterogeneity in BNF rates under 41 high  $N_r$  inputs and the importance of environmental parameters other than atmospheric  $N_r$  pollution in regulating 42 BNF. 43 44 45 Keywords: Peat, Sphagnum, nitrogen deposition, pollution, biological nitrogen fixation, BNF controls, 46 phosphorus limitation 47 48 49 **1. Introduction** 50 51 Nitrogen (N) is the limiting nutrient in most terrestrial environments. The amount and form of N available to 52 organisms (reactive N, N<sub>r</sub>) is controlled by biogeochemical processes (Vitousek and Howarth, 1991; LeBauer 53 and Treseder, 2008; Zhang et al., 2020; Davies-Barnard and Friedlingstein, 2020). A growing body of research 54 has focused on the role of biological  $N_2$ -fixation (BNF) as a source of  $N_r$  in pristine ecosystems, such as 55 subarctic tundra and boreal forests, with special attention given to ombrotrophic peat bogs and minerotrophic 56 fens (Hemond, 1983, Rousk et al., 2013, 2015; Larmola et al., 2014; Vile et al., 2014; Diakova et al., 2016; 57 Stuart et al., 2021; Yin et al., 2022). Globally, peatlands store between 20 and 30 % of total soil carbon and 58 approximately 15 % of total soil nitrogen (Wieder and Vitt, 2006; Gallego-Sala et al., 2018; Fritz et al., 2014). 59 Microbial N<sub>2</sub>-fixation helps to sustain C accumulation in peatlands and to remove carbon dioxide (CO<sub>2</sub>) from the 60 atmosphere (Vile et al., 2014, and references therein). Changes in BNF may affect the dynamics of climate 61 change. A combination of high anthropogenic Nr inputs with sustained N<sub>2</sub>- fixation may accelerate invasion of 62 vascular plants into peat bogs leading to the reduction of the C-N stocks. 63 64 The nitrogen budget at the peat bog scale results from a balance between N inputs [atmospheric deposition of  $N_r$ , 65 mostly nitrate ( $NO_3^{-}$ ) and ammonium ( $NH_4^{+}$ ), with a contribution of organic N and BNF] and N outputs [runoff 66 dominated by dissolved, colloidal, and particulate N, and emissions of gaseous N forms, mainly nitrous oxide 67  $(N_2O)$ , nitric oxide (NO), and  $N_2$  as products of denitrification; Souridis et al., 2021]. The atmospheric lifetime 68 of  $N_2O$ , a potent greenhouse gas, is relatively long (>100 yr; Frolking et al., 2011). In contrast, the atmospheric 69 lifetime of NO, another greenhouse gas, is short (days), and, along with N<sub>2</sub> as the final product of denitrification 70 with no warming potential, is not considered in climate warming scenarios. Atmospheric deposition of  $N_r$  in 71 high-latitude pristine bogs is 0.5-1.0 kg ha<sup>-1</sup>yr<sup>-1</sup> (Vitt et al., 2003). Bogs receiving less than 10 kg N<sub>r</sub> ha<sup>-1</sup>yr<sup>-1</sup> are 72 defined as low-polluted (Lamers et al., 2000). Bogs receiving more than 18 kg Nr ha<sup>-1</sup>yr<sup>-1</sup>are considered to be 73 highly polluted. Reactive N deposited on the surface of ombrotrophic peat bogs is vertically mobile (Novak et 74 al., 2014). 75

76 Nitrogen capture in rain-fed bogs is dominated by *Sphagnum* mosses (Limpens et al., 2006). Nitrogen-fixing

77 microbes (diazotrophs) mostly reside inside specialized *Sphagnum* cells (hyalocytes), although the mosses'

- 78 metabolic demands for N are supported also by free-living diazotrophs. In contrast, diazotrophs in feather
- 79 mosses, common in boreal forests, live epiphytically on the leaves (DeLuca et al., 2002; Rousk et al., 2015).
- 80 Endophytic diazotrophs are more protected against environmental fluctuations, including changes in Nr
- 81 deposition. BNF in bogs is associated mostly with cyanobacteria and methanotrophs (Larmola et al., 2014; Vile
- 82 et al., 2014; Leppanen et al., 2015; Holland-Moritz et al., 2021; Kolton et al., 2022). It follows that BNF may
- 83 affect potential methane (CH<sub>4</sub>) emissions in two opposing directions: while higher C accumulation due to
- 84 efficient BNF may lead to higher CH<sub>4</sub> emissions during peat decomposition, N<sub>2</sub>-fixing methanotrophs may
- 85 reduce emissions of  $CH_4$  by oxidizing this greenhouse gas.
- 86
- 87 Recent work in peatlands has quantified the relative roles of various biotic and abiotic controls over BNF.
- 88 Leppanen et al. (2015) reported than BNF rates were independent of the diazotroph community structure. The
- 89 effect of temperature was reviewed by Carrell et al. (2019), Zivkovic et al., (2022), and Yin et al. (2022). The
- 90 optimal temperature for BNF is 20-30 °C (Zielke et al., 2005). Dry conditions are generally unfavorable for
- 91 BNF, but the moisture–BNF correlation tends to be insignificant (Yin et al., 2022). The effect of phosphorus (P)
- 92 as a limiting nutrient was evaluated by Limpens et al. (2004), Larmola et al. (2014), Ho and Bodelier (2015), van
- den Elzen et al. (2017, 2020), and Zivkovic et al. (2022). In an interplay with other environmental and chemical
- 94 parameters, higher P availability may augment BNF. The role of the  $NH_4^+/NO_3^-$  ratio in atmospheric deposition
- 95 as a BNF control was evaluated by Saiz et al. (2021). A higher NH<sub>4</sub><sup>+</sup> proportion relative to the total N<sub>r</sub> deposition
- 96 may result in lower BNF rates. Stuart et al. (2021) stressed a strong interaction between moss identity,
- 97 temperature, moisture and pH as possible BNF drivers. Kox et al. (2018) reported higher BNF rates under
- 98 oxygen (O<sub>2</sub>) depletion. Wieder et al. (2019, 2020) and Kox et al. (2020) showed that BNF rates generally
- 99 increase in the presence of light.
- 100
- 101 In previous studies, BNF rates were measured under field conditions (*e.g.*, Vile et al., 2014; Rousk et al., 2018;
- 102 van den Elzen et al., 2020; Saiz et al., 2021; Zivkovic et al., 2022), or under controlled laboratory conditions
- 103 (*e.g.*, Knorr et al., 2015; van den Elzen et al., 2017; Warren et al., 2017; Stuart et al., 2021). According to
- 104 Myrold et al. (1999), an advantage of laboratory  ${}^{15}N_2$  experiments is related to easier preservation of a gas-tight
- 105 assay system. The rates of BNF are measured using an acetylene reduction assay (ARA),  ${}^{15}N_2$  isotope-labelling
- 106 incubations, or compound-specific amino acid <sup>15</sup>N probing (*e.g.*, Knorr et al., 2015; Chiewattanakul et al., 2022).
- 107 Recent studies have stressed the need for caution in ARA studies (Vile et al., 2014; Saiz et al., 2019; Soper et al.,
- 108 2021). Inhibition of the activity of methanotrophs by acetylene may lead to an underestimation of BNF rates.
- 109 These methods of direct measurements inevitably choose specific experimental conditions and thus provide
- 110 *potential instantaneous* BNF rates. A complementary, indirect evaluation of BNF can be based on natural-
- abundance  ${}^{15}N/{}^{14}N$  isotope systematics (Novak et al., 2016; Zivkovic et al., 2017; Saiz et al., 2021; Stuart et al.,
- 112 2021). *Sphagnum* taking up N through BNF would carry a  $\delta^{15}$ N signature close to 0 ‰, a value characterizing
- atmospheric N<sub>2</sub> ( $\delta^{15}$ N values are defined as a per mil deviation of the  ${}^{15}$ N/ ${}^{14}$ N ratio in the sample from a standard;
- 114 the widely used standard is atmospheric N<sub>2</sub>). With increasing BNF rates, the  $\delta^{15}$ N values of living *Sphagnum*
- 115 converge from the often negative  $\delta^{15}N$  value of atmospheric deposition to the 0 % value of the source N<sub>2</sub>. This
- simple approach is complicated by tight inner N cycling near the bog surface, involving open-system isotope
- fractionations. In particular, *Sphagnum* may additionally take up Nr resulting from mineralization of organic N.

118Because denitrification preferentially removes isotopically light N in a gaseous form, the residual  $N_r$  in bog119water may become isotopically heavy and supply high- $\delta^{15}N$  nitrogen for assimilation. Mineralized  $N_r$  in bog120water as another nutrient source may thus be isotopically similar to atmospheric  $N_2$  (Novak et al., 2019; Stuart et121al., 2021).

122

123 BNF is an energy-intensive process requiring 16 adenosine-triphosphate (ATP) molecules to fix 1 mol of N<sub>2</sub>. It 124 follows that, with an increasing input of pollutant  $N_r$  via atmospheric deposition, BNF should be rapidly 125 downregulated. However, experiments applying additional  $N_r$  to Sphagnum both in the laboratory and in the 126 field have indicated contradictory impacts on BNF. Some studies have shown a decrease in BNF rates in the 127 proximity of anthropogenic Nr sources (Wieder et al., 2019; Saiz et al., 2021), while others have indicated 128 continuing BNF even at N-polluted sites (van den Elzen et al., 2018). BNF data from natural settings with 129 known time-series of historical Nr deposition rates are rare (van den Elzen et al. 2018; Saiz et al., 2021). The aim 130 of the current study was to quantify BNF at high-elevation Sphagnum-dominated peatlands in an industrial part 131 of Central Europe, also known for intense agriculture. We combined <sup>15</sup>N<sub>2</sub>-tracer experiments with a natural 132 abundance N-isotope study at three peat bogs situated in the northern Czech Republic to provide qualitative 133 insights into the roles of individual BNF drivers. Our specific objectives were: (i) to investigate whether BNF 134 rates at the study sites correlate with well-constrained  $NO_3^-$  and  $NH_4^+$  deposition rates and P availability, and (ii) 135 to compare the results of experiments investigating <sup>15</sup>N-assimilation by Sphagnum with the results of a natural-136 abundance  $\delta^{15}$ N inventory of individual wetland pools and fluxes. We expected that convergence of *Sphagnum* N toward  $\delta^{15}N_{N2} = 0$  ‰ would corroborate the relative magnitude of instantaneous BNF rates in between-site 137 138 comparisons. Because, thus far, the natural-abundance <sup>15</sup>N approach has been rarely adopted in BNF studies, 139 compared to the more frequently used  ${}^{15}N_2$ -labelling approach, we generated a larger  $\delta^{15}N$  data set in the natural-140 abundance <sup>15</sup>N monitoring part of our study. 141

# 142 2. Materials and methods

143

145

146 The three studied Sphagnum-dominated peat bogs (Fig. 1, Tab. 1) are located in the northern Czech Republic, a 147 highly industrialized part of Central Europe with numerous coal-burning power plants. In the 1970s-1990s, 148 Norway spruce monocultures were affected by acid rain in the vicinity of Brumiste (BRU; Krusne Mts.) and 149 Uhlirska (UHL; Jizerske Mts.). At UHL, most spruce stands died back and were harvested. The third site, Male 150 Mechove Jezirko (MMJ; Jeseniky Mts.) is surrounded by relatively healthy mature spruce forests. The distance 151 between adjacent study sites is 160-190 km (Fig. 1). The studied high-elevation catchments are drained by small 152 streams. The studied peatlands are partly rain-fed, with a possible contribution of lateral water influx from the 153 surrounding segments of the catchments. The bedrock is composed of granite at BRU and UHL, and phyllite at 154 MMJ. The surface of each bog is characterized by a combination of hummock-hollows microtopography and 155 lawns (Dohnal, 1965). Moss species at BRU include S. cuspidatum, common in hollows and pools, S. 156 magellanicum, mostly occupying intermediate positions between the tops of the hummocks and the hollows, S. 157 rubellum, typical of dense carpets in rain-fed bogs, and S. papillosum, forming low hummocks and mats in bogs

**<sup>144</sup>** *2.1. Study sites* 

- and mires. At UHL and MMJ, the predominant moss species is shade-demanding *S. girgensohnii*, requiring
- 159 slight base enrichment (Tab. S1 in the Supplement). The growing season is more than seven months long, from
- 160 late March to early November. The measured density of living *Sphagnum* is 0.04 g cm<sup>-3</sup>. More details on BRU
- 161 are in Bohdalkova et al. (2013), and Buzek et al. (2019, 2020). Biogeochemical processes at UHL were studied
- 162 by Novak et al. (2005), Sanda and Cislerova (2009), Bohdalkova et al. (2014), Marx et al. (2017), Oulehle et al.
- 163 (2017, 2021a), and Vitvar et al. (2022). Further information on MMJ is in Novak et al. (2003, 2009).



- 165 Fig. 1. Location of the studied *Sphagnum*-dominated peat bogs. Nr deposition contours are by Czech
- 166 Hydrometeorological Institute (1998).
- 167
- **168 Table 1.** Study site characteristics.
- 169

| Site                               | Location             | Elevation<br>(m) | Long-term<br>precipitation<br>total<br>(mm yr <sup>-1</sup> ) | Mean annual<br>temperature<br>(°C) | Bog area<br>(ha) | Maximum<br>peat depth<br>(cm) | Atmospheric<br>vertical N <sub>r</sub><br>deposition<br>(kg ha <sup>-1</sup> yr <sup>-1</sup> ) <sup>1</sup> | Total<br>atmospheric N <sub>r</sub><br>deposition<br>(kg ha <sup>-1</sup> yr <sup>-1</sup> ) <sup>2</sup> | NH4 <sup>+</sup> -N/NO3 <sup>-</sup> -N<br>ratio |
|------------------------------------|----------------------|------------------|---|------------------------------------|------------------|-------------------------------|--|---|--|
| Brumiste,<br>BRU                   | 50°24′ N<br>12°36′ E | 930              | 1080  | 4.5                                | 17               | 200                           | 12.7   | 16.5  | 1.2  |
| Uhlirska,<br>UHL                   | 50°49′ N<br>15°08′ E | 830              | 1230  | 4.0                                | 50               | < 200                         | 15.5   | 20.2  | 1.2  |
| Male<br>Mechove<br>jezirko,<br>MMJ | 50°13′ N<br>17°18′ E | 750              | 1090  | 5.3                                | 195              | 660                           | 14.3   | 18.6  | 1.3  |

long-term average according to Oulehle et al., 2016 <sup>2</sup>including 30 % of horizontally deposited  $N_r$  (Novak et al., 2015)

- 170
- 171

#### **172** *2.2. Sampling*

173

174In our study, we compared long-term N isotope data (natural-abundance  ${}^{15}N$  monitoring in the field) with short-175term N isotope data ( ${}^{15}N_2$  laboratory moss incubations). Out of the 403  $\delta^{15}N$  measurements performed, 361 were

related to the field monitoring, and 42 were related to laboratory incubations.

177

178 Samples of rain and snow for  $\delta^{15}$ N determinations were collected between January 2016 and October 2019 using 179 a simplified protocol of Fottova and Skorepova (1998). Open-area precipitation was sampled by two rain 180 collectors placed five meters apart, 160 cm above ground. Spruce canopy throughfall was sampled using five 181 (UHL) or three (BRU, MMJ) collectors installed 10 m apart. Deposition samplers were polyethylene (PE) 182 funnels (surface area of 113 cm<sup>2</sup>) fitted to 1-L bottles. In winter, cylindrical PE vessels (surface area of 167 cm<sup>2</sup>) 183 were used to collect snow. At the end of cumulative one-month sampling, open area precipitation and throughfall 184 samples, respectively, were pooled prior to chemical and N-isotope analysis. One-liter samples of runoff were 185 collected in ~30-day intervals at BRU over a 25-month period, samples of runoff were collected at UHL and 186 MMJ in summer 2019 (see Tab. S2 for specific dates). Five replicates of surface bog water were collected 187 throughout each study site in June 2019. The depth of the water pools was less than 20 cm. The total number of water samples for  $\delta^{15}$ N analysis was 136. 188 189 190 A vertical peat core, 10-cm in diameter, 30-cm deep, was collected in a Sphagnum-dominated lawn at each of 191 the study sites in October 2018, kept vertically at 6 °C for 12 hours and then frozen. At the same time, 12 192 samples of living Sphagnum were collected randomly throughout each bog for species identification and N 193 isotope analysis. Additionally, 14 replicate samples of living Sphagnum were collected in various parts of each 194 of the peat bogs for a <sup>15</sup>N<sub>2</sub>-labelling experiment. Each replicate sample consisted of 30 individual 5-cm long

- 195 Sphagnum plants. S. girgensohnii was used in the UHL and MMJ experiments, a mix of S. magellanicum, S.
- 196 papillosum, and S. cuspidatum was used in the BRU experiment (cf., Tab. S1); Sphagnum samples were
- transported to the laboratory at a temperature of 6 °C. Transportation took 2 to 4 hours, the wet samples were

then kept at 6 °C, laboratory experiments started 2 to 6 days after moss collection. Prior to incubation, the moss
samples were kept at 22 °C for 4 hours.

200

In section 3.3.3, we will compare N isotope composition of living *Sphagnum* and surface bog water. These two
 sample types were collected *ca*. 8 months apart. Previous research based of <sup>210</sup>Pb peat-core dating has indicated
 that 5-cm long *Sphagnum* capitula and stems at 14 rainfed Central European sites represent more than a three year growth increment (Novak et al., 2003, 2008). Hence, N-isotope comparisons of living *Sphagnum* and bog
 water sampled less than 1 year apart may still provide useful information.

206

# 207 2.3. <sup>15</sup>N<sub>2</sub> <u>Sphagnum</u> incubation experiment

208 209 Measurements of potential N<sub>2</sub>-fixation rates were performed using a modified protocol of Larmola et al. (2014). 210 Four plant replicates per site were analyzed at time t = 0 without incubation (control no. 1). Ten replicates per 211 site were closed in 200-mL transparent PE containers with 5 mL of bog water collected at BRU, UHL and MMJ, 212 respectively. Out of these, four Sphagnum replicates with no  $^{15}N_2$  addition served as a procedural control no. 2 to 213 identify possible incubation artifacts after 168 hours. In the remaining six closed containers with Sphagnum 214 replicates, 24 mL of headspace air were removed at t = 0 and replaced with 32 mL of  $^{15}N_2$  tracer gas containing 215 98 atomic % of <sup>15</sup>N (Aldrich, Germany). Two <sup>15</sup>N-labelled replicates were incubated for 48 hours, another four 216 <sup>15</sup>N-labelled replicates were incubated for 168 hours. According to Zivkovic et al. (2022), BNF rates peak in 217 summer at relatively high ambient temperatures. We used data by Czech Hydrometeorological Institute to set the 218 incubation temperatures. The nearest high-elevation weather stations were Serak (15 km distance from MMJ, 219 1328 m a.s.l.) with day-time mean temperature for the June 21-September 23, 2017 period of 16.7 °C, and night-220 time temperatures of 9.4 °C, and Karlova Studanka (18 km distance from MMJ, 795 m a.s.l.) with analogous 221 temperature averages of 16.2 and 11.5 °C. Each day, the temperature in the growth chamber in our experiment 222 was kept at 18 °C for 16 hours at daylight, and at 10 °C for 8 hours under dark conditions. The duration of 223 daylight and darkness was unified with experimental conditions applied by van den Elzen et al. (2017). 224

Following N-isotope analysis, BNF rates were calculated according to Vile et al. (2014) and Knorr et al. (2015):

227 
$$N_{2fix} = \frac{\Delta at. \ \%^{15} N_{Sph}}{\Delta at. \ \%^{15} N_{gas}} \ x \ \frac{\text{total N } \%_{Sph}}{t^* 100} \quad (\text{g N g DW}^{-1} \text{ day}^{-1}), \tag{1}$$

228

where N<sub>2fix</sub> is the N<sub>2</sub>-fixation rate in g N g DW<sup>-1</sup>(*Sph*) day<sup>-1</sup>, t is incubation time (days), total N% <sub>*Sph*</sub>,  $\Delta$  at. % 1<sup>5</sup>N<sub>*Sphagnum*</sub> is the difference between atom % labeled and control sample,  $\Delta$  at. % <sup>15</sup>N<sub>*gas*</sub> is the difference between the concentration <sup>15</sup>N in the headspace and the natural abundance (at. %). The used *Sphagnum* density was 0.04 g cm<sup>-3</sup>.

233

234 We used larger sealed containers, compared to previous  ${}^{15}N_2$  experiments ( $\leq 96$  hours; Myrold et al., 1999) to

 $\label{eq:235} \mbox{minimize the effect of changing headspace concentrations of $O_2$ and $CO_2$ on the living moss and the microbiome}$ 

after 168 hours. While, for example, van den Elzen et al. (2017) used 30 mL containers, Saiz et al. (2021) used

50 mL containers, and Stuart et al. (2021) worked with a container volume of 60 mL, we used a sealed 200 mL
volume.

240 It bears mention that Dabundo et al. (2014) found a deviation from the declared  ${}^{15}N_2$  purity within commercially 241 available tracer tanks. We did not study the tracer purity and hence the observed BNF rates might be viewed as 242 maximum estimates. Because our incubation study was based on one-time measurements under laboratory 243 conditions, in the current paper we chose not to upscale the BNF rates to the entire peat bog and an annual time 244 span.

245

246 2.4. Chemical and isotope analysis

247

248 Frozen peat cores were sectioned to 2-cm thick segments. Samples of peat and Sphagnum were dried and 249 homogenized. Nitrogen concentrations in peat and Sphagnum samples were determined on a Fisons 1180 250 elemental analyzer with a 1.5 % reproducibility  $(2\sigma)$ . Ammonium and nitrate concentrations in water samples 251 were determined spectrophotometrically with a reproducibility of 0.1 mg L<sup>-1</sup>. About 0.5 L of each water sample 252 were used to separate NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> (Bremner, 1965). Nitrogen isotope composition was measured on a Delta V mass spectrometer and expressed in  $\delta^{15}N$  notation. IAEA isotope standards N1 ( $\delta^{15}N = 0.4$  ‰) and N<sub>2</sub> ( $\delta^{15}N =$ 253 20.3 ‰) were analyzed before every session, and two in-house standards (ammonium sulfate,  $\delta^{15}N = -1.7$  ‰, 254 255 and glycine,  $\delta^{15}N = 4.0$  ‰) were analyzed after every six samples. The reproducibility of the  $\delta^{15}N$ 256 determinations was 0.30 and 0.35 ‰, for the liquid and solid samples, respectively. Methods of concentration 257 analysis of other chemical species in October 2018 samples are given in Appendix I. 258 259 2.5. Historical rates of  $N_r$  deposition

260

Long-term data from 32 monitoring stations in the Czech Republic operated by the Czech Hydrometeorological
Institute, Prague, were used to assess temporal and spatial variability of NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> concentrations in vertical
deposition using a model by Oulehle et al. (2016). Median *z*-score values of NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> concentrations

derived from observations at the monitoring stations and nation-wide emission rates, published by Kopacek and

265Vesely (2005), and Kopacek and Posh (2011), showed significant relationships at the p < 0.001 level. Using266linear models, *z*-score values were expressed for the period 1900-2012 and then back-transformed to give

- 267 concentration estimates for the study sites. Annual rates of vertically deposited  $NH_4^+$  and  $NO_3^-$  were products of
- 268 modelled concentrations and precipitation quantities at BRU, ULH and MMJ.
- 269
- **270** *2.6. Statistical evaluation*

271

272 Statistical analysis was performed using the R software (R Core Team, 2019) version 3.6.2, and its contributed

273 packages *sandwich* (Zeileis, 2004) and *multcomp* (Hothorn et al., 2008). Comparisons of groups of N isotope

and concentration data (see sections 2.3 and 2.4)

275 were based on one-way analysis of variance with a sandwich estimator of covariance matrix to account for

276 heteroscedasticity among the groups (MacKinnon and White, 1985). *Post-hoc* multiple comparisons of the same

| 277 | groups were then performed according to Hothorn et al. (2008). Because of the largely uneven number of runoff   |
|-----|---|
| 278 | samples <i>per</i> site (50, 6, and 2 at BRU, UHL and MMJ, respectively), we did not include runoff $\delta^{15}$ N data in the   |
| 279 | statistical evaluation.   |
| 280 |   |
| 281 | 3. Results  |
| 282 |   |
| 283 | 3.1. Historical rates of atmospheric Nr inputs  |
| 284 |   |
| 285 | Vertical deposition of $NH_4^+$ reached a maximum in 1980, remained almost unchanged until 1990, and decreased  |
| 286 | thereafter (Fig. 2a). Nitrate-N deposition exhibited a wider maximum between ca. 1970 and 1990 (Fig. 2b). In  |
| 287 | the oldest modelled time period (1900-1930), ammonium in deposition dominated over nitrate. During the  |
| 288 | deposition peak, the contributions of $NH_4^+$ -N and $NO_3^-$ -N to total vertical N <sub>r</sub> deposition were similar (8 to 13 kg  |
| 289 | ha <sup>-1</sup> yr <sup>-1</sup> at individual sites). Across the modelled years, the NH <sub>4</sub> <sup>+</sup> -N/NO <sub>3</sub> <sup>-</sup> -N ratio in vertical deposition was |
| 290 | similar at all three sites (1.2 to 1.3; Tab. 1). Since ca. 1950, pollution at the study sites via total vertical  |
| 291 | deposition of inorganic $N_r$ increased in the order: BRU < MMJ < UHL (Fig. 2c). Fig. 2c shows that the between-  |
| 292 | site differences in the most recent years have been small (1-2 kg N ha <sup>-1</sup> yr <sup>-1</sup> ).  |
| 293 |   |



**Fig. 2.** Modelled long-term changes in atmospheric N<sub>r</sub> deposition according to Oulehle et al. (2016).

296

**297** 3.2.  ${}^{15}N_2$  incubation experiment

298

**299** There was no change in  $\delta^{15}$ N values of *Sphagnum* after 48 hours at BRU and UHL. (Fig. 3 and Tab.2). At MMJ,

**300** the average  $\delta^{15}$ N after two days increased to 3.0 ‰. (Fig. 3 and Tab. 2). There were no statistically significant

**301** differences between  $\delta^{15}$ N values of *Sphagnum* at time t = 0 and at time t = 168 hours following incubation in

atural atmosphere (controls no. 1 and 2; Tab. 2; p > 0.05). Mean  $\delta^{15}$ N values of the moss of the two controls

303 were similar among the sites (-3.6 to -4.1 %). At the end of the 168-hour <sup>15</sup>N<sub>2</sub> *Sphagnum* incubation, there was

still no change in the N isotope signature of the moss at BRU and UHL (p > 0.05). In contrast, there was a large

305 positive shift in  $\delta^{15}$ N values of *Sphagnum* collected at MMJ (59.2 to 467 ‰; Tab. 2; Fig. 3). The N<sub>2</sub> fixation rate

306 calculated from the N isotope systematics in the  ${}^{15}N_2$  labelling experiment was 0 at BRU and UHL, and 4.11 µg 307 N g<sup>-1</sup> d<sup>-1</sup>, or 8.20 mg N m<sup>-2</sup> d<sup>-1</sup> at MMJ.

308



**Fig. 3**. Results of a <sup>15</sup>N<sub>2</sub> incubation study using living *Sphagnum*. Means and standard errors are given.



309

**Table 2**. Positive  $\delta^{15}$ N shift in total moss nitrogen following the  ${}^{15}N_2$  assay incubation at MMJ.

313

| Site        | BRU                       |  |  | UHL  |                                       |  |  | ММЈ  |                            |  |  |  |
|-------------|---------------------------|--|--|--|---------------------------------------|--|--|--|----------------------------|--|--|--|
|             |                           |  |  |  |                                       | δ <sup>15</sup> Ν  | i (‰)  |  |                            |  |  |  |
|             | Sphagnum<br>control<br>to | Sphagnum<br>at $t = 48$ h<br>of $^{15}N_2$<br>incubation | <i>Sphagnum</i><br>control<br><i>t</i> = 168 h | Sphagnum<br>at the end<br>of <sup>15</sup> N <sub>2</sub><br>incubation<br>t = 168 h | Sphagnum<br>control<br>t <sub>0</sub> | <i>Sphagnum</i><br>at <i>t</i> = 48 h<br>of <sup>15</sup> N <sub>2</sub><br>incubation | <i>Sphagnum</i><br>control<br><i>t</i> = 168 h | Sphagnum<br>at the end<br>of <sup>15</sup> N <sub>2</sub><br>incubation<br>t = 168 h | Sphagnum<br>control<br>t o | <i>Sphagnum</i><br>at <i>t</i> = 48 h<br>of <sup>15</sup> N <sub>2</sub><br>incubation | <i>Sphagnum</i><br>control<br><i>t</i> = 168 h | Sphagnum<br>at the end<br>of <sup>15</sup> N <sub>2</sub><br>incubation<br>t = 168 h |
| Replicate 1 | -3.8                      | -3.8   |  |  | -4.1                                  | -3.9   |  |  | -3.6                       | 2.7  |  |  |
| Replicate 3 | -3.9                      | -5.4   | -4.0   | -41  | -3.7                                  | -5.5   | -3.8   | -39  | -2.7                       | 5.5  | -27  | 467  |
| Replicate 4 | -3.9                      |  | -4.1   | -3.9   | -3.9                                  |  | -3.7   | -3.7   | -4.0                       |  | -3.8   | 59.2   |
| Replicate 5 | -3.9                      |  | -4.2   | -4.3   | -4.4                                  |  | -4.0   | -4.2   | -3.8                       |  | -4.0   | 68.8   |
| Replicate 6 | -3.5                      |  | -3.8   | -3.6   | -4.7                                  |  | -4.6   | -4.6   | -3.8                       |  | -4.2   | 83.0   |
| Mean ± SE   | -3.8 ± 0.1                | -3.6   | $-4.0 \pm 0.1$                                 | $-4.0 \pm 0.2$   | <b>-</b> 4.1 ± 0.2                    | -3,7   | $-4.0 \pm 0.2$                                 | <b>-4</b> ,1 ± 0,2   | <b>-3</b> .6 ± 0.3         | 3,0  | <b>-3</b> .7 ± 0.4                             | 169 ± 99,2   |

- 314
- 315

317

# 318 3.3.1. Atmospheric deposition

319

320 Ninety-six per cent of the deposited inorganic N<sub>r</sub> species had negative  $\delta^{15}$ N values; *i.e.*, contained isotopically 321 light N (Tab. S2; Fig. S1). The mean  $\delta^{15}$ N value across all 181 samples of atmospheric deposition was -5.3 ± 0.3 322 % (SE). Mean  $\delta^{15}$ N values of both forms of atmospherically deposited N (NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup>) in an open area were 323 slightly higher than those in throughfall at BRU and MMJ, and slightly lower than those in throughfall at UHL 324 (Tab. 3). Nitrate-N in open-area deposition was on average slightly isotopically lighter than NH<sub>4</sub><sup>+</sup>-N at all three 325 sites. At the 0.05 probability level, however, the within-site differences among deposition sample types and 326 among N species at BRU and MMJ were insignificant. The only statistically significant difference was found 327 between  $\delta^{15}$ N values of open-area NO<sub>3</sub><sup>-</sup> and both N species in throughfall at UHL (*see* superscript letters in Tab. 328 3).

**<sup>316</sup>** *3.3. Natural-abundance N-isotope systematics* 

| 330 | <b>Table 3</b> . Multiple comparisons among $\delta^{15}$ N values of four sample types of atmospheric deposition. Different |
|-----|--|
| 331 | letters in superscript denote statistical difference ( $p < 0.05$ ).   |

|  |   | mean δ <sup>15</sup> N (‰) ± SD  | )  |
|--|---|--|--|
| Site   | BRU   | UHL  | MMJ  |
| open-area NH4 <sup>+</sup><br>open-area NO3 <sup>-</sup><br>throughfall NH4 <sup>+</sup><br>throughfall NO3 <sup>-</sup> | $\begin{array}{c} -5.18 \pm 3.63^{a} \\ -5.71 \pm 2.82^{a} \\ -6.86 \pm 3.10^{a} \\ -6.16 \pm 2.29^{a} \end{array}$ | $\begin{array}{l} -5.84 \pm 3.31^{ab} \\ -6.19 \pm 2.34^{b} \\ -3.15 \pm 1.66^{a} \\ -4.17 \pm 0.58^{a} \end{array}$ | $\begin{array}{l} -3.48\pm 6.01^{a}\\ -4.10\pm 3.18^{a}\\ -6.57\pm 6.40^{a}\\ -6.02\pm 4.14^{a} \end{array}$ |

332

333

**334** 3.3.2. Comparison of  $\delta^{15}N$  values of <u>Sphagnum</u> and atmospheric deposition

335

**336** The  $\delta^{15}$ N values of living *Sphagnum* were between -6.2 and -1.9 ‰ (Tab. S1). The  $\delta^{15}$ N values of living

337 *Sphagnum* at BRU were statistically different from the  $\delta^{15}$ N values of atmospheric deposition (means of -4.0 and

-5.9 %, respectively; p < 0.05; Fig. 4). At UHL (means of -4.3 and -5.6 %, respectively;) and MMJ (means of -

4.4 and -4.3 ‰, respectively), the differences between the  $\delta^{15}$ N values of living *Sphagnum* and the  $\delta^{15}$ N values

340 of atmospheric deposition were insignificant (p > 0.05; Fig. 4). At BRU (but also at UHL), *Sphagnum* N was on

341 average isotopically heavier than deposited N, *i.e.*, closer to the 0 ‰ value of atmospheric N<sub>2</sub>. Nitrogen

342 concentration in living *Sphagnum* was significantly higher at MMJ (mean of 1.1 wt. %) than at UHL (0.9 wt. %;

343 p < 0.05; Fig. 5). The mean N concentration in BRU *Sphagnum* was 1.0 wt. %, indistinguishable from the other 344 two study sites.



**346** Fig. 4. Histograms of  $\delta^{15}$ N values of atmospheric input of N<sub>r</sub> and living *Sphagnum*. Different letters in







**Fig. 5.** Nitrogen concentrations in living *Sphagnum*. Different letters in superscript mark statistically different sample types (p < 0.05).

**352** *3.3.3. Multiple*  $\delta^{15}N$  comparisons among sample types

353

The mean  $\delta^{15}$ N value of surface bog water was 0.9 ‰ at BRU, 1.8 ‰ at UHL, and -1.9 ‰ at MMJ. Nitrogen in surface bog water was isotopically significantly heavier than N in both *Sphagnum* and atmospheric input at all three sites (Fig. 6; p < 0.05). At BRU and UHL, the mean  $\delta^{15}$ N value of surface bog water was higher than the 0

- 357 % value of atmospheric N<sub>2</sub>. At MMJ, the mean  $\delta^{15}$ N value of surface bog water was lower than the N isotope
- 358 signature of atmospheric N<sub>2</sub>. In other words, all three sample types (deposition, *Sphagnum*, and bog water) at
- $359 \qquad \text{MMJ contained isotopically lighter N, compared to atmospheric N_2 (Fig. 6).}$
- 360
- 361 When averaged across all depths (0-30 cm), the mean  $\delta^{15}N$  value in the peat core was -2.4 ‰ at BRU, -0.4 ‰ at
- 362 UHL, and -1.9 ‰ at MMJ. At all three sites, the maturating peat in the vertical profile contained isotopically
- **363** significantly heavier N compared to living *Sphagnum* (p < 0.05; Fig. 6; Tab. S2).

**365** The mean  $\delta^{15}$ N value of runoff was -2.7 ‰ at BRU (combined NH<sub>4</sub><sup>+</sup> and NO<sub>3</sub><sup>-</sup> data; number of observations n =

**366** 50), -5.3 ‰ at UHL (n = 6), and -5.1 ‰ at MMJ (n = 2; Tab. S1). The N isotope signature of runoff was higher

367 compared to the atmospheric input at BRU, and similar with the atmospheric input at UHL and MMJ (small

solid squares in Fig. 6). At all three sites, runoff contained isotopically lighter N compared to bog water (Fig. 6).

369



370

**Fig. 6.** Between-site comparisons of  $\delta^{15}$ N values of studied N pools and fluxes. Horizontal lines in boxes correspond to median values. Different letters mark statistically different sample types (p < 0.05).

373

**374** *3.4. Chemistry of natural waters* 

375

Acidity. Surface bog water had lower pH than atmospheric deposition and runoff at all three sites. Mean bog
water pH was 4.0 at UHL, 4.3 at BRU, and 4.9 at MMJ (Tab. S3; data for October 2018). The pH of atmospheric
deposition was lower than 5.0 only at UHL.

379

380 *Nitrogen.* The maximum NH<sub>4</sub><sup>+</sup>-N concentration in open area precipitation was 1.7 mg L<sup>-1</sup> (UHL) and the

381 maximum NO<sub>3</sub><sup>-</sup>N concentration in open area precipitation was 7.1 mg L<sup>-1</sup> (BRU; Tab. S2). The maximum

382 concentration of  $NH_4^+$ -N in throughfall was 3.9 mg  $L^{-1}(MMJ)$  and the maximum concentration of  $NO_3^-$ -N in

throughfall was 9.7 mg  $L^{-1}$  (BRU; Tab. S2). The maximum concentration of NH<sub>4</sub><sup>+</sup>-N in bog water was 2.3 mg  $L^{-1}$ 

384  $^{1}$ (UHL) and the maximum concentration of NO<sub>3</sub><sup>-</sup>-N in bog water was 2.7 mg L<sup>-1</sup> (MMJ; Tab. S2). The maximum

385 concentration of  $NH_4^+$ -N in runoff was 1.3 mg L<sup>-1</sup> (BRU) and the maximum concentration of  $NO_3^-$ -N in runoff 386 was 7.1 mg L<sup>-1</sup> (BRU; Tab. S2).

387

388*Phosphorus.* The mean concentration of total P in atmospheric deposition increased in the order: BRU (below389 $6.0 \ \mu g \ L^{-1}$ ) < UHL (9.3  $\ \mu g \ L^{-1}$ ) < MMJ (15.5  $\ \mu g \ L^{-1}$ ; Tab. S3; data for October 2018). Phosphorus concentration

in surface bog water was roughly 30 times higher than in atmospheric deposition at BRU, more than 50 times higher at UHL, and more than 10 times higher at MMJ (Tab. S3). The UHL bog water contained as much as 490  $\mu$ g P L<sup>-1</sup>. The mean P concentration in runoff increased in the order: MMJ (12.4  $\mu$ g L<sup>-1</sup>) < BRU (29.4  $\mu$ g L<sup>-1</sup>) < UHL (40.2  $\mu$ g L<sup>-1</sup>; Tab. S3).

394

395 *Other chemical species.* Natural waters at UHL were richer in sulfate  $(SO_4^{2-})$  than those at the remaining two 396 sites (Tab. S3). UHL bog water and runoff contained as much as 47.4 and 33.7 mg SO<sub>4</sub><sup>2-</sup> L<sup>-1</sup>, respectively. Bog 397 water was richer in potassium (K<sup>+</sup>) at UHL (9.05 mg L<sup>-1</sup>) compared to BRU and MMJ (1.85 and 1.97 mg L<sup>-1</sup>, 398 respectively). The concentration of DOC in atmospheric deposition was 2-4 times higher at MMJ than at the 399 remaining two sites (Tab. S3). In contrast, surface bog water at MMJ had 1.4 to twice lower DOC 400 concentrations, compared to the remaining two sites. Detailed water chemistry in October 2018 is in Tab. S3.

401

402 *3.5. Vertical peat profiles* 

403

404 From peat surface to the depth of 15 cm, peat density exhibited a slight increase similar at the three sites (Fig. 405 7a). Deeper, peat density remained relatively low (~0.05 g cm<sup>-3</sup>) at MMJ, and continued increasing irregularly at 406 BRU and UHL. Ash content remained below 5 wt. % to a depth of 30 cm at MMJ, and, with one exception, also 407 at BRU (Fig. 7b). The highest ash content was observed at UHL. Below the depth of 20 cm, it increased 408 downcore to values greater than 10 wt. %. The total N concentrations in peat substrate increased downcore or 409 exhibited a zigzag pattern (Fig. 7c). The UHL peat core was the richest in N in most 2-cm peat sections. Down to 410 a depth of 15 cm, N concentration was the lowest in MMJ peat. This contrasts with Fig. 5 bottom showing that 411 the average N concentrations in replicated grab samples of living Sphagnum were relatively high. The apparent 412 paradox, however, suffers from an uneven replication in Figs. 5 and 7 (n = 21 and n = 1, respectively). The N 413 concentration data in the MMJ peat core may just illustrate the large [N] variability at the moss surface. At all 414 three sites, the vertical  $\delta^{15}$ N profile was characterized by a downcore increase near the surface flattening out in 415 the deepest peat sections (Fig. 7d). Generally, the  $\delta^{15}$ N values in peat cores increased in the order BRU < MMJ < 416 UHL.

417

The nearly constant carbon (C) concentrations in peat were similar at all three sites to the depth of 20 cm, and became more variable deeper (Fig. 7e). The sharpest downcore decrease in the C:N ratio was found at MMJ, with the exception of the 0-to-4 cm depth where the C:N ratio increased (7f). Throughout the vertical peat profiles, P concentration was the lowest at BRU, and the highest at UHL (Fig. 7g). The N:P ratio was close to 12 throughout the UHL peat profile, increased downcore at MMJ from 10 to 20, and exhibited an irregular pattern at BRU, ranging between 20 and 40 (Fig. 7h). Further information on vertical changes in peat composition is in Tab. S4.





428 Fig. 7. Vertical changes in physicochemical characteristics of *Sphagnum* peat.

# 430 4. Discussion

# *4.1. The role of horizontal* N<sub>r</sub> deposition in peatlands

| 455  |  |
|--|--|
| 434  | Using field experiments, we have recently shown a sizeable contribution of horizontally deposited $N_r$ to total   |
| 435  | atmospheric deposition in Central European Sphagnum peat bogs (Novak et al., 2015b). During 80-90 days of  |
| 436  | the spring and fall foggy seasons, horizontal deposition added another 45 % to vertical deposition at Kunstatska   |
| 437  | Kaple Bog (KB), a mountain-top site in northern Czech Republic, and 14 % at Blatenska Slat (BS) in the less  |
| 438  | polluted southern Czech Republic (see Fig. 1 for location). Additionally, Hunova et al. (2023) reported a  |
| 439  | relatively high horizontal contribution of nitrate-N to winter-time atmospheric deposition in Czech mountains by   |
| 440  | analyzing ice accretions (mean of $29 \pm 3$ %; data for December–March; number of sites $n = 10$ ). As a first  |
| 441  | approximation, we suggest that the upper limit of the contribution of horizontal deposition to vertical deposition   |
| 442  | at BRU, UHL and MMJ could have been 30 %. If so, the total average $N_r$ deposition was slightly higher than 18  |
| 443  | kg ha <sup>-1</sup> yr <sup>-1</sup> at UHL and MMJ, and 16.5 kg ha <sup>-1</sup> yr <sup>-1</sup> at BRU (Tab. 1). Our study sites can thus be considered as  |
| 444  | highly or medium-polluted (Lamers et al., 2000). The overall $N_r$ pollution decreased in the order UHL > MMJ >  |
| 445  | BRU.   |
| 446  |  |
| 447  | We note that total atmospheric deposition may also contain measurable amounts of total organic N (TON;   |
| 448  | Violaki et al., 2010; Cornell, 2011). TON fluxes have not been considered as part of the Nr input in existing  |
| 449  | peatland BNF studies. Open-area precipitation at BRU, UHL and MMJ contained an additional 15, 45, and 13 %   |
| 450  | of total organic N, respectively, relative to the sum of the two inorganic $N_r$ forms (Tab. S3; October 2018). More   |
| 451  | TON data in precipitation would be needed to realistically estimate annual deposition of organic N at our study  |
|  |  |
| 452  | sites.   |
| 452<br>453   | sites.   |
| 452<br>453<br>454  | sites.<br>4.2. Relationship between N <sub>r</sub> pollution and N <sub>2</sub> -fixation  |
| 452<br>453<br>454<br>455   | sites.<br>4.2. Relationship between N <sub>r</sub> pollution and N <sub>2</sub> -fixation  |
| 452<br>453<br>454<br>455<br>456  | sites.<br><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i><br>In theory, chronic atmospheric deposition of pollutant N <sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019,  |
| 452<br>453<br>454<br>455<br>456<br>457   | sites.<br><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i><br>In theory, chronic atmospheric deposition of pollutant N <sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019,<br>2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a   |
| 452<br>453<br>454<br>455<br>456<br>457<br>458  | <ul> <li>sites.</li> <li><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i></li> <li>In theory, chronic atmospheric deposition of pollutant N<sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup> from the atmosphere, these authors reported a 54 %</li> </ul>   |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459   | <ul> <li>sites.</li> <li><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i></li> <li>In theory, chronic atmospheric deposition of pollutant N<sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>, a 69 % decrease under the</li> </ul>   |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460  | <ul> <li>sites.</li> <li><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i></li> <li>In theory, chronic atmospheric deposition of pollutant N<sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>, a 69 % decrease under the deposition of 17 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>, and a 74 % decrease under the deposition of 27 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>. As seen in Fig. 3,</li> </ul>  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461   | <ul> <li>sites.</li> <li><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i></li> <li>In theory, chronic atmospheric deposition of pollutant N<sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>, a 69 % decrease under the deposition of 17 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>, and a 74 % decrease under the deposition of 27 kg N<sub>r</sub> ha<sup>-1</sup> yr<sup>-1</sup>. As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least</li> </ul>  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462  | sites.<br><i>4.2. Relationship between</i> N <sub>r</sub> <i>pollution and</i> N <sub>2</sub> - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant N <sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the deposition of 17 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least polluted peat bog exhibited no instantaneous BNF, while MMJ, whose N <sub>r</sub> inputs were lower than those at UHL  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463   | sites.<br><i>4.2. Relationship between</i> N <sub>r</sub> <i>pollution and</i> N <sub>2</sub> - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant N <sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019,<br>2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a<br>practically unpolluted site receiving 2 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 %<br>decrease in BNF rates under the atmospheric deposition of 6 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the<br>deposition of 17 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3,<br>our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least<br>polluted peat bog exhibited no instantaneous BNF, while MMJ, whose N <sub>r</sub> inputs were lower than those at UHL<br>and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i>  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463<br>464                                    | sites.<br><i>4.2. Relationship between N<sub>r</sub> pollution and N<sub>2</sub>-fixation</i><br>In theory, chronic atmospheric deposition of pollutant N <sub>r</sub> should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the deposition of 17 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg N <sub>r</sub> ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least polluted peat bog exhibited no instantaneous BNF, while MMJ, whose N <sub>r</sub> inputs were lower than those at UHL and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i> bogs reported non-zero BNF rates regardless of atmospheric N <sub>r</sub> deposition level ( <i>see</i> compilation in Tab. S5),  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463<br>464<br>465                             | sites.<br><i>4.2. Relationship between</i> $N_r$ <i>pollution and</i> $N_2$ - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant $N_r$ should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the deposition of 17 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least polluted peat bog exhibited no instantaneous BNF, while MMJ, whose $N_r$ inputs were lower than those at UHL and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i> bogs reported non-zero BNF rates regardless of atmospheric $N_r$ deposition level ( <i>see</i> compilation in Tab. S5), non-detectable BNF rates at BRU and UHL were surprising. The mean instantaneous BNF rate at MMJ was   |
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| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463<br>464<br>465<br>466<br>467               | sites.<br><i>4.2. Relationship between</i> $N_r$ <i>pollution and</i> $N_2$ - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant $N_r$ should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the deposition of 17 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least polluted peat bog exhibited no instantaneous BNF, while MMJ, whose $N_r$ inputs were lower than those at UHL and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i> bogs reported non-zero BNF rates regardless of atmospheric $N_r$ deposition level ( <i>see</i> compilation in Tab. S5), non-detectable BNF rates at BRU and UHL were surprising. The mean instantaneous BNF rate at MMJ was lower than BNF rates in unpolluted high-latitude bogs in Canada (Vile et al., 2014) and Patagonia (Knorr et al., 2015). Among the studies listed in Tab. S5, the mean BNF rates at MMJ were the fourth highest. Our data from   |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463<br>464<br>465<br>466<br>467<br>468        | sites.<br><i>4.2. Relationship between</i> $N_r$ <i>pollution and</i> $N_2$ - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant $N_r$ should suppress BNF in peatlands (Wieder et al., 2019, 2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a practically unpolluted site receiving 2 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 % decrease in BNF rates under the atmospheric deposition of 6 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the deposition of 17 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3, our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least polluted peat bog exhibited no instantaneous BNF, while MMJ, whose $N_r$ inputs were lower than those at UHL and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i> bogs reported non-zero BNF rates regardless of atmospheric $N_r$ deposition level ( <i>see</i> compilation in Tab. S5), non-detectable BNF rates at BRU and UHL were surprising. The mean instantaneous BNF rate at MMJ was lower than BNF rates in unpolluted high-latitude bogs in Canada (Vile et al., 2014) and Patagonia (Knorr et al., 2015). Among the studies listed in Tab. S5, the mean BNF rates at MMJ were the fourth highest. Our data from MMJ are consistent with a conclusion by Saiz et al. (2021) who suggested a development of diazotrophic  |
| 452<br>453<br>454<br>455<br>456<br>457<br>458<br>459<br>460<br>461<br>462<br>463<br>464<br>465<br>466<br>467<br>468<br>469 | sites.<br><i>4.2. Relationship between</i> $N_r$ <i>pollution and</i> $N_2$ - <i>fixation</i><br>In theory, chronic atmospheric deposition of pollutant $N_r$ should suppress BNF in peatlands (Wieder et al., 2019,<br>2020). Saiz et al. (2021) quantified downregulation of BNF along a geographical pollution gradient. Relative to a<br>practically unpolluted site receiving 2 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> from the atmosphere, these authors reported a 54 %<br>decrease in BNF rates under the atmospheric deposition of 6 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , a 69 % decrease under the<br>deposition of 17 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> , and a 74 % decrease under the deposition of 27 kg $N_r$ ha <sup>-1</sup> yr <sup>-1</sup> . As seen in Fig. 3,<br>our data did not confirm such an inverse correlation at Central European sites. Instead, the most and least<br>polluted peat bog exhibited no instantaneous BNF, while MMJ, whose $N_r$ inputs were lower than those at UHL<br>and higher than those at BRU, showed a high mean BNF rate. Given that most previous studies of <i>Sphagnum</i><br>bogs reported non-zero BNF rates regardless of atmospheric $N_r$ deposition level ( <i>see</i> compilation in Tab. S5),<br>non-detectable BNF rates in unpolluted high-latitude bogs in Canada (Vile et al., 2014) and Patagonia (Knorr et al.,<br>2015). Among the studies listed in Tab. S5, the mean BNF rates at MMJ were the fourth highest. Our data from<br>MMJ are consistent with a conclusion by Saiz et al. (2021) who suggested a development of diazotrophic<br>microbes' tolerance to high rates of atmospheric $N_r$ deposition in recent decades. Global assessments of the |

| 470 | dependence of BNF on total Nr deposition are difficult to make for several reasons: (i) few studies consider  |
|-----|---|
| 471 | horizontal Nr deposition which may be sizeable and depends not just on atmospheric pollution, but also on   |
| 472 | elevation; few studies have quantified atmospheric input of organic N (ii) there is a large within-site   |
| 473 | heterogeneity in BNF ( ${}^{15}N_2$ incubations should be performed using a large number of replicates, see $\delta^{15}N$  |
| 474 | differences between individual MMJ replicates in Tab. 2; cf., "BNF hotspots" in Stuart et al., 2021); and (iii)   |
| 475 | recalculation between two commonly used BNF units (µg N per 1 g of Sphagnum d <sup>-1</sup> , g N m <sup>-2</sup> yr <sup>-1</sup> ) in literature  |
| 476 | data requires information on additional site-specific parameters, such as peat density, seasonality in daily  |
| 477 | temperatures and snow cover duration. Additionally, it is often unclear to what maximum depth in peat bogs  |
| 478 | BNF proceeds and whether there is a gradient in BNF rates within this depth range (Vile et al., 2014; Knorr et  |
| 479 | al., 2015).   |
| 480 |   |
| 481 | Since the differences in $N_r$ deposition among sites were minor (Tab. 1; Fig. 2), we suggest that $N_r$ deposition was   |
| 482 | not the primary control of the BNF rates in our study at the time of Sphagnum sampling.   |
| 483 |   |
| 484 | 4.3. Chemical and environmental parameters as possible BNF controls   |
| 485 |   |
| 486 | 4.3.1. The role of the $NH_4^+$ -N/NO <sub>3</sub> <sup>-</sup> -N ratio in atmospheric deposition  |
| 487 |   |
| 488 | The impact of the two main $N_r$ forms in deposition on BNF can be different. Because BNF generates $NH_4^+$ , the  |
| 489 | need for BNF to complement metabolic demands of the moss may be lower if deposition of $NH_4^+$ -N exceeds the  |
| 490 | deposition of NO <sub>3</sub> <sup>-</sup> -N (van den Elzen et al., 2018; Saiz et al., 2021). At our study sites, the NH <sub>4</sub> <sup>+</sup> -N/NO <sub>3</sub> <sup>-</sup> -N ratios |
| 491 | were nearly identical (Tab. 1), slightly exceeding 1. It follows that this ratio was unlikely the driver of higher  |
| 492 | BNF potential at MMJ, compared to the remaining two sites.  |
| 493 |   |
| 494 | 4.3.2. The effect of temperature  |
| 495 |   |
| 496 | MMJ is situated at a lower elevation, compared to UHL and BRU, and its mean annual temperature is higher  |
| 497 | than at the remaining two sites (Tab. 1). This could positively affect the rate of BNF (Basilier et al., 1978;  |
| 498 | Schwintzer et al., 1983; Urban and Eisenreich, 1988; Zivkovic et al., 2022; Yin et al., 2022). By contrast, Carrell   |
| 499 | et al. (2019) argued that BNF rates may decrease with an increasing temperature due to lower microbial diversity  |
| 500 | and greater mineralization rates leading to more $N_r$ in bog water and hence lower demand for BNF. Under field   |
| 501 | conditions of the Czech sites and at the peatland scale, temperature likely is a key factor regulating BNF. In our  |
| 502 | <sup>15</sup> N assimilation study, however, the chosen temperature was identical for all three sites. Consequently,  |
| 503 | temperature was not the dominant control of the measured short-term BNF rates.  |
| 504 |   |
| 505 | 4.3.3. The effect of bog wetness  |
| 506 |   |
| 507 | Fig. S2 shows monthly measurements of water table level below bog surface at BRU (Bohdalkova et al., 2013)  |
| 508 | and UHL (Tacheci, 2002). The mean annual water table depth was -5.2 $\pm$ 2.3 and -7.5 $\pm$ 1.1 cm at BRU at UHL,  |

respectively. No water level monitoring data are available for MMJ, however, during our field sampling

| 510 | campaigns, numerous 10-to-20 cm deep water pools were observed near the bog center at MMJ, especially               |
|-----|---|
| 511 | during the growing seasons of 2017 and 2019. Other high-elevation peat bogs on crystalline bedrock previously       |
| 512 | studied in the Czech Republic exhibited water table fluctuation at shallow depths of 5-8 cm, similar to BRU and     |
| 513 | MMJ (Novak and Pacherova, 2008). Based on visual inspection, somewhat drier conditions were typical of              |
| 514 | UHL, compared to the other two sites. Hydrological monitoring (GEOMON network database, Czech                       |
| 515 | Geological Survey; Oulehle et al., 2021b) revealed significantly drier conditions at UHL in the water year 2018,    |
| 516 | compared to the long-term average given in Tab. 1. Precipitation totals at UHL were 1460 mm in 2016, 1370           |
| 517 | mm in 2017, mere 892 mm in 2018, and 1230 mm in 2019. The ecosystem suffered from chronic drought in                |
| 518 | 2018 also at other GEOMON sites, JEZ (the nearest site to BRU) and UDL (the nearest site to MMJ; for location       |
| 519 | see Fig. 1). While Sphagnum for the ${}^{15}N_2$ incubation was collected at all three study sites at the same time |
| 520 | (October 2018), site-specific moisture conditions could have affected microbial community structure and the         |
| 521 | BNF potential. In the laboratory experiment, however, similar wetness was ensured by the same volume of             |
| 522 | added bog water to Sphagnum from all three sites. Therefore, we suggest that water availability did not control     |
| 523 | the instantaneous BNF rates.  |
|     |   |

#### 525 4.3.4. The effect of <u>Sphagnum</u> species

526

527 Stuart et al. (2021) showed that host identity is often the primary driver of BNF in peatlands. Under low  $N_r$ 528 pollution, higher species-specific litter decomposability augments BNF by increasing nutrient turnover (van den 529 Elzen et al., 2020). Saiz et al. (2021) observed higher BNF rates in Sphagnum species typical of hollows than 530 those dominating hummocks. Specifically, S. fallax exhibited higher BNF rates than S. capillifolium and S. 531 papillosum. The reason for such systematics appeared to be that the anoxic environment of wet hollows is more 532 favorable for N<sub>2</sub> fixers (Leppanen et al., 2015; Zivkovic et al., 2022). By contrast, Vile et al. (2014) observed 533 higher BNF rates in the hummock species S. fuscum than in the hollows species S. angustifolium. All moss 534 samples for our <sup>15</sup>N assimilation experiment were collected in lawns. One exception was a subordinated number 535 of plants of S. cuspidatum typical of hollows in the BRU incubation. While the moss species were identical in 536 the UHL and MMJ incubation (S. girgensohnii), the BNF potential at these two sites was strikingly different 537 (Fig. 3). Therefore, we suggest that *Sphagnum* species was not a key BNF control in our  $^{15}N_2$  experiment. 538

- 539 4.3.5. Organic N availability
- 540

541Wang et al. (2022) stressed the positive effect of organic N on BNF. Assimilation cost of amino acids was542shown to be lower than that of  $NH_4^+$  (Liu et al., 2013; Song et al., 2016). Organic N molecules can also serve as543a C source for cyanobacteria, thus saving the cost of photosynthesis (Krausfeld et al., 2019). As seen in Tab. S3,544concentrations of total organic N (TON) in bog water increased in the order: MMJ < BRU < UHL, and were thus</th>

probably unrelated to augmented BNF at MMJ sensu Wang et al. (2022).

- 545 546
- 547 4.3.6. Possible P limitation
- 548

- 549 Phosphorus is needed for the synthesis of ATP playing a key role in symbiotic BNF (Rousk et al., 2017; Wieder
- et al., 2022). In plant tissues, N:P ratios greater than 16 may indicate P limitation, while N:P ratios lower than 16
- 551 correspond to N limitation (Koerselman and Meuleman, 1996). Caution must be exercised in interpreting N:P
- ratios in atmospheric deposition as potential controls of P or N limitation. In addition to atmospheric input
- 553 fluxes, bioavailable P and N in bog waters are strongly affected by a tight inner cycling with additional inputs
- from biomass decomposition (Walbridge and Navaratnam, 2006). Phosphorus input fluxes *via* atmospheric
- deposition into peat bogs may affect nutrient limitation in the long-run, depending on whether these input fluxes
- are large enough, compared to the frequently observed P leaching to deeper peat layers (Walbridge and
- 557 Navaratnam, 2006, and references therein). According to Tab. S3, atmospheric deposition at all three study sites
- is consistent with P limitation that might limit BNF (high N:P ratios of 169, 60, and 112 at BRU, UHL, and
- 559 MMJ, respectively). At the same time, N:P ratios in surface bog water were below 16 at two of the three sites,
- 560 UHL (7.6), and MMJ (15). At BRU (N:P = 24), P limitation inferred from bog water chemistry would provide an
- explanation of non-detectable instantaneous BNF. At UHL, we found no indication of a relationship between P
- availability and zero BNF. The relatively P-rich bog water (165-490  $\mu$ g P L<sup>-1</sup>; Tab. S3) at all sites may contain,
- in addition to deposited P and mineralized P released during peat degradation, also, to some extent, geogenic P.
- 564 Bedrock granite (BRU, UHL) contains P in accessory apatite and K-feldspar whose weathering was probably
- 565 more efficient during the recent 40 years of acid rain. Phosphorus in phyllite (MMJ) is concentrated in apatite.
- Phosphorus concentrations in fresh bedrock were similar at BRU and MMJ (52-55 ppb), and twice lower at UHL
  (29 ppb; Gurtlerova et al., 1997; Pecina, 1999). The possible input of bioavailable geogenic P depended on local
  hydrology and could be site-specific.
- Living *Sphagnum* had N:P ratios of 31, 12, and 7 at BRU, UHL, and MMJ, respectively (Tab. S4), indicating
  conditions favorable for BNF at the latter two sites. As seen in Fig. 7h, N:P < 16 marking N-limitation was</li>
  characteristic of the entire vertical peat profile at UHL, and downcore to a depth of 15 cm at MMJ. In contrast,
  the N:P ratio was above 16 throughout the vertical peat profile at BRU. Phosphorus availability inferred from
- bog water and living *Sphagnum* gave consistent results with respect to possible BNF. As mentioned above, P
- 574 likely limited BNF only at BRU.
- 575
- 576 Recently, measurements of regional P deposition started in headwater catchments of the GEOMON network
- 577 (Oulehle et al., 2017). In the time period 2014-2018, UHL, a site directly included in the GEOMON network,
- 578 exhibited lower P concentrations in the atmospheric input, compared to JEZ in the west (a proxy of BRU) and
- 579 UDL in the east (proxy of MMJ; *see* Fig. 1 for catchment locations; the distance between JEZ and UDL, and
- between BRU and MMJ was approximately 70 km). Four-year average P concentrations at UHL were 72 and 36
- 581  $\mu g L^{-1}$  in open-area precipitation and spruce throughfall, respectively. At JEZ, analogous P concentrations were
- 582 103 and 87  $\mu$ g L<sup>-1</sup>, At UDL, these sample types contained on average 110 and 91  $\mu$ g P L<sup>-1</sup>. The high P uptake by
- tree canopy resulting in low P concentration in throughfall might indicate P deficiency in UHL inputs. At the
- same time, the N:P ratio in total vertical atmospheric deposition was lower than 16 at all three sites (13.1 at JEZ,
- 585 15.5 at UHL, and 13.7 at UDL (GEOMON Hydrochemical Database, Czech Geological Survey).
- 586
- 587 4.3.7. Possible Mo limitation
- 588

- 589 Nitrogenase requires molybdenum (Mo) in its active center to reduce  $N_2$  to bioavailable  $NH_4^+$  (Rousk et al.,
- 590 2017; Bellenger et al., 2020). In principle, Mo limitation of BNF may have played a role in the contrasting BNF
- 591 potentials observed at our sites. We do not have data on Mo concentrations in the studied ecosystems, except for
- trace metal analysis of the prevailing rock types ( $\leq 1$  ppm; Gurtlerova et al., 1997). However, known Mo
- 593 contents in coal massively mined/burnt in the Central European industrial region could shed some light on Mo
- availability *via* atmospheric deposition: North Bohemian soft coal (Sokolov basin close to BRU; Fig. 1) contains
- on average 18 ppm Mo, whereas Upper Silesian stone coal (Ostrava close to MMJ; Fig. 1) contains only ~0.6
- ppm Mo, *i.e.*, 30 times less (Bouska et al., 1997). Since UHL is situated downwind of the North Bohemian

Mo significantly influences the contrasting BNF potentials at our study sites.

- cluster of coal-burning power plants, and very close to Turow (soft coal mining in the Polish part of the Lusatianbasin; Fig. 1), atmospheric Mo inputs at UHL may be relatively high. Preliminarily, it appears to be unlikely that
- 599

600
601 4.3.8. The role of SO<sub>4</sub><sup>2-</sup> deposition

602

603 Large atmospheric inputs of acidifying sulfur forms (SO<sub>2</sub>, H<sub>2</sub>SO<sub>4</sub>), characterizing northern Czech Republic since 604 the 1950s (Hunova et al., 2022), can affect BNF in two ways: by suppressing methanogenesis, and by reducing 605 the pH. Sulfate in peat bogs under high S deposition becomes an important electron acceptor (Pester et al., 2012) 606 and bacterial sulfate reduction is thermodynamically favored relative to methanogenesis and fermentative 607 processes (Vile et al., 2003). It not only decreases gross CH<sub>4</sub> production in peat, mitigating the flux of CH<sub>4</sub> to the 608 atmosphere and minimizing climate warming, but also lowers the supply of  $CH_4$  to methanotrophs that, at some 609 sites, represent a major BNF pathway (Dise and Verry, 2001; Vile et al., 2014). Large  $SO_4^{2-}$  inputs may thus 610 suppress BNF in peat bogs. In this context, is should also be mentioned that a  ${}^{34}S/{}^{32}S$  isotope study has 611 documented post-depositional vertical mobility of S in industrially polluted peat bogs (Novak et al., 2009). 612 While long-term vertical S deposition, calculated according to Oulehle et al. (2016), was similarly high at UHL 613 and MMJ (means of 18.6 and 17.0 kg ha<sup>-1</sup> yr<sup>-1</sup> for the 1900-2012 period), higher than at BRU (12.2 kg ha<sup>-1</sup> yr<sup>-1</sup>), 614 UHL bog water at the time of this study was nearly 70 times richer in SO<sub>4</sub><sup>2-</sup> than MMJ bog water, and eight 615 times richer in  $SO_4^{2-}$  than BRU bog water (Tab. S3). Runoff at UHL was 4-5 times richer in  $SO_4^{2-}$  than runoff at 616 MMJ and BRU. The zero instantaneous BNF at UHL in our <sup>15</sup>N<sub>2</sub> incubation can be related to the highly elevated 617 S deposition in the case that UHL primarily hosts methane oxidizing diazotrophs.

618

619 UHL waters were characterized by lower pH, compared to those at MMJ and BRU (Tab. S3). Runoff pH at UHL

620 was 4.48, while runoff pH at MMJ was 7.40. Bog water pH at UHL was 4.02, while pH at MMJ was 4.88.

- 621 Downregulation of BNF in more acidic environment has been reported, *e.g.*, by Basilier (1979) and van den
- Elzen et al. (2017). Accordingly, lack of BNF at UHL may be related to its lower pH, compared to the other twostudy sites.
- 023 St
- 624
- 625 4.4. Natural-abundance N isotope systematics
- 626
- **627** Sphagnum metabolizes bioavailable  $NH_{4^+}$  approximately eight times faster than  $NO_3^-$  (Saiz et al., 2021). Because
- 628 there were nonsignificant differences between  $\delta^{15}N$  values of  $NH_4^+$  and  $NO_3^{-1}$  in rainfall at our study sites (Fig.

- 629 S1), it is reasonable to use the entire  $δ^{15}N$  data set for a comparison with  $δ^{15}N$  values of living *Sphagnum* (Fig.
- 630 4). Slow lateral mixing of surface bog waters may bring throughfall N from the forested margins of each bog to
- 631 the central unforested area and, therefore, we additionally included throughfall  $\delta^{15}N$  data in Fig. 4 comparisons.
- 632 The isotopically analyzed living *Sphagnum* plants represented on average a one-to-two-year increment (*cf.*,
- 633 Wieder and Vitt, 2006). We found a statistically significant shift from isotopically light N of the deposition to
- 634 isotopically heavier N of *Sphagnum* only at BRU (p < 0.05). This might indicate mixing with even heavier
- $\label{eq:states} \textbf{635} \qquad \text{atmospheric $N_2$ taken up by diazotrophs. At BRU, BNF might have intermittently proceeded over the most}$
- $\label{eq:constraint} 636 \qquad \text{recent growing seasons even though the $^{15}N_2$ experiment did not corroborate this process in October 2018.}$
- 637
- 638 A straightforward attribution of the N isotope pattern at BRU to BNF, however, is hampered by the fact that 639 mineralization is a likely alternate source of dissolved  $N_r$  for assimilation by the moss (Zivkovic et al., 2022, and
- 639 mineralization is a likely alternate source of dissolved N<sub>r</sub> for assimilation by the moss (Zivkovic et al., 2022, and 640 references therein). The often found high  $\delta^{15}$ N values of mineralized N<sub>r</sub> remaining in the bog ecosystem result
- for an isotope fractionation accompanying denitrification, a process known to occur especially in peat bogs
- from an isotope fractionation accompanying denitrification, a process known to occur especially in peat bogsthat are not extremely acidic. Gaseous products of denitrification contain isotopically light N both in wetlands
- that are not extremely acidic. Gaseous products of denitrification contain isotopically light N both in wetlands(Denk et al., 2017; for data from Czech peat bogs *see* Novak and al., 2015a, 2018), and aerated forest soils
- 644 (Houlton and Bai, 2009; for data from Czech upland soils *see* Oulehle et al. 2021a). Nitrogen in surface bog
- 645 water at BRU had a positive mean  $\delta^{15}$ N value of 0.9 % (Fig. 6). Isotope systematics at BRU are thus consistent
- 646 with incorporation of mineralized  $N_r$  into moss biomass during assimilation instead of uptake of N resulting from
- 647 BNF.
- 648
- 649 Advancing mineralization accompanying peat maturation with mobilization and export of gaseous low- $\delta^{15}N$ 650 nitrogen is also responsible for the increasing  $\delta^{15}N$  values of the residual peat substrate downcore (Fig. 7d).
- 651
- Fig. S3 summarizes two general scenarios, under which a difference between N isotope composition of
- atmospheric input, *Sphagnum* and bog water indicates BNF: (1) the mean  $\delta^{15}$ N values increase in the order:
- deposited  $N_r < bog$  water  $N_r < Sphagnum N_r < atmospheric N_2$ , or (2) the mean  $\delta^{15}N$  values decrease in the
- order: deposited  $N_r > bog$  water  $N_r > Sphagnum N_r > atmospheric N_2$ . Whereas the  $\delta^{15}N$  value of bulk
- atmospheric deposition in Central Europe is mostly negative, positive mean  $\delta^{15}N$  values have been reported from
- other regions. One example is isotopically heavy N of dry-deposited HNO<sub>3</sub> in an industrial part of the U.S.
- **658** (Elliott et al., 2009). Fig. S3 assumes that the magnitude of potential N isotope fractionations during uptake of
- inorganic N into plant biomass is relatively small and does not overprint the larger N isotope differences
- between the above discussed mixing endmembers.
- 661
- 662 It remains to be seen how to reconcile the relatively high instantaneous BNF rate at MMJ, measured in the
- laboratory, with the non-existence of a positive  $\delta^{15}N$  shift from atmospheric deposition (mean of -4.3 ‰) to
- **664** Sphagnum (mean of -4.4 ‰; Fig. 4; p > 0.05). Given that we explained the positive  $\delta^{15}$ N shift from deposition to
- 665 Sphagnum at BRU by mixing of low- $\delta^{15}$ N rainfall with high- $\delta^{15}$ N bog water, and that bog-water N at MMJ is
- isotopically heavy, a similar positive N isotope shift from rainfall to *Sphagnum* would be expected also at MMJ.
- 667 Such was not the case. This observation is important because it might indicate that uptake of recently
- 668 mineralized Nr from bog water at sites hydrologically similar to MMJ (and also BRU) may not control the N

- isotope signature of living *Sphagnum*. An input of isotopically light N<sub>r</sub> for assimilation by the MMJ moss could,
- 670 in principle, originate from shallow groundwater upwelling or lateral water inflow from other segments of the
- 671 catchment possibly bringing legacy low- $\delta^{15}$ N nitrogen from the peak acid-rain period throughfall. Such within-
- 672 site water inputs could affect the intermediate  $\delta^{15}$ N value of *Sphagnum* at MMJ.
- 673

### 674 Conclusions

675

676 Based on hydrochemical monitoring data and statistical modelling, the three studied Sphagnum peat bogs located 677 in the industrial northern Czech Republic received close to 18 kg Nr ha<sup>-1</sup> yr<sup>-1</sup> via atmospheric deposition. Since 678 1900, the atmospheric input of  $N_r$  affected the study sites in the order: UHL > MMJ > BRU. In the most recent 679 years, the annual N<sub>r</sub> inputs via vertical deposition between the sites differed by mere 1 to 2 kg ha<sup>-1</sup> yr<sup>-1</sup>. The sites 680 can thus be classified as highly to medium-polluted. A 168-hour  $^{15}N_2$  assimilation experiment revealed relatively 681 high but variable rates of BNF at MMJ, and non-detectable BNF at the remaining two sites, characterized by 682 slightly higher and slightly lower  $N_r$  depositions, respectively, compared to MMJ. We investigated in all 10 683 different parameters that might have served as controls of the presence or absence of instantaneous BNF in 684 living moss. In addition to bulk  $N_r$  deposition fluxes, these parameters included:  $NH_4^+-N/NO_3^--N$  ratio in 685 atmospheric input, temperature, wetness, Sphagnum species, organic-N availability, possible P limitation, 686 possible Mo limitation,  $SO_4^{2-}$  deposition, and pH. Using the available data, we argue that P deficiency was the 687 likely inhibitor of BNF at BRU. Assuming that methanotrophic bacteria represented a major type of diazotrophs, 688 extremely high  $SO_4^{2-}$  inputs may have been the key control of the absence of BNF at UHL. While the long-term 689 temperature and wetness at UHL were also lower, compared to the remaining two sites, they probably did not 690 affect the results of the  ${}^{15}N_2$  experiment since the incubation was performed under the same temperature and 691 wetness for all sites. In general, higher concentrations of decomposition-inhibiting metabolites could be causally 692 related to BNF rates. Such a control of BNF was unlikely since the same Sphagnum species from MMJ and UHL 693 was used for the  ${}^{15}N_2$  experiment that showed contrasting results for these two sites. The large  $\delta^{15}N$  differences 694 between moss replicates that were collected from various segments of MMJ at the end of the <sup>15</sup>N<sub>2</sub> incubation 695 suggested an existence of BNF hotspots.

696

697 The use of natural-abundance N isotope ratios to corroborate the observed instantaneous BNF rates was 698 hampered by isotopically heavy N of surface bog water. The bog water contained secondary  $N_r$  forms which 699 could have resulted from partial *Sphagnum*/peat decomposition and removal of the complementary low- $\delta^{15}$ N 700 products of denitrification. At BRU, we found statistically significant differences in  $\delta^{15}$ N values in the order: 701 deposited  $N_r < Sphagnum N_r < atmospheric N_2 < bog water N_r$ . Stable isotope ratios could not unambiguously 702 distinguish between assimilation of bog-water  $N_r$  and atmospheric  $N_2$  to form the observed N-isotope signature 703 of Sphagnum. At UHL and MMJ,  $\delta^{15}$ N differences between Sphagnum and the atmospheric input were 704 statistically insignificant. The natural-abundance approach as a test of BNF presence may give more promising 705 results at high-latitude sites often characterized by greater (30-40 cm) depth of the water table level below 706 Sphagnum capitula than the Central European sites. 707

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|-------|--|
| 709   | editing; M. Novak: conceptualization, data interpretation, writing - original draft; B. Cejkova: methodology,    |
| 710   | nitrogen fixation data acquisition, data interpretation; I. Jackova: methodology, concentration and isotope data |
| 711   | acquisition; F. Buzek: methodology, data interpretation, validation; F. Veselovsky: field work; J. Curik: field  |
| 712   | work; E. Prechova: formal analysis, resources; A. Komarek: statistical analysis; L. Bohdalkova: data             |
| 713   | interpretation   |
| 714   |  |
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