Re-assessing the vertical distribution of testate amoeba communities in surface peats: Implications for palaeohydrological studies

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Abstract

Testate amoeba-derived transfer functions are frequently used in peatland palaeohydrological studies and involve the development of training sets from surficial peats. However, within acrotelmic peats, considerable vertical variation in assemblage composition can occur, particularly along Sphagnum stems, which may limit the representation of the associated ‘contemporary’ testate amoeba samples as analogues for the peatland surface. This paper presents contiguous testate amoeba assemblage data from nine monoliths collected from different peatland microforms (hummock, hollow, lawn) in three Sphagnum dominated ombrotrophic peatlands in Ontario and Quebec, eastern Canada. The aim is to: (i) gain a greater understanding of the vertical distribution of xerophilous/hygrophilous taxa along Sphagnum stems; (ii) determine the vertical extent of live/encysted taxa along this gradient; and (iii) assess the significance of this distribution on surface sampling protocols. The results show that testate amoeba communities in the uppermost acrotelmic peat layers display considerable variability. This may reflect a complex interplay of abiotic and biotic controls, including moisture, temperature, light and other characteristics, food availability, and mineral particle availability for test construction. These findings underline the complexity of testate amoeba community structure and highlight the importance of analysing both living and dead Sphagnum stem sections when developing calibration sets.

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Introduction

Testate amoebae have been widely used as proxies for hydrological change and for inferring palaeoenvironmental conditions in peatlands (e.g. Hendon et al. 2001; Lamentowicz et al. 2015; Loisel and Garneau 2010; Payne et al. 2011; Swindles et al. 2009). Surface wetness reconstructions are commonly inferred from transfer functions, which aim to model the relationship between contemporary testate amoeba assemblages and depth to water table (Charman et al. 2007; Hendon and Charman 2004; Woodland et al. 1998). These are in turn applied to down core assemblages in contemporary peatland sites. However, it has long been known that considerable variability exists within testate amoeba communities in surficial peat layers (Buttler et al. 1996; Heal 1962; Mitchell et al. 2008). The ecology of testate amoebae inhabiting Sphagnum stems and in surficial peats has been shown to be controlled by a number of physi-
cal parameters, for example, moisture, light, food resources, temperature and the availability of mineral material for test construction (Booth 2002; Charman et al. 2000; Heal 1962; Meisterfeld 1977; Mieczan 2010). Below the surface, the acrotelm represents the aerobic uppermost layer of peatlands and varies in depth from between 5–50 cm, whereas the underlying catotelm has a lower rate of residual peat decay (Belyea and Clymo 2001; Clymo 1984). The vertical depth of the acrotelm is dependent upon surface microtopography (e.g. hummock, hollow and lawn microforms), the interaction between water tables and processes of peat accumulation (Belyea and Clymo 2001; Quinty and Rochefort 2003; Rydin and Jeglum 2006). However, the concept of two distinctive layers is arguably too simplistic, as the acrotelm contains anaerobic sections and aerobic channels that can extend into the catotelm via vascular plant roots (Hayward and Clymo 1983; Morris et al. 2011). Notwithstanding this, the boundary between the acrotelm and catotelm has been considered to occur at the minimum water table in summer (Clymo 1984). Other criteria that have been used to distinguish the boundary include changes in humification (Charman et al. 1999) and bulk density of the peat (Yu et al. 2003). The acrotelm layer also shows considerable internal variation. The organic material here may be fibrous or pseudofibrous (with plant remains such as stems, rhizomes and root matter recognisable), or may be lost but the peat may still retain integral structure. As mosses and other plant communities die, fibrous material is added to the peat, providing physical structure for the new upwards growth of mosses (Malmer et al. 1994). Decomposing litter content also varies within the oxic, acrotelmic peat layers, and changes with depth, eventually becoming covered in the catotelm by rising water tables (Rydin et al. 2006).

In order to quantify hydrological changes within peatlands through transfer function models, modern analogues of testate amoeba assemblages must be sampled from these surficial peats. A number of methodological approaches have been employed to achieve this. In many European studies, for example, the uppermost 1 cm of surficial peat has been used to process testate amoeba samples after the removal of the Sphagnum capillifolium (e.g. Swindles et al. 2009; Woodland et al. 1998). This narrow vertical interval has been selected to ensure that a sample representing only the very recent period is obtained. A slightly modified approach was applied by Swindles et al. (2015) in which the entirety of the green (living) moss fraction and 1 cm of the underlying brown Sphagnum were sampled. In other studies, assemblages have been characterised by identifying an ‘upper’ surficial peat section, which comprised living, vertical Sphagnum stems, and a ‘lower’ section, comprising brown, humified Sphagnum with collapsed stems (e.g. Booth 2002; Schönborn 1963). Such studies have aimed to explore vertical variations in assemblage composition in regions where there is considerable thickness of living Sphagnum present. The study of Booth (2002), for example, which was undertaken in Michigan, USA, involved sampling the entire green Sphagnum layer and a ~10 cm$^3$ lower sample from immediately below the green stems. In this case only the lower samples were included in the training set used for calibration model development (Booth 2002).

In spite of the large number of studies that have examined surficial peat samples, the vertical distribution of living testate amoeba communities in surficial Sphagnum stems remains largely unexplored, but is critical for understanding species–environment relationships. Environmental gradients (e.g. moisture) have been shown to control the vertical variation of ‘live’ and ‘dead’ assemblages, which can be distinguished through the staining of living protoplasm with Rose Bengal (Meisterfeld 1977, 1978; Scott and Medioli 1980; Schönborn 1963). Light appears to be a particularly important factor influencing the distribution of species containing symbiotic zoochlorellae, such as Archecrella flavum (Gilbert and Mitchell 2006; Mazei and Bubnova 2007; Schönborn 1963). Similar symbiotic species include Hyalosphenia papilio and Heleopera sphagni, which live in the chlorophyllous upper few centimetres of peatlands (i.e. living green stems) to enable photosynthesis of algae to occur. Taxa that use xenosomes to build tests such as Diffugia spp., Trigonopyxis arcula and Centropyxis spp. have been observed in lower stem positions (Payne and Pates 2009) than idiosomic taxa such as Euglypha spp. (Heal 1962; Schönborn 1962).

Testate amoebae also have the ability to encyst for long periods of time (Foissner 1987; Ogden and Hedley 1980). However, there is a limited understanding of the vertical distribution of encysted testate amoebae and the potential factors controlling this state, as encysted species have typically not been distinguished from live or dead individuals in most modern (i.e. surficial peat) and palaeoecological studies (Jassey et al. 2011; Vincke et al. 2004). Encystment is nevertheless considered to be a response to unfavourable environmental conditions such as desiccation (Booth and Zygmunt, 2005; Lousier 1974; Woodland et al. 1998), frost (Mitchell et al. 2000; Warner et al. 2007) or decreases in food availability (Laminger and Sturm 1984). One challenge in understanding how testate amoeba communities respond to such unfavourable conditions relates to a lack of seasonal monitoring of species–environment relationships. A notable exception was a study by Sullivan and Booth (2011) who employed data loggers to measure relative humidity within the upper few centimetres of Sphagnum stems in 11 peatlands in Pennsylvania and Wisconsin, USA. The results were used to determine the influence of short-term environmental variability on testate amoeba communities. This study has highlighted the role of surface vegetation, as well as changes in precipitation and evapotranspiration in controlling subannual variation in testate amoeba community composition.

Many other studies have examined testate amoeba assemblages from short (<50 cm) cores collected from acrotelmic peats to reconstruct recent hydrological change. Testate amoeba-inferred water table fluctuations from dated cores
of acrotelmic peats, for example, have been compared to instrumental records of climate change to validate calibration models (Charman et al. 2004, 2009; Charman 2007). There is, however, a poor understanding of the relationship between live/dead and encysted testate amoebae in deeper acrotelmic peats and the significance that this might have for interpreting model-inferred hydrological changes. There is also a need to examine the vertical variation in testate amoeba assemblages in regions with deep acrotelms, for example, in dry, continental areas where the living parts of mosses can span a large vertical range. In this paper we examine the live/dead assemblage composition of different microforms (hummock, hollow, lawn) from three ombrotrophic peatlands in continental eastern Canada to make new inferences about the ecology and distribution of testate amoeba taxa in surficial peat environments. The three study sites in Ontario and Quebec are all characterised by high mean summer temperatures (July averages range from 26.1 to 26.5°C) (Supplementary Table 1A), and deep acrotelmic peats. Floristically, they are dominated by a similar range of Sphagnum species and vascular plants. As well as providing a new assessment of the ecology of testate amoebae communities in surficial peatlands, the study provides new insights into the selection of sampling ranges in these environments for quantitative palaeohydrological reconstructions.

Material and Methods

Site description

The three study peatlands, Mer Bleue Bog, Mirabel Bog and Alfred Bog, are all ombrotrophic peatlands in the central St Lawrence Lowlands region of eastern Canada (Fig. 1). This region is characterised by thick acrotelmic peat deposits that typically extend up to 50 cm below the surface, which is ideal for sampling along a vertical hydrological gradient within hummock, hollow and lawn topographies. Mer Bleue Bog (45°24′N; 75°31′W), located ca. 15 km east of Ottawa, covers 28 km² and is characterised by three drainage ‘arms’ separated by alluvial sand ridges (Elliott et al. 2011; Mott and Camfield 1969). The northern arm is domed (Joyal 1970), with peat depths of 5–6 m, decreasing to 1–2 m around the margins. Surface vegetation is dominated by Sphagnum species (e.g. S. capillifolium, S. fuscum, S. magellanicum and S. angustifolium) and ericaceous shrubs such as Ledum groenlandicum, Chamaedaphne calyculata and Kalmia angustifolia.

Mirabel Bog (45°41′N; 74°02′W), formerly known as Saint-Canut Bog (MacPherson 1967; Muller et al. 2003), is a 2.15 km² ombrotrophic peatland situated ca. 45 km northwest of Montreal (Fig. 1). The central part of the bog comprises a well-defined hummock and hollow topography dominated by S. capillifolium and S. magellanicum. Other vegetation includes C. calyculata, Polytrichum strictum, Eriophorum spissum, K. angustifolia, Kalmia polifolia and Viburnum cassinoides. The peatland also supports stunted forms of Picea mariana, Betula populifolia and Larix laricina. Alfred Bog (45°29′N; 74°48′W) is situated 75 km east of Ottawa. It is a 42 km² peatland, the largest in the Southern Ontario Canadian Shield. Dominant moss species include S. capillifolium, S. fuscum, S. magellanicum, Polytrichum commune and Pohlia nutans. Ericaceous shrubs dominate the open peatland and include C. calyculata, K. polifolia, K. angustifolia, L. groenlandicum and Vaccinium oxycoccus.

Sampling methods

To investigate the vertical distribution of testate amoebae in the uppermost sections of peat, nine monoliths (10 × 10 × 20 cm) were collected from the three study sites, Mer Bleue, Mirabel and Alfred Bogs representing a range of hummock, hollow and lawn microsites (Fig. 2). The rationale behind sampling over a wide hydrological gradient was to gain insights into species–environment relationships and is similar to previous investigations that have examined the surficial distribution of testate amoebae in mires (Jassey et al. 2011; Lamarre et al. 2013; Lamentowicz et al. 2008). The sampling extent of the monoliths varied (between ca. 15–20 cm) and was dependant on the depth of the living (green) Sphagnum stems at each sampling point. Each sampling site covered a range of vegetation species and water table depths. Vegetation was identified in the field and the laboratory using the keys of Crum and Anderson (1981), Daniels and Eddy (1990) and Bastien and Garneau (1997). To measure water table depths, ‘single-shot’ readings were taken from the monolith sampling stations using wooden rods. Holes were excavated using a soil auger and sharp knife. The rods were left in the holes for 20 min to allow the readings to equilibrate. The use of single-shot water table measurements for testate amoeba-based hydrological studies has been debated previously (e.g. Bobrov et al. 1999; Booth 2008; Swindles et al. 2015). Whilst the method only provides a
Fig. 2. Schematic diagram showing hummock, lawn and hollow microforms. The transition layer between the live and more decomposed plant stems is indicated with arrows. (For interpretation of the references to colour in the text, the reader is referred to the web version of this article.)

‘snapshot’ of the seasonal water table variability, it has been shown to adequately represent the relative hydrological status of sample locations and is sufficient to drive a hydrological gradient for transfer function development (Swindles et al. 2015). Sampling should not be carried out, however, during periods of extreme weather such as summer drought or after heavy rain (Charman et al. 2007; Swindles et al. 2009). To this end, the monoliths were all collected on dry days during the month of May at the same time of day (before noon). Sampling was undertaken after a period of approximately two weeks when no rain had fallen in the region. After collection, the monoliths were carefully sealed in polythene bags prior to transportation.

Laboratory analyses

In the laboratory, each monolith was split in half vertically to expose the structure and vegetation composition (cf. Buttler et al. 1996). Preliminary results showed that there was a transition between the live and more decomposed stems, which further justified the need to sample at high resolution. Each half was then cut at contiguous 1 cm intervals to analyse the vertical variation of testate amoebae (cf. Meisterfeld 1978; Niinemets et al. 2011). Sub-samples from one half were used to determine moisture content, bulk density and colorimetric peat humification, while those from the other half were stained using Rose Bengal and used for testate amoeba analysis. Samples were oven dried at 105 °C for 24 h and re-weighed to determine moisture content (Chambers et al. 2011). A known volume of peat was also measured at 1 cm intervals and oven dried at 105 °C for 24 h, then weighed to determine bulk density (Tolonen et al. 1992). Rose Bengal stain was added ca. four hours after monolith collection to stain the living protoplasm (Bernhard 2000; Scott and Medioli 1980). Samples were prepared following a modified version of the method by Hendon and Charman (1997) and Charman et al. (2000). Instead of boiling, the samples
were heated gently for 10 min as the ethanol would denature the living tests (cf. Booth et al. 2010). Testate amoebae were identified largely with reference to the key of Charman et al. (2000), although a number of other keys, plates and descriptions were consulted, including Penard (1902), Loeblich and Tappan (1961), Corbet (1973), Ogden and Hedley (1980) and Booth (2002). Subtle differences in morphology were noted during routine counting, for example, the number of pores in specimens of *Hyalosphenia papilio* (Booth and Meyers, 2010). Since the study was undertaken, phylogenetic work has led to some re-classifications (for example, several taxa within the *Nebela tincta-bohemica-collaris* group have been re-classified; Kosakyan et al. 2013). In this paper, however, we adhere to the terms *Nebela tincta* and *Nebela collaris sensu stricto* (cf. Charman et al., 2000).

The number and type of taxa that were encysted from the monoliths was also recorded. Encystment was identified on the basis of a closed aperture and/or with a mass of protoplasm (cf. Corbett 1973; Heal 1962). More specifically, encysted forms were recognised for having an operculum consisting of a plug of organic material surrounding the inner aperture of the test wall and a thickened anterior cyst wall. This has been recognised in forms of *Cryptodifflugia oviformis* and *Cryptodifflugia leachi* (Nicholls 2006). As in previous studies, encysted taxa were summed with the total live count because individuals normally represent <2% of the sample (Beyens et al. 2009; Vincke et al. 2004).

In addition to plotting the live and dead testate amoeba assemblages with depth for each of the nine monolith profiles, Detrended Correspondence Analysis (DCA) was carried out to explore the inter- and intra-microform characteristics of the testate amoeba communities and to examine associations between species. As the main emphasis in undertaking this analysis was to better understand the distribution of living testate amoeba species, only samples that yielded >5% live specimens for at least one taxon were included in the analysis.

**Results and Discussion**

In general, the zone where live testate amoebae were present varied according to microform (hummock, hollow, lawn) in the three study sites and with dominant *Sphagnum* type (e.g. *S. fuscum*, *S. capillifolium*, *S. magellanicum* and *S. angustifolium*). The relationship between live and encysted taxa in each of the nine monoliths is shown in Figs 3–5. The DCA plot (Fig. 6), shows that the testate amoeba assemblages were well clustered for the hummock samples but there was some overlap between the hollow and lawn samples. Axis one in the plot appears to be strongly influenced by depth to water table, with the species that are common indicators of dry conditions (e.g. *Assulina seminulum*, *Trigonopyxis arcula*) plotting on the left of the figure and established wet indicator species (e.g. *Archerella flavum*, *Hyalosphenia elegans*) in the centre and right (Fig. 6). The overlap and apparent reverse ordering of the lawn and hollow samples may reflect the similarity in water table measurements between these microforms (Supplementary Table 1B). It may also suggest that other ecological gradients e.g. light and temperature had an influence on assemblage composition. A summary of the vertical distribution of the total live taxa in the hummock, hollow and lawn microsites is given in Fig. 7, whilst Fig. 8 provides a simplified summary of the relationship between live testate amoeba species, water table depths and microform vegetation. Emphasis is placed in the following sections on examining the assemblages from the upper and middle sections of the monolith profiles for each microform, i.e. from the green (=living) or ‘transitional’ zone samples that yielded living testate amoebae (Figs 3–5). The deeper, semi-humified to humified parts of the profiles that primarily yielded dead specimens may represent different environmental conditions to those that prevailed at the time of sampling, potentially spanning several decades. As we have no chronological control for the profiles (such as that afforded by 210Pb dating; Appleby 2001), and no means of determining the rates at which the surficial, green *Sphagnum* layer decomposes and becomes incorporated to the humified peat, only brief reference is made to the lower assemblages.

**Hummock microforms**

Counts above 150 specimens were difficult to achieve in the hummock profiles, particularly when water table depths exceeded 50 cm below the peat surface (Fig. 3B and C). Xerophilous communities dominated by *Assulina muscorum* and *A. seminulum* were typical of the *S. fuscum* and *S. capillifolium* hummock microforms (Fig. 3). Other taxa that favour dry conditions included *Hyalosphenia subflava*, *Trigonopyxis arcula* and *Cyclopyxis arcelloides*. Xenosomic species increased in abundance with depth and photophilic species were found in the upper aerobic *Sphagnum* that contain symbiotic algae (zoochlorellae). For example, *Phryganella acropodia* was only present in the lower depths of the *Sphagnum* profile (Fig. 3C). This could be due to the increased availability of material for test construction (e.g. minerals and diatoms) (cf. Booth 2002; Mitchell and Gilbert 2004). As depth increased, the apertures of *P. acropodia* became increasingly armoured with mineral particles, making it difficult to distinguish them from small forms of *Cyclopyxis arcelloides*.

Interestingly, in the hummock profiles (Fig. 3), there is evidence for a zonation in the assemblages between the upper 5 cm of the green *Sphagnum* stems, and the underlying ‘transitional’ layer where the stems typically were decomposing and where colour changes (notably an increased dominance of red-brown stems) were observed (Figs 2, 3). For example, in the hummock profile in Fig. 3A, *A. muscorum* reached peak abundances of 60–80% in the uppermost 3–4 cm (green layer), in contrast to *H. elegans* which attained peak frequencies (10–30%) between 5–9 cm (transitional layer). In a study of the vertical micro-zonation of testate amoebae from re-
colonizing Sphagnum peatlands in the Swiss Jura Mountains, Mitchell and Gilbert (2004) suggest that competitive exclusion may occur between species that are closely related. They note that A. flavum, A. muscorum and H. papilio reach higher abundances in the capitulum compared with Amphi trema wrightianum, A. seminulum and H. elegans. Patterns of species micro-distribution may also be explained by specific environmental constraints. For example, the observed changes could be a response to the increased sensitivity of taxa to desiccation (e.g. A. muscorum). A decrease in moisture at the capitulum can lower pH and ion concentrations relative to the Sphagnum stem (cf. Mitchell et al. 2000). Some taxa are known to have a preference for low pH and moisture environments, including A. muscorum and A. flavum (cf. Lamentowicz and Mitchell 2005; Swindles and Roe 2007), whilst A. seminulum can reproduce more successfully at greater depths where there is a thicker water film along the Sphagnum branches (Mitchell et al. 2008). It is interesting to note that the sub-division noted above between the assemblages from the upper stems (above 4 cm) and from a few centimetres below is not particularly apparent in the DCA plot; the samples from different hummock depths are well mixed. This suggests that in comparison to the variation in the dataset as a whole, the assemblages are not that different.

Fig. 3. Hummock profiles: A (Sphagnum capilifolium) monolith. B (Sphagnum fuscum) monolith. C (Sphagnum fuscum) monolith. Abundance of species >2% shown. Red bars indicate % live with only live >2% shown. Water table depth measurements taken at the time of sampling are indicated. The upper ‘green’ profile sections were dominated by stems in vertical growth position; the lower ‘brown’ monolith sections typically showed increasing humification with depth. For a fuller description of these zones and the intermediate transitional layer see Fig. 2. Taxonomic note: some N. American testate amoeba studies have not identified Hyalosphenia ovalis because of confusion with N. tintca and H. papilio (see Booth and Meyers, 2010). We used the criteria detailed in Charman et al. (2000). (For interpretation of the references to colour in this figure legend, the reader is referred to the web version of this article.)
The vertical distribution of living testate amoebae (>2%) in the hummock microforms spanned a maximum of 12 cm, and comprised up to 55% of the total assemblage. Assulina muscorum dominated the live specimens up to a depth of 10 cm (Fig. 3). Other live taxa included A. seminulum, H. elegans, Heleopera sylvatica, H. subflava, Nebela militaris and Trinema lineare (Figs 3A and B, 8). In a study in the Middle Volga region (Russia), Mazei and Tsyganov (2007) recorded maximum live abundances of testate amoebae in dry habitats, such as hummocks of between 22–27%. Robroek et al. (2007) and Sullivan and Booth (2011) suggest that despite the increasing distance of the hummock surface from the water table, hummocks can contain a greater seasonal storage of moisture due to their compact structure. The common hummock-forming species Sphagnum fuscum, for example, has the ability to retain water more effectively than some lawn and hollow-forming Sphagnum species (e.g. S. rubellum, S. magellanica) because of its high capitula density and efficient capillary system (Robroek et al. 2007; McCarter and Price 2012). As a result, this would reduce the impacts of short-term environmental variability (e.g. desiccation) on these microforms in comparison to hollows or lawns, and may explain why a high abundance of live taxa were found in the uppermost 5–10 cm of the hummocks.

**Lawn microforms**

Counts between 70–100 specimens were achieved in the upper 0–3 cm of the lawn profiles, rising to >150 below this (Fig. 4). There was a co-dominance of xerophilous taxa such as A. muscorum and H. subflava, as well as hygrophilous taxa such as Heleopera petricola and H. elegans. Within the deeper, unhumified (green) parts of the lawn profiles the testate amoeba communities become increasingly uniform in composition (Fig. 4). This could be due to a decrease in light and/or temperature (Mitchell and Gilbert 2004; Jassey et al. 2011). Heleopera sphagni and H. papilio, for example, both attain highest abundances in the uppermost 5 cm of the S.
encysted specimens then decline rapidly below this (Fig. 4A and B). Both taxa have symbiotic zoochlorellae and require light, therefore their maximum abundance would be at the top of the Sphagnum stem (cf. Booth 2002). Cyclopyxis arcelloides, however, appears more frequently below 8 cm in the transitional layer where the stems are starting to decompose (Fig. 2). This may reflect the enhanced availability of mineral particles for test construction or increased availability of food sources.

In the more humified samples below the transitional layer the assemblages show further changes, most notably increases in Nebela minor and Nebela penardiana (Fig. 4). A centimetre of peat in these deeper parts of the profiles is likely to represent a longer time interval than the same thickness of sample in the green, stem-dominated horizons because of the influence of decompositional processes and the compression of the stems that occurs with decay. Without chronological control it is not possible to ascertain, however, what time interval the lower assemblages represent, or what relationship, if any, they show with the environmental conditions that prevailed at the time of sampling.

Co-dominance of live wet and dry indicator taxa was most notable in the lawn profiles. This may reflect the fact that the living Sphagnum stems in the lawn habitats are characterised by intermediate moisture characteristics for most of the year and are thus able to satisfy the niche requirements of species with tolerance ranges spanning both ends of the moisture spectrum. As water-table measurements were only taken at the time of sampling it is not possible to assess this further. Sullivan and Booth (2011) note that it is the increase in short term variability that results in the appearance of taxa at opposing ends of the moisture gradient in fossil assemblages from these environments. The presence of mixed wet and dry assemblage constituents in the lawn samples may explain why these samples showed a weaker degree of clustering in the DCA plot than the hummock and hollow samples (Fig. 6). This pattern may also suggest that factors other than hydrology are influencing the assemblages.
Fig. 6. Detrended correspondence analysis (DCA) results showing principal patterns of variation in the testate amoeba assemblages. The analyses were run on the total counts from the unhumified (green living) monolith samples (n = 65) that yielded >5% live specimens for at least one taxon. Clusters are encircled for the three microform environments and core depth information is given.

Fig. 7. Live testate amoeba abundances (>2%) in the hummock, lawn and hollow microforms from the three peatland sites.
The vertical extent of living testate amoebae (≥2%) in the lawn microforms spanned a maximum of 19 cm, and comprised up to 50% of the total assemblage (Fig. 4A). This is the largest depth range encountered during the study (Fig. 7). The majority of the live taxa consisted of *H. petricola* and *H. sphagni*, which occurred within the upper 5 cm. As noted above, the dominance of these species in the green, chlorophyllous *Sphagnum* stems probably reflects near-surface light availability, which would have enabled photosynthesis of algal symbionts. Interestingly, live specimens of *H. elegans*, *N. penardiana* and *N. minor* were also encountered in the more humified sections of the lawn profiles between ca. 10–18 cm, suggesting that light was able to penetrate deeper into the acrotelmic peats in the lawn environments, possibly along vertical channels created by stems or root action. Taphonomic displacement of the extant specimens down profile (for example, as a result of percolating rainwater or short-lived water table fluctuations) is also a possibility, but is less likely given the relatively even distribution of the living specimens of the three noted taxa within the 10–18 cm depth range (Fig. 4A and C).

A number of *Euglypha* taxa recorded in the lawn samples (notably *Euglypha strigosa* and *Euglypha tuberculata*) were recorded as having a narrow/closed aperture and were classified as encysted based upon the descriptions by Heal (1962). These encysted specimens occurred exclusively in the upper living (green stem) sections of the profiles at depths of up to 5 cm below the surface (Fig. 4).

**Hollow microforms**

The testate amoeba assemblages from the hollow profiles were characterised by a range of hygrophilous taxa such as *H. elegans* and *H. papilio*, and some xerophilous taxa, for example, *A. muscorum* and *H. subflava* (Figs 5, 8). Spined forms of *Euglypha* (e.g. *E. strigosa*) were also abundant in the upper 10 cm, which are typical of wet habitats (Mitchell et al. 2008; Payne 2007). The rotifer *Habrotrocha angusticollis* achieved maximum abundance in the upper 1–3 cm in the capitulum of *S. magellanicum* and *S. angustifolium*. The dominant taxon in these environments was *H. elegans*, which also attained peak abundance in the uppermost 3 cm (Fig. 5).
This suggests that its vertical distribution is influenced by light (cf. Mieczan 2009), but its presence also corresponds to a decrease in bulk density through the profile and an increase in moisture content (Fig. 5). Overall, there were more taxa in hollow environments (between 25–29) than in the hummocks and lawns, an observation also noted in Sphagnum dominated peatlands by Mieczan (2010).

An interesting minor constituent of the hollow microsites is Nebela tincta. In profile C from Mirabel Bog, for example, 4% live specimens of N. tincta were recorded at 2.5 cm (Fig. 5C). This species has been described as a dry indicator taxon in a regional study of 31 eastern North American peatlands (Booth 2008), although it has also been reported in ‘very wet’ conditions in peatlands in southwestern Ontario (Warner 1987). The presence of N. tincta, albeit in low abundances in the hollow microsites, may also relate to other controls such as food availability, as the species is known to ingest a wide range of organic remains, diatoms and other testate amoebae (e.g. Euglypha and Assulina spp.) (Booth 2002; Gilbert et al. 2003). Decompositional processes may also have been more active in the hollow habitats relative to the other microforms, producing greater availability of other food sources, e.g., fungi and bacteria.

A further characteristic of the N. tincta record that is worthy of note is the slightly different distribution shown between this taxon and Nebela minor, which has been suggested to be synonymous with N. tincta (sensu stricto) based on recent phylogenetic work (Kosakyan et al. 2013). Nebela minor increases in the lower, more humified parts of the profiles, whilst N. tincta shows a more variable distribution (Fig. 5). When live specimens of the two taxa are present, they do not occur in the same samples (see for example, Figs 5B and C, 4A and C). These findings are interesting, suggesting that the two morphologically distinct forms may be ecophenotypes. Further work is required to assess this in other peatlands.

Overall, the vertical distribution of live testate amoebae (>2%) in hollow microforms spanned a maximum of 12 cm, and comprised up to 90% of the total assemblage (Fig. 4). These values were the highest recorded in the study (Fig. 7). The majority of the live taxa consisted of H. elegans, particularly within the upper 4 cm (Fig. 5). The high percentage of live specimens in the hollow profile from Mirabel Bog (Fig. 5C), might be attributed to the presence of a tamarack tree in proximity to this sampling station, which provided shade. As well as reducing light penetration, this may have resulted in changes in temperature or moisture availability that provided favourable conditions for live testate amoeba, particularly H. elegans. Previous studies have also found that small environmental changes such as shading and light intensity can affect the growth of the Sphagnum capillifolium (Buttler et al. 1996; Mitchell et al. 2000) and Sphagnum stem length (Hayward and Clymo 1983), which may further impact testate amoeba communities. Notwithstanding this, the inferred shaded conditions at this Mirabel Bog sampling station nevertheless permitted the growth of non-compact forms of S. angustifolium. Other live taxa (>2%) present in the hollow microforms included Placocista spinosa, A. muscorum, Euglypha compressa type, E. strigosa type, H. subflava, Nebela militaris, N. minor, N. tincta and Diffugia leidyi (Figs 5, 8).

One species that was notably absent from the live assemblage was H. papilio, a common hygrophilous taxon. This may reflect the fact that the hollow sites selected were not particularly moist at the time of sampling (water tables ranged from 14 to 22 cm below the surface), or, as discussed above, it may reflect other factors, such as the availability of food sources. Interestingly, Jassey et al. (2012) observed that H. papilio has a different feeding strategy to N. tincta (which was present in the live assemblage) based on a study of digestive vacuole content. The former species was found to preferentially ingest ciliates, the latter fungi.

As variability in surface moisture explains the majority of testate amoeba community variance in peatlands, increased moisture typically results in higher species abundance, diversity and decreased encystment (Laminger 1978; Payne 2011). However, in this study, samples from a S. capillifolium-dominated hollow microform (Fig. 5B) yielded 63 encysted forms, which included A. muscorum and A. catinus. Encysted forms of A. muscorum were identified because of a narrowing aperture with a rounded mass of dark protoplasm. The majority of the encysted species of this taxon (44 specimens) occurred in the more humified parts of the profiles between 8–14 cm. As A. muscorum is characteristic of dry conditions, its encystment stage could reflect unfavourable, moist conditions given the proximity of the water table (22 cm). In the case of A. catinus, encystment was based on an infilled and dome-like aperture. It only occurred in a few specimens from the upper portions of the Sphagnum stem, where conditions may have become too dry. A study by Heal (1964) also found that encystment occurred in response to unfavourable environmental conditions, for example, freezing periods which exceeded the biophysical tolerances of certain taxa (Foissner 1987). Whilst encysted forms of other taxa were not found in this study, Heal (1962) noted the encystment of many forms of H. papilio concentrated at the Sphagnum capillifolium in wet habitats. Sullivan and Booth (2011) comment on the importance of understanding the life history characteristics of individual taxa, for example, rate of reproduction and rate of encystment/uncystment. Those species that encyst have a competitive advantage over those that do not in environments with highly variable moisture availability such as peatlands in a continental setting (Booth and Zygmunt, 2005). During periods of favourable conditions, specimens that can encyst may have the potential to exploit food and suitable materials for test construction (Sullivan and Booth 2011).

Conclusions

This study has provided new insights into the vertical distribution of live and encysted testate amoebae in surface Sphagnum-dominated peats, particularly in the green moss fraction and the partly decomposed ‘transitional’ layer.
which occurs between the living stems and the more humified peat below. These micro-environments have hitherto received relatively little attention. The assemblages as a whole (live and dead) were similar to those reported previously in Sphagnum-dominated peatlands, with the greatest abundances of xerophilous taxa occurring in hummock microforms (e.g. *A. muscorum*), and with increased proportions of hygrophilous taxa such as *H. elegans* and *H. papilio* in hollow microsites. The peat profiles were collected in late Spring during a prolonged period (ca. 2 weeks) of dry conditions. The community structure of the extant (live) species is thus likely to reflect seasonal (or sub-seasonal) variations in moisture availability and other physical controls that prevailed in the weeks to months prior to sampling.

The high-resolution, microform-specific data presented shows that the testate amoeba assemblages of the sampled peatlands display considerable vertical variability. This was apparent for all three microform environments investigated (hummock, hollow and lawn), with the most pronounced changes typically occurring at around 4–5 cm in the profiles (Figs 3–5). In the hummock samples, for example, live specimens of *A. muscorum* dominated the upper 4 cm, whilst live *H. elegans* peaked in the transitional layer (5–9 cm). In the lawn microforms, live specimens of *H. petricola*, *H. papilio* and *H. sphagni* attained maximum abundances in the uppermost 5 cm and then declined below this depth. Other taxa such as *Placocista spinosa*, *Euglypha rotunda* type, and several Diffugid taxa also peaked in lower sections of the green *Sphagnum* fraction and the underlying deeper, more humified sections of the hollow profiles. Marked vertical changes in assemblage composition have been noted previously in the upper sections of *Sphagnum*-dominated peat profiles (Gilbert and Mitchell 2006; Mazei and Bubnova 2007; Payne and Bates 2009) and have been explained by changes in light, temperature, moisture and oxygen availability, the availability of minerals for test construction, and the differential availability of food sources. Further work is required to better understand how these complex micro-habitats change with depth. The influence of multiple abiotic controls, as well as the availability of food sources, should be assessed over a full range of seasons. The use of data loggers to obtain continuous measurements throughout the year has considerable benefits in this regard as demonstrated previously (Sullivan and Booth 2011).

The results of the study also have implications for sampling testate amoeba assemblages for the development of calibration (training) sets in areas with deep acrotelmic peats. In particular, the vertical fluctuations noted within the green stem layer, and the lack of strong clustering of samples from specific depths for the three microforms in the DCA (Fig. 6) reinforce the need to sample across a vertical range that includes the full living stem layer to ensure that assemblage variability is captured. Some previous studies have removed the upper 1–2 cm of the green layer, spanning the capitulum, prior to testate amoeba analysis as this uppermost stem section has been found to display stronger assemblage variation in response to light, temperature, oxygen, moisture and mineral material for test construction (Booth 2002; Booth et al. 2010; Heal 1962; Meisterfeld 1977; Mieczan 2010). We found no clear justification for removing this layer; in some cases the uppermost (0–2 cm) samples yielded assemblages that were different to those at 3–4 cm, in other cases the assemblages were similar to those immediately below (Figs 3–6). It would seem more logical to include the capitulum as part of an integrated analysis of the green stem fraction.

The study also highlights the need to better understand the relationship between the assemblages of the green layer and those of the uppermost brown layer, which has been often used in addition to (e.g. Swindles et al. 2015), or in place of (e.g. Booth 2002), the green fraction in calibration (training) sets, as it may provide a more representative, time-averaged signal for hydrological reconstructions than the green layer alone. Our results show that significant assemblage changes can occur between these two layers. The rapid decline and disappearance of *H. sphagni*, *H. petricola* and *H. papilio* between the uppermost stem sections of *S. capillifolium* and the underlying more humified peat in the profiles shown in Fig. 4A and B provides a case in point. Further work is required to more accurately delimit the two layers, to understand the decompositional and taphonomic processes that influence the associated testate amoeba assemblages, and to statistically compare the assemblages to assess how they relate to key environmental variables such as moisture availability.

The occurrence of live testate amoebae up to a depth of 19 cm within the profiles also has implications for peatland-based palaeohydrological studies. This wide vertical range may reflect the presence of vertical channels within the peat, which permitted light and/or oxygen to penetrate. The introduction of significant numbers of live specimens into the deeper, well humified sections of the peat could obscure reconstructions that are based on the total assemblage from these horizons. This underlines the need to stain testate amoeba samples from surficial peat cores to provide a refined understanding of assemblage derivation. Analysis of the controls on encystment would also provide additional insights into testate amoeba ecology in near surface peat deposits. In this study, encysted species were found most commonly in the lawn and hollow profiles. Most notably, encysted forms of the xerophilous taxon *A. muscorum* were more commonly found in hollow samples with higher moisture content than in adjacent samples (Fig. 5B). This preliminary finding confirms earlier observations that encystment is a response to stressed conditions (Foissner 1987; Heal 1962, 1964). Additional sampling could be undertaken during specific seasons to assess how taphonomic processes influence live and encysted testate amoebae inhabiting acrotelmic *Sphagnum* stems and how communities are subsequently integrated into the palaeoecological record.
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Appendix A. Supplementary data

Supplementary data associated with this article can be found, in the online version, at http://dx.doi.org/10.1016/j.ejop.2017.03.006.

References


