1 Soil stabilisatizion for DNA metabarcoding of plants and fungi. Implications

2 for sampling at remote locations or via third-parties

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11 Abstract

12 Storage of soil samples prior to metagenomic analysis presents a problem. If field sites are remote or 13 if samples are collected by third parties, transport to analytical laboratories may take several days or 14 even weeks. The bulk of such samples and requirement for later homogenisation precludes the 15 convenient use of a stabilisation buffer, so samples are usually cooled or frozen during transit. There 16 has been limited testing of the most appropriate storage methods for later study of soil organisms 17 by eDNA approaches. Here we tested a range of storage methods on two contrasting soils, 18 comparing these methods to the control of freezing at -80°C followed by freeze-drying. To our 19 knowledge this is the first study to examine the effect of storage conditions on eukaryote DNA in 20 soil, including both viable organisms (fungi) and DNA contained within dying/dead tissues (plants). 21 For fungi, the best storage regimes (closest to the control) were storage a 4°C (for up to 14 d) or 22 active air-drying at room temperature. The worst treatments involved initial freezing followed by 23 thawing which led to significant later spoilage. The key spoilage organisms were identified as 24 Metarhizium carneum and Mortierella spp., with a general increase in saprotrophic fungi and 25 reduced abundances of mycorrhizal/biotrophic fungi. Plant data showed a similar pattern but with 26 greater variability in community structure especially in the freeze-thaw treatments, probably due to 27 stochastic variation in substrates for fungal decomposition, algal proliferation and some seed 28 germination. In the absence of freeze drying facilities, samples should be shipped refrigerated but 29 not frozen if there is any risk of thawing.

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31 Keywords: Freeze-drying; chitinolytic fungi; Freeze-thaw; sample preservation

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34 Introduction

35 The use of eDNA metabarcoding (amplicon sequencing) has transformed our knowledge of the 36 structure and composition of soil biological communities (Geml et al., 2014; Williams, 2020), with 37 more recent metagenomic studies enhancing our understanding of the metabolic processes 38 mediated by these organisms (Keepers et al., 2019; Ogwu et al., 2019). However, the methods used 39 to sample soils (Epp et al., 2012; Lindahl et al., 2013; Taberlet et al., 2012), store and extract total 40 soil DNA/RNA (Kennedy et al., 2014; Soliman et al., 2017) can exert a strong influence of the data 41 obtained, as can the barcoding loci/primers and sequencing platforms used. Thus there is a need for 42 a standard operating procedure (SOP) before metabarcoding analyses conducted in different 43 laboratories can be compared (Lindahl et al., 2013; Orgiazzi et al., 2015).

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As the use of eDNA metabarcoding has extended to the study of soils in more remote locations (Detheridge et al., 2020; Tedersoo et al., 2014) and to more applied deployment in nature conservation site monitoring by statutory organisations (Detheridge et al., 2018; Geml et al., 2014; Latch, 2020; Valentin et al., 2020), the issue of soil storage between sampling and subsequent analysis has become an important consideration. Such concerns have interested soil scientists for many decades but usually in relation to the metabolic status of soil organisms, assessed via community level physiological profiles (CLPP), soil respiration etc. (Lee et al., 2007). However, for

52 metabarcoding / metagenomic studies, the preservation of DNA/RNA unchanged from the natural 53 state presents its own distinctive challenges.

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There are established international guidelines recommending refrigerated storage or freezing (OECD, 2000) when chemical or microbiological analyses cannot be undertaken on fresh soil. However, the preservation of soils by air-drying dates back to the origins of soil science and remains the simplest method for long-term stabilisation. Examination of air-dried soil archive samples dating back over a century by Clark et al. (2008) found that whilst long-term storage of air-dried soil reduced the amount of DNA present, differences in bacterial populations according to soil plot treatment were still detectable.

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63 Current standards have been established through the International Organization for Standardization (ISO), for example ISO 11063 (2012) ("Soil quality - Method to directly extract DNA from soil 64 65 samples") and ISO-10381-6 (2002) ("Soil quality—Sampling—Part 6: Guidance on the collection, 66 handling and storage of soil for the assessment of aerobic microbial processes in the laboratory"). 67 However, these are focused predominantly on bacteria (Petric et al., 2011), which appear to respond 68 in storage rather differently from fungi and other groups of biota (Terrat et al., 2015). However, it is 69 unclear whether this is due to the greater tolerance of bacterial cells to disruption by freezing etc. or 70 the lower taxonomic resolution of the standard 16S metabarcoding procedures.

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Martí et al. (2012) examined the effects of different soil storage and found that the stability of bacterial DNA (assessed via DGGE) varied according to soil type. Similarly Lauber et al. (2010), used DNA metabarcoding of bacterial populations to examine the effects of storage of soil or faeces at a range of temperatures (-80°C, -20°C, 4°C, 20°C) for 3 or 14 days. Even after 14 d at 20°C (in sealed container), they found only small changes in bacterial communities. However, Rubin et al. (2013), also using DNA metabarcoding to assess changes in soil bacterial populations, found that there was a progressive loss of diversity associated with storage under warmer conditions.

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80 Freeze drying (with subsequent frozen storage) is widely considered by many to be the best available 81 option for the stabilisation of soils prior to nuclei acid extraction (Castaño et al., 2016; Straube and 82 Juen, 2013; Weißbecker et al., 2017). Initial freezing inactivates biological processes and later 83 removal of water via sublimation at low pressure, stabilises the soil in an inactive dry state. 84 However, Bainard et al. (2010) found a reduction in arbuscular mycorrhizal fungal DNA in roots 85 following long-term storage of freeze-dried roots at ambient temperature, so frozen storage 86 following initial processing is important here too. There are significant additional advantages to 87 freeze-drying. Freshly-collected soils can be frozen immediately without prior processing and after 88 freeze-drying stored frozen in a stable state, convenient (unlike directly frozen soil) for later 89 homogenisation/grinding. The ease with which freeze-dried soils can be finely ground, allows for 90 efficient homogenisation of larger samples. The latter point is important, since it is crucial that 91 subsampling for DNA extraction (commonly 200 mg from 500g samples) is representative of the 92 whole (Lindahl et al., 2013). Repeated DNA extractions from a fully homogenised soil would 93 therefore result in the same community structure derived from metabarcoding. However, 94 availability of suitably large freeze-drying capacity can be an important limiting factor at many 95 institutions.

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97 Where it is not possible to freeze samples within a few hours of collection, the question remains as 98 to what pre-treatment is best to preserve the nucleic acids of the soil communities during shipping 99 from field sites (often sampled by third parties) to analytical labs. Here we compare the effect of 100 immediate freezing to a range of different soil DNA stabilisation methods, using equipment available 101 outside laboratories (freezers, fridges, fans and ovens). The resulting effects of these soil storage 102 methods are examined using eDNA metabarcoding profiles for plants and fungi, hypothesising that 103 inferior storage conditions would lead to: a general loss of diversity in both plants and fungi, due to 104 DNA degradation; proliferation of a subpopulation of faster-growing fungi, well-suited to growth in

105 particular storage conditions, which would be associated with greater levels of DNA degradation.

107 Methods.

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108 Soil was collected from an upland (grazed) grassland immediately adjacent to the Brignant longterm 109 grazing experimental field site (lat/long: 52.3648°N, 3.8214°W; 367m asl.). Brignant soil is an acidic 110 (pH 5) Manod Series (loam over shale Palaeozoic slate, mudstone and siltstone; well-drained fine 111 loamy or fine silty soils over rock; (Hallett et al., 2017)) with an organic carbon content of 7.3%. Turf 112 was removed to a depth of 3-5 cm from a single 0.25 m^2 area and approximately 10 kg of soil collected from the remaining 10 cm depth of topsoil avoiding large stones. A contrasting less acidic 113 114 and lower organic matter, alluvial soil type (Conway series 0811b [Fluvic Eutric Gleysols] 115 groundwater gley (silt to silty-clay loam. Depth to gley layer varies @ 10 - 20 cm.)(Hallett et al., 2017)) pH 6.1, organic carbon 3.6%, was also collected (15 kg) from the edge of an arable field at 116 117 Gogerddan (52.4364°N, 4.0313°W; 15 m a.s.l.), with removal of vegetation, as above, and processed 118 in the same way. The soils were transported within 2 h to the laboratory and sieved (3 mm) to 119 remove further stones and to enable thorough homogenisation of the soil/roots. Samples of 200 g 120 were weighed into Ziploc bags (40 in total) and divided into treatments with 4 replicates per 121 treatment. Bags for the control treatment (Treatment 1) were immediately frozen at -80°C.

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123 The soil treatments are shown in Table 1. Treatments T2, T3 and T4 were designed to test the effect 124 of initial freezing followed by thawing during shipping either refrigerated for 14 d (T2) or 5 d (T3), or 125 stored at ambient temperature (23°C, T4). Treatments T5, T6 and T7 were all dried for 5 d after an 126 initial "shipping" period (storage for 3d at 4°C). T5 was dried more rapidly by blowing ambient air 127 into open bags (hairdrier but no heat; as shown in SuppFig 1), whereas T6 and T7 were passively 128 dried in open bags at room temperature and 37°C respectively. Treatments T8, T9 and T10 involved 129 storage in closed bags either at 4°C for 14 d (T8) or at ambient temperature (T9 for 14 d and T10 for 130 5 d). Five of these 10 treatments (T1, T3, T6, T8 and T9) were also applied to the alluvial soil to see if 131 a contrasting soil responded to storage in a similar manner.

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133 Table 1. Storage treatments tested in this study using soil adjacent to the Brignant longterm 134 experiment. An * indicates treatments also tested with soil from the Gogerddan alluvial plain.

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SuppData 1. Photo of the hairdrier apparatus used for active air-drying of soil (treatment T5)

138 After the storage treatments were completed, all bags were frozen at -80°C. Samples were then 139 freeze-dried before sieving at 0.5 mm and thoroughly homogenised according to our standard lab 140 procedure (Detheridge et al., 2016). DNA was extracted from 200mg of freeze dried soil using the 141 Power Soil DNA extraction kit (Qiagen), as described by Detheridge et al. (2016). The ITS2 region of 142 plants and fungi were amplified using a mix of primers. For fungi, the forward primers were those 143 used by Tedersoo et al. (2014), with an equimolar mix of 6 primers (SuppData 2). To this mix the 144 plant primer Chen S2F (ATGCGATACTTGGTGTGAAT) (Chen et al., 2010) was added in a ratio of 3 145 fungal mix to 1 plant primer. This ratio was chosen to ensure that the majority of sequences 146 returned were fungal as this is the prime aim of the analysis and fungal communities are generally more complex than plant communities. The reverse primer was the universal ITS4 primer. The 147 148 forward primers were linked at the 5' end to the Ion Torrent B adapter sequence 149 (CCTCTCTATGGGCAGTCGGTGAT). The ITS4 primer was linked at the 5' end to the Ion Torrent A-150 adapter sequence (CCATCTCATCCCTGCGTGTCTCCGAC), the TCAG key and an IonXpress Barcode.

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152 SuppData 2. Fungal forward primer sequences with target group to amplify all fungal groups, and 153 also Stramenopiles (Ooomyces) devised by Tedersoo et al. (2014)

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PCR was carried out using PCR Biosystems Ultra polymerase mix (PCR Biosystems Ltd, London UK).
Each reaction contained 250 nM of the forward primer mix and 250 nM of reverse primer.
Amplification conditions were initial denaturing 15 min at 95°C, followed by 30 cycles at 95°C for 30
s, 55°C annealing for 30 s, 72°C extension for 30 s, and a final extension of 5 min at 72°C.

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After PCR, samples were processed, sequenced and sequence data processed as detailed in Detheridge et al. (2016; 2018; 2020). Fungal sequences were identified using a database build from v8.0 of UNITE (Abarenkov et al., 2019) and plants sequences using a database built as detailed in Detheridge et al. (2020). Sequence data have been submitted to the European Nucleotide Archive with reference number XXXXXXX. Data were expressed as relative abundance of each of the species detected (separately for plants and fungi).

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167 Principal coordinate ordination (PCO) visualised differences in community structure using square 168 root transformed abundances and a Bray-Curtis distance matrix; these analyses were undertaken in 169 R (R Core Team, 2013). PERMANOVA determined whether there were significant differences in 170 fungal and plant communities between treatments and the pairwise test used to determine which 171 treatments differed and their degree of separation. An analysis of similarity (SIMPER) was used to 172 determine which OTUs varied between treatments. These analyses were conducted in PRIMER-173 PERMANOVA + v6. ANOVA was used to determine the significance of treatment effects on relative abundance and these were carried (in R), after any appropriate transformations to meet 174 175 requirements of analyses.

176

177 Results

After quality checking there were a total of 2 772 707 ITS2 sequences with a maximum of 94 109 sequences per sample and a minimum of 52 931 (Mean 69 318). After rarefying to the lowest number of sequences per sample, dropping singleton sequences and trimming 5.8S and 28S regions, clustering resulted in 848 plant and fungal OTUs for the upland (Brignant) soil and 769 for the alluvial (Gogerddan) soil.

183

184 Upland (Brignant) soil: Principal coordinate ordinations of the fungal community data (Fig 1 A), show 185 the clear outlier relative to T1 (control) is T4 (Freeze-thaw, left at 23°C for 14 d), with T2/T3 (Freeze-186 thaw, left at 4°C for 5 d/14 d respectively) and T7 (storage at 4°C followed by passive warm air-187 drying at 37°C) being the next most divergent treatments. PERMANOVA analyses showed a 188 significant effect of storage treatment for the fungal community data (Pseudo F = 5.1728 P= 0.001). 189 Pairwise Permanova comparisons of the fungal populations in each treatment with the control (Fig. 190 2A) confirm that treatments T2, T3, T4, and T7 had a significant effect on the fungal populations 191 present at the end of the storage period. However, for the other pre-treatments there were no 192 significant differences relative to control.

193

Fig. 1. Principal coordinate diagrams of the fungal community data (A) and plant community data (B)
highlighting the difference in community between the different soil storage treatments. Points show
the mean axis scores and error bars show standard error of the mean.

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Fig. 2. Levels of t-statistic from pairwise Permanova of the control treatment compared to all other
 treatments. The p value is shown above the bar with significant (P<0.05) values shown in red. (A)
 fungal community data (B) plant community data.

201

Similar analyses for the effect of different treatments on the plant DNA (including algae:
 Chlorophyta) remaining after storage show that the general level of divergence from the control was
 less than for fungi but still significant (Permanova Pseudo F = 2.5117 P = 0.001). Here too freeze-thaw

treatments were also the most divergent (Fig. 1B, Fig. 2B) and there is similarity in the PCO ordinations for plant and fungal data. This similarity was corroborated by a Mantel test of the difference matrices, which revealed a Pearson correlation coefficient of 0.54 (P=0.0001). In contrast to the broader trend, the warm air-drying treatment (T7) had very different effect on the trajectory of the resulting plant and fungal communities later detected, causing significant change to the fungal community present (Fig. 2A) but very little effect on the plant DNA later recovered.

211

212 Apart from the treatments mentioned above, most treatments involving storage of soil at 4°C or at 213 ambient temperature for up to 14 d did not result in significant changes to the plant or fungal 214 populations later detected. In PCO ordination, the 4°C for 14 d (T8) treatment was closest to the 215 control for both plants and fungi (Figs. 1A/1B). For the fungal community data, treatment 10 (closed 216 bag, ambient temp) in particular showed an increase in the spread of data in both primary axis 217 dimensions. For plants, in contrast to the fungi, the variance of axis scores (especially PCO1) for most 218 treatments were larger than for the control treatment(Fig. 1B). This reduces the ability of statistical 219 analyses to find a significant effect between treatments and may be related to the proliferation of 220 spoilage fungi during storage.

221

222 Apart from green algae (Chlorophyta) which comprised <1% of total plant DNA in most treatments, 223 the plant DNA present in the sieved soils was mainly within dead or dying tissues (e.g. fine roots). In 224 contrast, a significant component of the fungal community would likely remain viable in the short 225 term, with some species proliferating if storage conditions are conducive to their growth. Since fungi 226 are the main decomposers of plant-derived lignocellulose in soil in terrestrial ecosystems, it is likely 227 that proliferation of certain fungi would be associated with more rapid degradation of plant DNA. 228 The relative sequence abundance of fungi and plants did significantly vary between treatments (Fig. 229 3). There were significant increases in the representation of fungi (and thus decrease in plants) 230 relative to the control (T1), which was higher in two of the freeze-thaw treatments (T2 and T4) and 231 T9 (23°C for 14 d). The relative proportions of fungi and plant sequences (Fig. 3) is clearly 232 determined by the the exact mix of plant and fungal primers used but in our experiments the same 233 mix was used for all samples, and therefore the changes reflect actual patterns of DNA 234 degradation/proliferation.

235

239

Fig. 3. Relative sequence abundance of fungi to plants by treatment. Letters on the bars indicate
 significant groupings as determined by Tukey's HSD post hoc test and error bars show standard error
 of the mean.

Some storage treatments led to a reduction in fungal species diversity (SuppData 3A/3B) relative to control, notably the freeze-thaw treatments (T2, T3, T4) and T7 (warm air-drying). In addition to significant differences in diversity between treatments, some treatments notably T5, T7 and T10, showed an increased spread of index values by replicate within treatment, as can be seen by the larger error bars. For plants (SuppData 3C/3D) reductions in species diversity were less pronounced, with only T4 differing significantly from control.

246

SuppData 3. Variations in diversity indices by storage treatment for the upland (Brignant)
 soil A) Fungi Simpson diversity index B) Fungi Shannon diversity index. C) Plant Simpson
 diversity index D) Plant Shannon diversity index. Letters above the bars indicate significant
 groupings as determined by Tukey's HSD post hoc test and error bars show standard error
 of the mean. Note that Shannon and Simpson indices are scaled inversely (i.e. higher index = lower
 diversity).

253

254 More detailed examination of the differences in the fungal community composition with treatment 255 reveal a large increase in abundance of Ascomycota relative to Basidiomycota for treatment 4 (Mean

256 2.83) compared to the control treatment (Mean 0.70) with all other treatments remaining very
257 similar to the control (Fig. 4A). This change was mainly due to the ca. 10-fold (19.4 vs. 1.5%)
258 increased abundance of the ascomycete *Metarhizium* (formerly *Paecilomyces*) *carneum* (UNITE
259 species hypothesis SH1552520.08FU) following freeze-thaw and storage for 14d at 23°C (Fig. 4B).
260 Mortierellomycota showed a similar 8-fold increase relative to control (17.8 vs 2.2%) in the freeze261 thaw treatment stored at 4°C for 14 d and were also more abundant in treatments T4 and T8 (Fig.
262 4C).

263

Analysis of the functional grouping, as determined unambiguously by FUNguild (Nguyen et al., 2016),
 revealed that 'saprotrophic fungi' demonstrated a similar but less pronounced trend, with a higher
 relative abundance in T4 relative to control (22.85% vs. 17.01%) (Fig. 4D). It should be noted that
 Mortierellomycota and *Metarhizium* spp. are classed as 'symbiotroph' and 'animal pathogen'
 respectively in FUNGuild.

269

Fig. 4. Variations in relative abundance of key fungal groups by storage treatment A) Ratio of
 Ascomycota to Basidiomycota; B) *Metarhizium carneum*; C) Mortierellomycotina; D) Saprotrophic
 fungi; E) Glomeromycotina (Arbuscular mycorrhizal fungi); F) Grassland fungi (CHEGD). Letters above
 the bars indicate significant groupings as determined by Tukey's HSD post hoc test and error bars
 show standard error of the mean.

275

276 As might be expected following disruption of active plant hosts, abundance of arbuscular 277 mycorrhizal fungi (AMF; subphylum Glomeromycotina) was reduced under the three freeze-thaw 278 storage conditions, with a 6-fold reduction in T4. Other fungi suspected to be mycorrhizal or with 279 intricate biotrophic association with higher plants also showed large reductions in abundance, 280 notably the CHEGD fungi. These fungi, mainly basidiomycetes and comprising members of the 281 families <u>C</u>lavariaceae, <u>H</u>ygrophoraceae, <u>E</u>ntolomataceae and <u>G</u>eoglossaceae, are dominant 282 components of undisturbed grassland habitats. Combined abundance of CHEGD fungi was 5-fold 283 lower in treatment T4 and also significantly lower in treatments T2, T3 and T7 (Fig. 4D). Analysis of 284 the individual components of the CHEG fungi revealed that the Clavariaceae, Hygrophoraceae, 285 Geoglossomycetes varied by treatment with significantly lower relative abundances in treatment T4. 286 However, there was no significant difference by treatment for Entolomataceae (SuppData 4). The 287 two dominant CHEGD species in the original Brignant soil were Clavulinopsis laeticolor (UNITE 288 SH1611741.08FU) and Hygrocybe chlorophana (SH1546991.08FU) with mean abundance in the 289 control (T1) soil of 21.5% and 9.4% respectively, with these levels being 4-fold and 15-fold lower in 290 the most unfavourable storage regime (T4; freeze-thaw followed by 14d at 23°C).

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SuppData 4. Relative abundance of CHEG fungi by storage treatment for upland (Brignant) soil A)
 Hygrophoraceae B) Clavariaceae. C) Geoglossomycetes D) Entolomataceae. Letters above the bars
 indicate significant groupings as determined by Tukey's HSD post hoc test and error bars show
 standard error of the mean.

296

297 In the Brignant soil, grasses (Poaceae; 9 spp.) were dominant (mean 88.8% of plant sequences in 298 control treatment T1) followed by Brassicaceae (Cardamine pratensis; 5.57%), Asteraceae (3 spp.; 299 2.58%) and Trifolium repens (Fabaceae; 1.33%), with algae (Chlorophyta) comprising 0.41% of the 300 plant sequences in the control soil. The turf layer was removed during sample collection, so the 301 higher plant tissues comprised mainly (live or dead) root tissues. Several species (e.g. Crepis 302 capillaris, Hypochaeris radicata, Ranunculus repens, Cerastium glomeratum) were detected in three 303 or fewer of the initial 40 sieved soil samples, probably due to heterogeneous distribution of larger 304 pieces of taproot tissue. The abundance of Poaceae varied by treatment (Suppdata 5) but this 305 variation was not significant because of the broad range of data in some treatment replicates (e.g. 306 3.7% to 68.2% in treatment 4). The greatest treatment effect on plant populations was the ca. 20fold increase in abundance of Chlorophyta in T4, likely due to tolerance of these microbes to freezing
 and later proliferation inside the clear plastic bags when incubated under ambient indoor lighting.

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Suppdata 5. Relative sequence abundance of most abundant plant orders. Error bars show standard
 error of the mean.

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Alluvial (Gogerddan) soil: A subset of the storage treatments (T1, T3, T6, T8, T9) were applied to a contrasting soil type from an arable field in Gogerddan. The organic matter content of this soil was much lower (3.6% vs 7.3%) and initial plant and fungal populations of the original soils were very different. For example, Ascomycota fungi comprised ca. 70% of the initial fungal population at Gogerddan (vs 37% at Brignant), mostly due to the much lower abundance of CHEGD fungi (16% vs 41%, with Hygrophoraceae absent).

319

As with the Brignant soils, freeze-thaw storage (T3; freeze-thaw followed by 5 d at 23°C) resulted in the greatest difference in fungal populations relative to control (Fig. 5A), with a reduction in diversity (Suppdata 6A,6B) compared to control and the other treatments. The pattern of divergence of plant DNA composition followed a similar pattern to the fungi (Fig. 5B) but with no significant decrease in diversity indices (Suppdata 6C,6D). The most divergent treatment was storage at 23°C for 14 d (T9) which, in contrast with the other findings, showed higher diversity compared to the control but not the other treatments (Suppdata 6D).

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Fig. 5. Principal coordinate diagrams of the fungal community data (A) and plant community data (B)
 in Gogerddan soil, highlighting the difference in community between the different soil storage
 treatments. Points show the mean axis scores and error bars indicate standard error of the mean.
 The control (immediate freezing at -80^o₂C) is indicated with black arrow.

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SuppData 6. Variations in diversity indices by storage treatment for the alluvial (Gogerddan) soil A)
 Fungi Simpson diversity index B) Fungi Shannon diversity index. C) Plant Simpson diversity index D)
 Plant Shannon diversity index. Letters above the bars indicate significant groupings as determined by
 Tukey's HSD post hoc test and error bars show standard error of the mean. NS indicates no
 significant treatment effect. Note that Shannon and Simpson indices are scaled inversely (i.e. higher
 index = lower diversity).

339

340 In contrast to the Brignant soil, the relative abundance of Ascomycota: Basidiomycota did not 341 increase following freeze-thaw storage (Suppdata 7), in large part because the dominant ascomycete 342 at Gogerddan (Chaetothyriales sp:SH1512803.08FU accounting for 21% of all fungal sequences in 343 T1) was 4-fold lower in T3 and several basidiomycetous soil yeasts (e.g. Solicoccozyma spp.) 344 increased several fold in abundance. In both the Gogerddan and Brignant soils, mycorrhizal fungi 345 (AMF) declined in abundance following freeze-thaw treatment and fungi categorised as saprotrophic 346 in FUNguild exhibited a 2-fold increase relative to control. Metarhizium carneum and 347 Mortierellomycota (mainly Mortierella elongata) both increased 2-fold in abundance following 348 freeze-thaw storage, as was found with the Brignant soil.

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SuppData 7. Relative abundance of fungal groups by storage treatment for the alluvial soil
 (Gogerddan) A) Ascomycota to Basidiomycota ratio; B) *Metarhizium carneum*; C) Mortierellomycota;
 D) Saprotrophic fungi; E) Glomeromycotina (Arbuscular mycorrhizal fungi); F) Grassland fungi
 (CHEGD). Letters above the bars indicate significant groupings as determined by Tukey's HSD post
 hoc test and error bars show standard error of the mean.

355

The plant community was also less diverse in the alluvial soil, with *Ranunculus bulbosus* (36.9%), the chlorophyte *Coelastrella sp.* (22.4%), *Holcus lanatus* (7.4%) and *Polygonum aviculare* (5.7%) found as

358 the dominant plant species. Plant sequences comprised 14%-42% of all the sequences retrieved 359 suggesting a similar ratio of plant to fungal biomass to that found in the Brignant soil, although with 360 a greater abundance of Chlorophyta. The dominant chlorophyte, Coelastrella sp, is a ubiquitous 361 species found in many substrates, including soils, worldwide (Wang et al., 2019). As with the 362 Brignant soil, plant community data increased in variation with treatment especially after freeze 363 thaw (T3) and longer storage at 4°C (T8) and room temperature (T9) (Fig. 5). As also observed for the 364 Brignant soil, all treatments showed greater variability in plant community than the control (Fig. 5B-365 error bars), suggesting that DNA degradation was occurring and that this was due to the treatment 366 effects rather than any initial inter-replicate variability. Seed germination was observed in some 367 samples stored at room temperature (T9) and this may have contributed to this variability.

368

369 Discussion

370 In this investigation we have tested the effectiveness of different soil storage conditions in stabilising 371 fungal and plant DNA prior to later storage (-80°C) and DNA extraction. This is a concern for soil 372 ecologists, since transport from remote and field sites to research laboratories requires interim 373 storage in transit. This may also be a concern where soil sampling is undertaken by third parties and 374 require transport by mail or courier. For example, the authors recently studied the soils of endemic 375 woodlands in St. Helena and transport of samples to Wales involved storage of the samples for up to 376 8 d at 4°C in sealed plastic bags (Detheridge et al., 2020). On other occasions we receive samples 377 from third party collaborators who may send soil samples in batches with delays of up to a week 378 between collection and final frozen storage.

379

The data presented here shows clearly that refrigerated storage for up to 14 d (T8) prior to frozen storage at -80°C has little effect on the fungal or plant DNA later extracted. In contrast, samples initially frozen but allowed to thaw show the most rapid deterioration, presumably due to initial icedamage from freezing and subsequent enzymatic degradation of DNA.

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385 Air-drying (sometimes with the aid of silica gel) is widely used in botanical fieldwork for preservation 386 of plant tissues (Chase and Hills, 1991; Liston et al., 1990) and has been shown to be superior to other methods for many (Pyle and Adams, 1989) but not all (Thomson, 2002) species. For the 387 388 preservation of plant DNA in soil, this method was also highly effective, more so with passive, warm 389 (T7) than active ambient air-drying (T5) (Fig. 1B). However, warm (37°C) air-drying caused large 390 changes to fungal communities, presumably due to degradation of the DNA of certain fungi, notably 391 CHEGD fungi (Fig. 4F). Relative to other groups of biota, fungi are heat-sensitive and only a few 392 species can grow at 37°C (Robert et al., 2015) and no such effect was apparent with ambient air-393 drying. Mean relative abundance of saprotrophic fungi was greater for both air-dry treatments than 394 in the control (Fig. 4D) but there was no significant treatment effect.

395 396 A few fungi were observed to increase in abundance under some storage treatments, presumably 397 because the storage conditions were conducive to their growth. Metarhizium (formerly 398 Paecilomyces) carneum (Kepler et al., 2014) showing the greatest increase in abundance, notably in 399 the freeze-thawed soils which were then incubated at ambient temperature (Fig. 4C). This species is 400 strongly chitinolytic and was frequently recovered from soil baited with chitin (Gray and Baxby, 401 1968; Jackson, 1965). Another closely related species, Metarhizium marguandii (Inglis and Tigano, 402 2006), showed the same patterns of relative abundance but was present at lower levels. Both these 403 species are entomopathogenic to Lepidoptera (Bakeri et al., 2009; Magda and Said, 2014; Shin et al., 404 2013). However, in this experiment the soils were sieved did not have a high content of soil fauna 405 and it is likely that proliferation of *M. carneum* was due to its ability to degrade the cell walls of 406 recently dead fungi (e.g CHEGD fungi and AMF). 407

408 Of the 15 Mortierella spp. detected within the Brignant soil, all but one increased in abundance in 409 the freeze-thawed soil incubated at 4°C (Fig. 4C). Mortierella spp. are psychro-tolerant (Melo et al., 410 2014; Widden, 1987), exhibit ice nucleation activity (Fröhlich-Nowoisky et al., 2015) and are abundant in recently thawed glacier forefront soils (Dresch et al., 2019). As has been found for 411 412 Metarhizium spp., Mortierella spp. are also chitinolytic and frequently isolated in soil baiting 413 experiments with chitin (Gray and Baxby, 1968; Jackson, 1965). Mortierella alpina, the most 414 abundant Mortierella species found here is also reported to be parasitic on soil fungi (Rhizoctonia 415 spp.) and nematodes (Al-Shammari et al., 2013), as well as occurring as endophytes of plant roots 416 (Bonfante, 2020). Thus, it is likely that, like M. carneum, the Mortierella spp. benefit from the 417 increased abundance dead hyphae (AMF/ CHEGD etc.) and are able to exploit these at low 418 temperature. Mortierellomycotina (all Mortierella spp.) were also elevated 4-fold in soil stored at 419 4°C for 14 d (T8), the treatment least changed from the control. Thus the presence of elevated 420 populations of these fungi provides a useful indication that soils have potentially been stored sub-421 optimally. Also highly elevated (10-fold) in freeze-thawed soils was Myxotrixchum (anamorph is 422 Oidiodendron); members of this genus are also most commonly encountered in boreal soils rich in 423 organic matter (Rice and Currah, 2005).

424

425 Of the taxa which declined substantially following freeze-thawing, Glomeromycotina (AMF) showed

the greatest decline. However, within this subphylum some taxa were more heavily affected than

427 others. For instance, *Acaulospora* sp. showed a >4-fold decline (treatments T2 and T4), whereas

428 *Claroideoglomus* spp. declined less than 2-fold. This is consistent with the findings of Klironomos et 429 al. (2001) who found *Claroideoglomus* to be tolerant of freeze-thaw cycles compared to other AMF

430 spp.

431

432 The CHEGD fungi (barring Entolomataceae) also showed substantial decline in relative abundance in 433 freeze-thaw treatments. Like AMF, these fungi are obligate root-associated biotrophs (Halbwachs et 434 al., 2013; Halbwachs et al., 2018), and are negatively affected by killing of host vegetation (Griffith et 435 al., 2014). the fact that Entolomataceae were differently affected to other CHEGD fungi suggests that 436 they are nutritionally more flexible, potentially with some saprotrophic ability. Together the CHEGD 437 fungi comprised >40% of the total fungal biomass at the Brignant site and are recognized to be the 438 dominant fungi of undisturbed mesotrophic grasslands (Griffith et al., 2019; Halbwachs et al., 2013). 439 Their susceptibility to freeze-thaw treatment and resultant increase in fungal necromass is likely the 440 cause of the large proliferation of the chitinolytic M. carneum and Mortierella spp. in freeze-thaw 441 treatments.

442

443 Conclusions

444 To our knowledge this is the only study to have examined the effects of sub-optimal soil storage on 445 eukaryotic eDNA using a high resolution method. When analysing fungal communities, for those 446 situations where freezing samples and freeze drying are impractical, such as remote locations 447 without equipment and requiring length shipping times, the analysis indicates that the best options 448 available are to ship cold or, if impractical, to air dry at room temperature prior to shipping. Air 449 drying can be enhanced by using an unheated active air source, such as a blower or a fan. Pre-450 freezing a sample prior to shipping is not recommended, nor is drying with a heat source such as a 451 drying oven. For two contrasting soil types, the results of suboptimal storage were similar, 452 suggesting broad applicability of these guidelines.

453

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460

461 <u>Author's contribution</u>

462 GWG, JS, APD conceived the study. Experiments were undertaken by LAC and APD.

- 463 Manuscript was drafted by GWG/APD and all authors contributed to editing of the 464 manuscript.
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629

Table 1. Storage treatments tested in this study using soil from the Brignant longterm experiment. An * indicates treatments also tested with soil from the Gogerddan alluvial plain.

Treatment	Initial Processing	Duration	Secondary Processing	Duration	Final Processing
T1*	Freeze -80°C				Freeze Dry
T2	Freeze -20°C	Overnight	Thaw. Cold room 4°C closed bag	14 days	Freeze -80°C & Freeze Dry
T3*	Freeze -20°C	Overnight	Thaw. Cold room 4°C closed bag	5 days	Freeze -80°C & Freeze Dry
T4	Freeze -20°C	Overnight	Thaw. RT 23°C closed bag	14 days	Freeze -80°C & Freeze Dry
T5	Cold room 4°C closed bag	3 days	RT 23°C active air dry	5 days	Freeze -80°C & Freeze Dry
T6*	Cold room 4°C closed bag	3 days	RT 23°C open bag	5 days	Freeze -80°C & Freeze Dry
Т7	Cold room 4°C closed bag	3 days	Warm dry 37°C open bag	5 days	Freeze -80°C & Freeze Dry
T8*	Cold room 4°C closed bag	14 days			Freeze -80°C & Freeze Dry
T9*	RT 23°C closed bag	14 days			Freeze -80°C & Freeze Dry
T10	RT 23°C closed bag	5 days			Freeze -80°C & Freeze Dry



PCO 1 25.82% of total variation

Fig. 1. Principal coordinate diagrams of the fungal community data (A) and plant community data (B) in Brignant soil, highlighting the difference in community between the different soil storage treatments. Points show the mean axis scores and error bars indicate standard error of the mean. The control (immediate freezing at -80°C) is indicated with black arrow.



Fig. 2. Levels of t-statistic from pairwise Permanova of the control treatment compared to all other treatments (Brignant soil). The p value is shown above the bar with significant (P<0.05) values shown in red. (A) fungal community data (B) plant community data.



Fig. 3. Relative sequence abundance of Fungi to plants by treatment (Brignant soil). Letters on the bars indicate significant groupings as determined by Tukey's HSD post hoc test and error bars show standard error of the mean.



Fig. 4. Variations in relative abundance of key fungal groups by storage treatment in Brignant soil. A) Ratio of Ascomycota to Basidiomycota; B) *Metarhizium carneum*; C) Mortierellomycota; D) Saprotrophic fungi; E) Glomeromycotina (Arbuscular mycorrhizal fungi); F) Grassland fungi (CHEGD). Letters above the bars indicate significant groupings as determined by Tukey's HSD post hoc test and error bars show standard error of the mean.



PCO 1 47.89% of total variation

Fig. 5. Principal coordinate diagrams of the fungal community data (A) and plant community data (B) in **Gogerddan** soil, highlighting the difference in community between the different soil storage treatments. Points show the mean axis scores and error bars indicate standard error of the mean. The control (immediate freezing at -80°C) is indicated with black arrow.