Diversity of Testate Amoebae as an Indicator of the Conservation Status of Peatlands in Southwest Europe

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Abstract: Testate amoebae are one of the most studied groups of microorganisms in Sphagnum peatland ecosystems and, therefore, one of the most reliable bioindicators of their ecological status. Peatland ecosystems are supported by a delicate biogeochemical balance that leads to the formation of peat, one of the main sinks of C, as a result of soil–atmosphere interaction, but currently they are one of the most threatened wetland types at their southern distribution limit. In the European continent, where climatic conditions limit peat formation, they have endured significant anthropic pressure for centuries, and the risk of loss of biodiversity linked to these ecosystems is critical. In addition, peatlands are poorly known ecosystems in the Iberian Peninsula compared with other wetlands; therefore, we have studied the chemical parameters of water and the diversity patterns of testate amoebae in the western Iberian Peninsula to better understand the current status of these ecosystems. The analysis of testate amoeba communities showed an inverse relationship between the diversity and conservation status of these peatlands, both in relation to chemical parameters (i.e., pH, electrical conductivity, phosphates) and to the proportion of anthropized area, with a marked geographical pattern in the degree of anthropogenic disturbance.

Keywords: testate amoebae; diversity; human impact; peatlands water chemistry; southern Europe

1. Introduction

Peatland ecosystems develop as a result of a complex biogeochemical balance, generating a sink for atmospheric C as a result of the soil–atmosphere interaction [1–3]. However, the disruption of peat generation processes due to changes in climatic conditions (e.g., droughts or high temperatures) or anthropic impact (e.g., drainage, nutrient input, peat extraction, and construction of infrastructures) leads to the release of greenhouse gasses due to peat mineralization [4,5]. This is especially critical in the context of climate change because of its positive feedback effect on global warming [6]. In addition, the loss of peatland ecosystems has a direct impact on the biodiversity associated with these ecosystems [7]; for this reason, they have been catalogued as a priority habitat of community interest by the Habitats Directive 92/43/EEC of the European Union [8] in order to establish protection areas and promote monitoring programs that guarantee their conservation, especially compromised by climatic conditions in southern Europe due to the greater fragility of the balance that allows for peat formation and due to the limited size and high fragmentation of this type of habitat [9,10].

Testate amoebae are one of the most studied microorganisms in Sphagnum peatlands due to their ubiquity and great diversity, but especially because their close relationship with environmental factors makes them one of the most informative ecological indicators in peatland ecosystems [11–13]. Testate amoebae as predators play a key role in the microbial food web structure, which critically determines the C cycle in semi-aquatic ecosystems [14,15]. The high specific diversity of testate amoebae is linked to the microhabitats originated...
by microtopography and the consequent spatial heterogeneity patterns that determine biogeochemical processes, therefore leading to important macro- and microenvironmental gradients [15–18]. Therefore, the composition and structure of testate amoeba communities are highly sensitive to ecological conditions of peatland ecosystems, as shown by numerous studies on ecological succession, seasonal spatio-temporal variability, and interaction between testate amoebae and vegetation [19–27]. For these reasons, testate amoebae are widely used for paleoecology inferences in peatland studies [22,28–31].

The use of testate amoeba diversity as a bioindicator in *Sphagnum* peatlands constitutes a powerful tool to assess the conservation status of peatland ecosystems and to understand the magnitude of disturbances entailed by anthropogenic impacts and environmental changes [16,22]. In this sense, the study of testate amoeba communities is essential due to the poor knowledge of the current conservation status of Iberian peatland ecosystems [10]. Therefore, we have studied the diversity of testate amoebae in western Iberian peatlands to identify the main chemical parameters of water that determine the composition and structure of testate amoeba communities.

2. Materials and Methods

2.1. Study Site

A total of 37 peatlands representative of those present in the western Iberian Peninsula were studied in the period between March and June 2017 (Table 1; Figure 1). Of these, 33 peatlands were studied within the context of a program to monitor the ecological conditions of peatlands of Galicia (NW Spain). In addition, four peatlands in the Serra da Estrela (Portugal) were included in this study. Most Iberian peatlands are located in the Atlantic region, followed by the Mediterranean and Pyrenean geographical areas. Fens constitute the predominant peatland typology, while bogs (both raised and blanket bogs) are scarcer and are mainly found in the Atlantic area. The NW Iberian Peninsula is the geographical area with the highest density of peatlands and the highest diversity of peatland typologies [10]. According to the Habitats Directive 92/43/EEC of the European Union [8], twenty-nine are acidic fens (7140), two are blanket bogs (7130), two are raised bogs (7110), two are meso-eutrophic fens (7230), and the para-peaty wetland AG_G5, a 50-cm-thick organic soil but with a proportion of organic carbon <15% [32]. The western Iberian Peninsula is formed by the Iberian Massif, mainly consisting of Paleozoic metamorphosed sedimentary formations intruded by plutonic rocks [33]. Although the dominant lithology is constituted by siliceous rocks, some peatlands are located on local gabbro outcrops, such as CO_ALC, or on ultramafic rocks, such as CA_CF and CA_QUE. The characteristics of the Iberian Peninsula define it as a climatic hotspot in terms of regional modulation of global climate models [34]. The western Iberian Peninsula has a temperate climate with a dry summer season, and average temperatures of the coldest month ranging between 4 and 9 °C. A gradient exists from the NW Iberian Peninsula, which includes most of Galicia and which shows a greater oceanic influence (CsB), with mild temperatures and high rainfall (900–1200 mm), towards the S Iberian Peninsula (Csa), with average temperatures over 22 °C in the warmest months and with low rainfall (400–800 mm). However, the effect of the orography and NW–SW orientation of the high mountains of Serra da Estrela, with a maximum elevation of 1993 m a.s.L, result in higher mean rainfall values (900–1100 mm) due to the interception of oceanic humid air masses that enter the Iberian Peninsula [35]. Finally, anthropogenic impacts and the degrees of transformation of peatland ecosystems have been evaluated in situ by direct observation (i.e., livestock, drainage channels, plantations), and the percentages of the affected area with respect to the total area have been estimated.
| Peatlands                  | Code   | Locality (Province)                  | Country | Geographic Area | EU Habitat Directive         | Coordinates (Datum ETRS89) | Altitude (m a.s.l.) |
|---------------------------|--------|-------------------------------------|---------|----------------|------------------------------|-----------------------------|---------------------|
| Cruz do Frade (1)         | CA_CF  | Cariño (A Coruña, Galicia)          | Spain   | Serra da Capelada (CA) | Meso-eutrophic               | 7230 29 T 583605 4837869 | 530                 |
| Queizal (2)               | CA_QUE | Cariño (A Coruña, Galicia)          | Spain   | Serra da Capelada (CA) | Meso-eutrophic               | 7230 29 T 584005 4837885  | 490                 |
| Rego do Tremoal (3)       | X_VRRT | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Acidic fens                  | 7140 29 T 626494 4812597  | 570                 |
| Tremoal de Penas Gordas (4)| X_TPG  | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Acidic fens                  | 7140 29 T 626534 4811601  | 625                 |
| Veiga do Rial (5)         | X_VR   | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Acidic fens                  | 7140 29 T 616311 4810992  | 750                 |
| Tremoal da Revolta (6)    | X_TR   | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Acidic fens                  | 7140 29 T 619398 4812159  | 700                 |
| Chao de Veiga Mol (7)     | X_CVM  | O Valadouro (Lugo, Galicia)         | Spain   | Serra do Xistral (X) | Raised bog                   | 7110 29 T 620745 4822070  | 695                 |
| Tremoal do Fiuco (8)      | X_TF   | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Raised bog                   | 7110 29 T 618996 4811886  | 700                 |
| Veiga do Tremoal (9)      | X_VT   | Abadín (Lugo, Galicia)              | Spain   | Serra do Xistral (X) | Acidic fens                  | 7140 29 T 620106 4811318  | 700                 |
| Chao de Lamoso (10)       | X_CL   | O Valadouro (Lugo, Galicia)         | Spain   | Serra do Xistral (X) | Blanket bog                  | 7130 29 T 617074 4817170  | 1020                |
| Pena da Cedela (11)       | X_PC   | Muras (Lugo, Galicia)               | Spain   | Serra do Xistral (X) | Blanket bog                  | 7130 29 T 616449 4816677  | 1014                |
| Rego de Pena Grande (12)  | TC_PG1 | Xermade (Lugo, Galicia)             | Spain   | Terra Chá (TC)     | Acidic fens                  | 7140 29 T 605556 4692824  | 480                 |
| Rego de Pena Grande (13)  | TC_PG3 | Xermade (Lugo, Galicia)             | Spain   | Terra Chá (TC)     | Acidic fens                  | 7140 29 T 605785 4692891  | 485                 |
| Lagoa de Fabás (14)       | TC_GZ  | Vilalba (Lugo, Galicia)             | Spain   | Terra Chá (TC)     | Acidic fens                  | 7140-pond 613605 4799346  | 435                 |
| Lagoa de Alcaíán (15)     | CO_ALC | Coristanco (A Coruña, Galicia)      | Spain   | Chairas Occidentais (Western Plains) (CO) | Acidic fens | 7140-6410 Moléine meadows-Atlantic wet heath  | 29 T 520943 4775406  | 400                 |
| Peatlands                        | Code   | Locality (Province) | Country | Geographic Area | EU Habitat Directive | Coordinates (Datum ETRS89) | Altitude (m a.s.l.) |
|---------------------------------|--------|---------------------|---------|----------------|----------------------|-----------------------------|---------------------|
| Poza do Lameiro (16)            | CO_PL  | Lousame (A Coruña, Galicia) | Spain   | Chairas Occidentais (Western Plains) (CO) | Acidic fens | 7140 | 29 T | 515709 | 4733290 | 450 |
| Rego Grande (17)                | BO_B32 | Toques (A Coruña, Galicia) | Spain   | Montes do Bocelo (BO) | Acidic fens | 7140- grasslands | 29 T | 581205 | 4759180 | 600 |
| Rio Ameneiros (18)              | BO_RA31| Toques (A Coruña, Galicia) | Spain   | Montes do Bocelo (BO) | Acidic fens | 7140 | 29 T | 581388 | 4760452 | 705 |
| Barranqueira da Devesa da Graña (19) | BA_118.2 | Boiro (A Coruña, Galicia) | Spain   | Serra do Barbanza (BA) | Acidic fens | 7140 | 29 T | 505495 | 4727159 | 540 |
| Rego da Puntaña (20)            | BA_118.3 | Boiro (A Coruña, Galicia) | Spain   | Serra do Barbanza (BA) | Acidic fens | 7140 | 29 T | 505318 | 4726200 | 555 |
| Chan das Brañas (21)            | BA_118.6 | Boiro (A Coruña, Galicia) | Spain   | Serra do Barbanza (BA) | Acidic fens | 7140 | 29 T | 502641 | 4723368 | 650 |
| Rego das Brañas (22)            | BA_118.7 | Boiro (A Coruña, Galicia) | Spain   | Serra do Barbanza (BA) | Acidic fens | 7140 | 29 T | 503665 | 4722536 | 525 |
| Campa dos Extremos (23)         | ANC_CE1 | Cervantes (Lugo, Galicia) | Spain   | Serra dos Ancares (ANC) | Acidic fens | 7140 | 29 T | 675506 | 4743659 | 1615 |
| As Pontellas 1 (24)             | SU_C19 | A Lama (Pontevedra, Galicia) | Spain   | Serra do Suido (SU) | Acidic fens | 7140-7110 | 29 T | 554323 | 4696467 | 855 |
| As Pontellas 2 (25)             | SU_C21 | A Lama (Pontevedra, Galicia) | Spain   | Serra do Suido (SU) | Acidic fens | 7140 | 29 T | 554364 | 4696930 | 860 |
| Costa de Uxía (26)              | SU_C26 | A Lama (Pontevedra, Galicia) | Spain   | Serra do Suido (SU) | Acidic fens | 7140 | 29 T | 551521 | 4695120 | 820 |
| As Landras (27)                 | SU_C32 | A Lama (Pontevedra, Galicia) | Spain   | Serra do Suido (SU) | Acidic fens | 7140 | 29 T | 553636 | 4696430 | 870 |
| Brañas da Groba (28)            | AG_G5  | Oia (Pontevedra, Galicia) | Spain   | Serra da Groba (AG) | Acidic fens | 7140 | 29 T | 513096 | 4657227 | 525 |
| A Portela (29)                  | AG_V10 | Oia (Pontevedra, Galicia) | Spain   | Serra de Valga (VA) | Acidic fens | 7140 | 29 T | 512525 | 4648061 | 350 |
### Table 1. Cont.

| Peatlands                     | Code     | Locality (Province) | Country | Geographic Area | EU Habitat Directive | Coordinates (Datum ETRS89) | Altitude (m a.s.l.) |
|-------------------------------|----------|---------------------|---------|-----------------|----------------------|-----------------------------|---------------------|
| Gándaras de Budíño (30)       | LOR_GB   | Porriño (Pontevedra, Galicia) | Spain   | Louro River Valley (LOR) | Acidic fens | 7140 29 T 530591 4660889 | 30                  |
| Veiga de Gopellás (31)        | LIR_VB124 | Calvos de Randin-Muños (Ourense, Galicia) | Spain   | Limia River Valley (LIR) | Acidic fens | 7140 29 T 593321 4647552 | 865                 |
| Val de Deus (32)              | LIR_VD129 | Muños (Ourense, Galicia) | Spain   | Limia River Valley (LIR) | Acidic fens | 7140 29 T 589484 4645252 | 875                 |
| Veiga do Calvelo do Rio (33)  | LIR_VC32 | Baños de Molgas (Ourense, Galicia) | Spain   | Limia River Valley (LIR) | Acidic fens | 7140 29 T 612374 4675533 | 520                 |
| Covao da Clareza (34)         | F_CC     | Seia (Beira Alta)    | Portugal | Serra da Estrela (E) | Acidic fens | 7140 29 T 617938 4466145 | 1820                |
| Cabeço do Talao (35)          | E_CT     | Seia (Beira Alta)    | Portugal | Serra da Estrela (E) | Acidic fens | 7140 29 T 616955 4466623 | 1850                |
| Pedras Guiereiras (36)        | E_PGU    | Seia (Beira Alta)    | Portugal | Serra da Estrela (E) | Acidic fens | 7140 29 T 614406 4468721 | 1590                |
| Lagoa Seca (37)               | E_LS     | Seia (Beira Alta)    | Portugal | Serra da Estrela (E) | Acidic fens | 7140 29 T 616302 4469814 | 1625                |

#### 2.2. Sampling and Testate Amoeba Extraction

Testate amoeba samples were collected from the *Sphagnum* stratum in analogous microhabitats (20- to 30-cm-tall moist *Sphagnum* hummocks); five spatial replicas were collected in each peatland. The selection criteria for *Sphagnum* hummocks as the target microhabitat are due to the greater dominance of *Sphagnum* and the high sensitivity to hydrological alteration, especially to the duration and intensity of the dry period. Testate amoebae were extracted from a volume of 10 cm³ of *Sphagnum*, using only the section between 5 and 10 cm below the top of the capitulum, thus avoiding the vertical microdistribution of species in the upper centimeters [11,36]. Moss samples were disaggregated in distilled water and shaken for 10 min to maximize the release of testate amoebae. Testate amoebae were extracted using a standard sieving method [37] through nested 355 µm and 10 µm sieves. The sieved size fraction was washed by centrifugation at 900 × g for 5 min and resuspended in a volume of 5 cm³. Slides were mounted fresh, and a minimum of 200 thecae were counted under light microscopy with a Nikon Eclipse E600 equipped with Plan-Apochromat 40x (N.A. 0.95). The relative abundance of each taxon was calculated as a percentage of the total number of thecae counted. A total of 67 testate amoeba taxa have been found in western Iberian peatlands. The list of testate amoeba taxa and their codes is summarized in Supplementary Table S1, and abundance data (%) of the main taxa (>10% in at least one peatland) of testate amoeba communities in the western Iberian peatlands are summarized in Supplementary Table S2.
2.2. Sampling and Testate Amoeba Extraction

Figure 1. Known peatland ecosystems (yellow) in the Iberian Peninsula (SW Europe) according to Pontevedra-Pombal et al. [10], and locations of the thirty-seven western Iberian peatlands where testate amoeba communities were studied (orange) (Table 1).

2.3. Water Chemistry Analysis

Electrical conductivity and pH were measured in situ using a Hanna HI 98,194 multiparametric probe. Three 50 mL replicates of water samples for nutrient and cation chemical analysis were collected from water-saturated moss surfaces at the same points where samples for testate amoeba analysis were collected (Table 1; Figure 1). Previously to nutrient sample collection, bottles were treated with 8% HNO₃ (c/v) for 48 h and repeatedly washed with distilled water. Water samples for nutrient analysis were frozen at −20 °C until analysis, and samples for cation determination were kept at 4 °C, avoiding direct sunlight.

Content of cations (calcium and magnesium) in water was measured using a Perkin Elmer 2100 flame atomic absorption spectrometer. Nutrients in water (i.e., nitrate, nitrite, ammonium, and phosphates) were analyzed with an Integral Futura autoanalyzer system (Alliance Instruments), which uses separate analytical lines to determine nitrates, nitrites, and ammonium according to standard colorimetric methods [38]. Total reactive phosphorus was measured in unfiltered water samples using colorimetric methods according to Murphy and Riley [39]. Measurement precision was assessed using relative standard deviation and always remained below 5%. Detection limits, expressed as three times the standard deviation of 10 replicated measurements of reagent blanks, were 0.28 µM for nitrate, 0.10 µM for nitrite, 0.10 µM for ammonium, and 0.16 µM for phosphate. Dissolved inorganic nitrogen (DIN) was calculated from the results obtained for all the inorganic nitrogen sources measured (i.e., DIN = nitrates + nitrites + ammonium). The chemical parameters of peatlands water are summarized in Supplementary Table S3.
2.4. Statistical Analysis

Spatial parameters (i.e., UTM X (m), UTM Y (m), and altitude (m a.s.l.)) have been log-transformed to base 10 so that their orders of magnitude were similar to those of water chemistry data and relative abundances of functional groups for statistical treatment. Calcium and magnesium (meq L\(^{-1}\)) and nutrients (i.e., nitrate, nitrite, ammonium, and phosphates; µmol L\(^{-1}\)) were transformed to equivalent stoichiometric units for statistical analysis. The nominal variable “typology” has been coded according to degree of ombrotrophy as follows: para-peaty ecosystems = 5, meso-eutrophic peatlands = 4, acidic fens = 3, raised bogs = 2, and blanket bogs = 1. Four functional groups were considered according to Bonnet [40]: mixotrophs (e.g., Amphitrema wrightianum, Archerella flavum, Heleopera sphagnicola, Hyalosphenia papilio), filose (r-strategists) (e.g., Euglypha, Corythion, Trinema), lobose (K-strategists) (e.g., Arcella, Nebela, Centropyxis, Difflugia), and a fourth category called Indifferent, which grouped all other unassigned taxa. Additionally, the L (lobose) to F (filose) ratio was calculated.

The diversity of testate amoeba taxa in each peatland ecosystem was evaluated using the Shannon diversity index (H) [41]:

\[
H = - \sum_{i=1}^{s} p_i \ln (p_i)
\]

where \(p_i\) is the proportion of the number of individuals within a taxon with respect to the total number of individuals.

The relationship between the three datasets of variables in western Iberian peatlands (i.e., functional groups of testate amoebae, physicochemical parameters of water, and geospatial data) has been explored using Spearman correlation analysis, applying a Student’s t-test to determine the level of statistical significance of correlations between parameters. Moreover, the datasets were explored using Redundancy Analysis (RDA) to summarize the main patterns of variation in the structure of testate amoeba communities as a function of the dataset of the chemical parameters of the water and environmental variables, considering those taxa with mean abundances greater than 5% in at least one peatland, or in functional groups that could be explained by the set of main physicochemical variables.

Chemical data, anthropized area, Shannon’s diversity index, functional groups of testate amoebae, and L:F ratio were analyzed as a function of the previously defined classes based on the structure of testate amoeba communities, applying the non-parametric Kruskal–Wallis test (H) [42]. Moreover, the affinities among the structures of testate amoeba communities from the thirty-seven peatland ecosystems were analyzed using non-metric multidimensional scaling (NMDS) and the analysis of similarities (ANOSIM) to test the statistical differences between the groups obtained. All statistical analyses were performed using XLSTAT [43], with the exception of NMDS and ANOSIM analysis, which was performed using the vegan package [44] with R [45].

3. Results

3.1. Water Quality

The pH showed a statistically significant positive correlation with peatland typology, with acidity increasing from minerotrophic to ombrotrophic conditions, and a decrease in the anthropized area. The pH range of the studied peatlands was divided according to the three main typologies: (i) pH 3.90–4.5 for blanket bogs and raised bogs (ombrotrophic), (ii) pH 4.5–5.43 for acidic fens (minerotrophic) and (iii) pH 5.60–6.79 for meso-eutrophic fens or degraded acidic fens (minerotrophic). The first category (i) includes two blanket bogs (X_PC, X_CL), two raised bogs (X_CVM, X_TF), and two headwater fens (X_VR, TC_PG3). The second category (ii) is comprised of most of the acidic fens studied. The third category (iii) includes two typologies, meso-eutrophic fens on ultramafic rocks, serpentinites (CA_CF, CA_QUE) or gabros (CO_ALC), and degraded acidic fens with a
high percentage of human-altered areas, such as TC_GZ, LIR_VB124, and LOR_GB (81.4%, 33.5%, and 43.8% of anthropized area, respectively; Table S3).

pH and phosphates showed a statistically significant positive correlation with electrical conductivity (Supplementary Table S4). The range of electrical conductivity was <100 µS cm$^{-1}$ in most peatland ecosystems, although it was higher in some peatlands with a high proportion of anthropized area (25.3–100%), such as TC_GZ (108.72 µS cm$^{-1}$), CO_ALC (111.60 µS cm$^{-1}$), LIR_VD129 (265.00 µS cm$^{-1}$), and LIR_VB124 (325.00 µS cm$^{-1}$). Low calcium and magnesium levels were consistent with low electrical conductivity values, typical of soft waters (<9 mg L$^{-1}$) (Table S3). Ammonium showed a statistically significant positive correlation with spatial arrangement (UTM Y), and nitrite correlated positively with pH and anthropized area within the peatland ecosystems, while nitrate did not correlate with any other environmental variables. Therefore, DIN was positively correlated with spatial arrangement (UTM Y) and anthropized area (Table S4). Phosphates ranged from 0.01 to 0.09 mg L$^{-1}$, and levels of dissolved nitrogen nutrients were low in all peatland ecosystems, with values in the following ranges: nitrates: 0.03–3.98 mg L$^{-1}$, nitrites: 0.02–0.58 mg L$^{-1}$; ammonium: 0.02–0.39 mg L$^{-1}$, and DIN: 0.09–4.10 mg L$^{-1}$ (Table S3). The anthropized area in peatland ecosystems showed significant correlation with spatial variables, suggesting that the magnitude of the anthropic impact follows a differential geographical pattern, with higher degrees of impact on peatland ecosystems at low altitudes and near the coast than on those located in inland mountain areas, and also linked to DIN inputs (Table S4).

3.2. Testate Amoeba Assemblages

Non-metric multidimensional scaling (NMDS) analysis on the structure of testate amoeba communities revealed three significant classes (ANOSIM R = 0.620, p-value = 0.001): class 1, mainly composed of the testate amoeba communities in the Suído, Bocelo and Terra Cha peatlands, as well as in the Xistral blanket bogs (X_PC, X_CL) and in fens LIR_VB124 and LIR_VD129; class 2, mainly composed of the testate amoeba communities in the Xistral, Capelada, Ancares, Estrela, and LOR_GB, including fen X_TR and the Xistral raised bogs (X_CVM and X_TF); and class 3, mainly composed of the testate amoeba communities in the fens of Barbanza, CO_ALC and LIR_VC32. Fen CO_PL and para-peaty peatland AG_G5 are highly anthropized and drained ecosystems whose taxa diversity levels were very different from the remaining peatlands studied (Figure 2).

The proportions of the different functional groups in the three cluster classes were the following: 0–5% mixotrophic taxa, 24–37% r-strategist taxa, 35–45% K-strategist taxa, and 18–30% indifferent taxa. Mixotrophic taxa were absent from class 3 and present in small proportions (∼5%) in classes 1 and 2. Meanwhile, the proportions of r- and K-strategists were 1:1 in class 2, while K-strategist taxa were dominant (40–45%) in classes 1 and 3, although differences were very small (Figure 3). According to peatland typology, testate amoeba communities in blanket bogs (X_PC, X_CL) and in the only raised bog in Suído (SU_C19) were grouped in class 1; those in the Xistral raised bogs (X_CVM, X_TF) and meso-eutrophic peatlands (CA_CF, CA_QUE) were grouped in class 2, and the communities in the para-peaty ecosystem AG_G5 and in the highly degraded fen CO_PL were not classified (Figure 2).

The composition of testate amoeba communities in peatlands showed low percentages of the dominant taxa due to the important spatial variability within each ecosystem. However, in general, class 1 was dominated by Nebela collaris, with high proportions of Assulina muscorum, Corythion dubium, and Euglypha spp., with low L:F ratios (0.52–1.63); class 2 showed co-dominance of Nebela collaris, Assulina seminulum, Heloepora petricola, Centropyxis aculeata, Corythion dubium, and Trinema lineare, with high L:F ratios (0.66–11.6); class 3 was dominated by Centropyxis aerophila, with high proportions of Corythion dubium and Euglypha strigosa and with high L:F ratios (0.78–3.05); and unclassified ecosystems were dominated by Alabasta longicollis, Padaungiella tubulata, and Microcorycia radiata, with
a high L:F ratio (3.01), in the highly anthropized fen CO_PL and exclusively by the Nebela genus in the para-peaty ecosystem AG_G5, with a high L:F ratio (2.13) (Table S3; Figure 2).

Communities of testate amoebae according to peatland types showed some differences with respect to their assigned classes: in the case of Xistral (X_CL, X_PC), blanket bogs included in class 2 had a greater proportion (~10%) of mixotrophs (i.e., Amphitrema wrightianum, Helicopera sphagni and Hyalosphenia papilio). However, testate amoeba communities in the meso-eutrophic fens of Capelada (CA_CF, CA_QUE) and in the raised bogs of Xistral (X_CVM, X_TF) and Suido (SU_C19), both of which were included in class 1, did not show any significant differences compared with acidic fens, although, in the latter, differences were found between Sphagnum samples from the ombrotrophic dome and from the marginal fen (Table S2).

Peatland ecosystems as a function of testate amoeba communities showed the following mean altitudes: class 1 1036.8 ± 583.8 m a.s.l. (625–1820), with the exception of the acidic fen Gándaras de Budiño in the Louro River (LOR_GB), located at an altitude of 30 m a.s.l.; class 2 670.8 ± 197.8 m a.s.l. (350–875), with the exception of the Xistral blanket bogs, located at 1014 m a.s.l. (X_PC) and 1020 m a.s.l. (X_CL); and class 3 534 ± 89.4 m a.s.l. (400–650). Finally, those unclassified were located at 450 m a.s.l. (CO_PL) and 525 m a.s.l. (AG_G5) (Table 1; Figure 2).
Figure 3. Biplots of the Redundancy Analysis (RDA) performed on the dataset of environmental parameters and testate amoeba community structures in western Iberian peatlands, both based on (A) testate amoeba community structures and (B) testate amoeba functional groups (i.e., K-strategists, r-strategists, mixotrophic, indifferent, L:F ratio and Shannon diversity index).
The Kruskal–Wallis test on Shannon index values and L:F ratios as a function of the three cluster classes based on testate amoeba communities showed significant differences between class 1, on one hand, and classes 2 and 3, on the other. The Shannon index showed significantly lower values in class 3 (1.28–2.74) and class 2 (1.45–2.96) than in class 1 (2.76–3.55) (H = 23.867; p-value < 0.0001). Additionally, L:F ratios were significantly higher in class 2 (0.66–11.6) and class 3 (0.78–3.05) than in class 1 (0.52–1.63) (H = 16.776; p-value = 0.001).

3.3. Effects of Environmental Variables on the Testate Amoeba Assemblages

A total of 67 testate amoeba taxa were identified in western Iberian peatland ecosystems (Table S1); the structure of testate amoeba communities is summarized in Table S2. The RDA analysis showed that six environmental variables (i.e., pH, electrical conductivity, DIN, phosphates, UTM Y and anthropized area) explained 61.48% of the total variance of the structure of testate amoeba communities, both in terms of taxa abundances (Figure 3A) and functional groups, respectively (Figure 3B). All the environmental variables were included in the RDA analysis, with exception of altitude and UTM X because these variables show a poor relationship with the other variables considered in the correlation test, or redundant variables (i.e., calcium and magnesium with electrical conductivity; ammonium, nitrite and nitrate with DIN). The first component (RDA1) explained 35.62% of the total variability of the dataset, showing an environmental gradient along an axis defined by peatland ecosystems with a higher ombrotrophy and testate amoeba diversity; on one end, and those with increased nutrient levels and minerotrophic conditions, on the other. The Xistral and Serra da Estrela peatland ecosystems had greater testate amoeba diversities, while the peatland ecosystems of Barbanza, A Grova mountains, and Chairas Occidentais (Western Plains) showed increasing degrees of anthropogenic impact, with the poorest water quality and the lowest testate amoeba diversity (Figure 3A,B). The Limia River valley peatlands are mainly coupled to a river system, and water chemical composition was highly influenced by runoff water (Figure 3A,B). The taxa *Trinema lineare*, *Trinema enchelys*, *Centropyxis aculeata*, *Hyalosphenia subflava*, *Euglypha heterospina*, *Nebela flabellulum*, *Amphitrema wrightianum*, and * Archerella flavum* (Figure 3A), and therefore the r-strategist and mixotroph functional groups (Figure 3B), were linked to increased values of the Shannon diversity index and to ombrotrophic conditions.

The second component (RDA2) explained 25.86% of the total variability of the dataset, which we interpreted as an increase in nutrients and minerotrophic conditions either due to (i) hydrological conditions with a greater runoff contribution (i.e., electrical conductivity, phosphates), and/or (ii) the degree of anthropogenic impact (i.e., pH, DIN, UTM Y, anthropized area) (Figure 3). The abundance of the taxa *Nebela collaris*, *Assulina muscorum*, *Diffugia sp. 1*, *Hyalosphenia ovalis*, *Heleopera rosea*, and *Heleopera sphagni* (Figure 3A) (and, therefore, of the Indifferent functional group) and the L:F ratio (Figure 3B) are linked to the increase in (i) runoff contribution to peatland ecosystems. The taxa *Centropyxis aemphila*, *Diffugia oblonga*, *Padaungiella tubulata*, *Alabasta longicollis*, *Nebela barbata*, *Nebela militaris*, *Argynnia crenulata*, *Euglypha compressa*, *Euglypha rotunda* and *Euglypha strigosa* (Figure 3A) (and, therefore, the K-strategists functional group; Figure 3B) are linked to the greater magnitude of the (ii) anthropogenic impacts on peatland ecosystems.

The correlation between the variables in the datasets indicates a decrease in the human impact with altitude and along the West–East axis, since the same trend has been observed for the Shannon diversity index; conversely, there was an increase in variables linked to runoff water (i.e., phosphates, electrical conductivity) or to human impact (i.e., pH, DIN, anthropized area) (Table S4; Figure 3). The correlations between functional groups of testate amoebae and environmental datasets also reflected the effect of human impact on peatland ecosystems, with decreasing ratios of mixotrophic and r-strategist (filose) taxa with respect to K-strategist (lobose) taxa and the subsequent increase in the L:F ratio. Mixotrophic taxa positively correlated with the Shannon diversity index and negatively correlated with high trophic conditions of peatlands. Mixotrophic and K-strategist taxa correlated with
spatial arrangement (i.e., UTM X, UTM Y) and the magnitude of the human disturbance of peatland ecosystems. The proportions of K-strategist taxa correlated positively with anthropized area and negatively with the Shannon diversity index, while r-strategist taxa negatively correlated with phosphates, calcium, and magnesium and, conversely, indifferent taxa positively correlated with the same water chemical parameters (Table S4; Figure 3). The L:F ratio correlated negatively with more ombrotrophic typologies and positively with electrical conductivity and phosphates (Table S4). Therefore, the proportion of K-strategist and indifferent taxa correlated negatively with the proportion of r-strategist and mixotrophic taxa; moreover, the proportion of r-strategist taxa negatively correlated with indifferent taxa (Table S4; Figure 3).

4. Discussion

4.1. Iberian Peatlands Water Chemistry

The anthropogenic transformation of peatland ecosystems entails both the alteration of the hydrological regime (e.g., drainage for agricultural use or for water supply) and of the geochemical balance of water by increasing pH and electrical conductivity due to the increased contributions of runoff water and nutrients. However, land uses (e.g., agriculture, livestock, and wastewater discharge) also involve greater nutrient inputs into peatland ecosystems, causing the disruption of complex and fragile biogeochemical cycles in peatlands [46–50]. In addition, experiments on Swedish peatland ecosystems showed that increases in pH, electrical conductivity, dissolved inorganic nitrogen, and phosphorus are associated with the degree of anthropogenic transformation of peatland ecosystems; conversely, a decrease in these parameters has been observed in long-term peatland restoration programs [50].

The water chemistry data measured in western Iberian peatlands were within the range expected for Sphagnum-dominated peatlands [51–53]. pH is one of the most important chemical parameters of water for the biogeochemistry of peatlands. Water pH ranges indicate the main source of water: blanket bogs (X_PC, X_CL) and raised bogs (X_CVM, X_TF, SU_C19), with pH ranging between 3.5 and 4.5, are fed exclusively by rainfall, while peatland ecosystems with higher pH ranges (4.5–7) also receive runoff contributions (Table S3) [54]. pH in acidic fens, the most common type of peatland, ranges from 4.5 to 5.5, suggesting that their main water sources are rainwater and very weakly mineralized runoff water from headwater mountain streams. In other peatland ecosystems with pH ranges of 5.5–7, such as para-peaty ecosystems (AG_G5), meso-eutrophic peatlands (CA_QUE, CA_CF), highly anthropized acidic fens (CO_ALC, CO_PL), and fens coupled to rivers (LIR_VB124, LOR_GB) or ponds (TC_GZ), mineralized runoff water is their main water source (Table S3) [54].

The water composition in western Iberian peatlands is fairly similar to that of weakly mineralized soft waters of the western Iberian peninsula (<400 µS cm⁻¹), with a primary contribution from oceanic rainwater and a poor contribution of ions from the dominant siliceous lithology [55,56], although their low electrical conductivity ranges (<100 µS cm⁻¹; Table S3) are similar to those found in other Iberian fens, such as those in the Sierra Segun- dera Mountains [46] or the Bassa Nera fen and Tresurco fens complex in the Pyrenees [57,58], as well as to other European and North American peatlands [51–53]. Western Iberian peatlands are very poor in nutrients compared with other peatlands in temperate regions of Europe and North America [51–53], both in terms of phosphorus (phosphates < 0.10 mg L⁻¹) and of dissolved inorganic nitrogen (i.e., nitrate < 2.40 mg L⁻¹, nitrite < 0.58 mg L⁻¹, ammonium < 0.39 mg L⁻¹, and DIN < 2.75 mg L⁻¹). However, phosphates are linked with electrical conductivity, suggesting the influence of runoff water on peatland water composition (Table S3; Figure 3), very evident in fens associated with the Limia River (LIR_VB124, LIR_VD129). DIN and nitrites are exclusively linked to the anthropized area (Table S4; Figure 3). Traditionally, human impacts on western Iberian peatlands are related to livestock burden (both of cattle and of wild horses) due to the use of green grass as pasture during the summer season, which remains important nowadays in virtually all
the studied peatlands. The partial drainage of peatlands is due to several causes: hydrological control due to the construction of water supply reservoirs, the construction of infrastructures (Gándaras de Budiño), and, more generally, drainage for agricultural use, particularly important in the desiccation of Lake Alcaián, or, more recently, for tree plantations (i.e., Pinus, Eucalyptus). Both livestock and agriculture, or even hydrological alteration, directly or indirectly cause an increase in nitrogen input in peatlands, impacting ecological processes and the carbon storage function of peatland [59].

Some peatlands with a high proportion of anthropized area showed a deviation from normal electrical conductivity values (>100 µS cm⁻¹) (i.e., TC_GZ; CO_ALC; LIR_VD129; LIR_VB124) and nutrients, with slightly high mean nitrate (i.e., X_TPG = 2.56 mg L⁻¹; SU_C32 = 2.80 mg L⁻¹; G_V10 = 3.98 mg L⁻¹) and ammonium (i.e., CA_CF = 0.24 mg L⁻¹; TC_PG1 = 0.21 mg L⁻¹; TC_PG3 = 0.23 mg L⁻¹; TC_GZ = 0.15 mg L⁻¹; CO_ALC = 0.39 mg L⁻¹; BA_SB118.3 = 0.2 mg L⁻¹). Phosphate levels were low in all the western Iberian peatlands (<0.05 mg L⁻¹) (Table S3), similarly to other Iberian fens [46,57,58], although some peatlands coupled to rivers have slightly high levels of phosphates (0.08–0.09 mg L⁻¹; TC_PG1, X_TR, LIR_VB124, and LIR_VD124) with respect to the threshold value of 0.05 mg L⁻¹ of total phosphorus for peatlands to be considered oligotrophic, according to Eurela and Holappa [60] and Malmer [61] based on the study of interactions between water chemistry and peatland vegetation in north-western Europe and in Finland, respectively.

4.2. Ecological Drivers of Testate Amoeba Communities

The known diversity of testate amoebae in Iberian peatlands is around one hundred taxa [62–68]; therefore, the number of 67 taxa found in western Iberian peatlands is representative of these ecosystems and widely overlaps with the known testate amoeba fauna of *Sphagnum* peatlands in Europe [11,12,19,20,26,36,53,69]. Moreover, some rare taxa have not been previously cited in the Iberian Peninsula, such as *Microcorycia radiata*, *Wailesella eboracensis*, *Corythion asperulum*, *Amphitrema wrightianum* and *Nebela flabellulum*; the latter two are closely linked to ombrotrophic habitats in European peatland ecosystems [19,70,71].

In addition, the climatic patterns of the Iberian Peninsula establish a SE–NW gradient in terms of the frequency and extension of peatland ecosystems, which are more widely represented in the NW Iberian Peninsula [10]. Abundance peaks of testate amoebae in peatlands have been mainly reported in spring [16,72].

The diversity and structure of testate amoeba communities are closely linked to water availability and water chemistry [19,26,73,74], both of which are in turn influenced by the geographic location of the peatlands, showing affinities that are consistent with the peatland human disturbance pattern along the geographical gradient caused by human impact on the western Iberian landscape (Table S4; Figure 3). Moreover, landscape disturbance and chemical alterations caused by human impact on *Sphagnum* peatlands have also been reported to inversely relate to the altitudinal gradient in the northern Swiss Alps [11]. Human disturbance of *Sphagnum* peatland ecosystems is mainly associated with agricultural uses of the landscape, since this usually entails hydrological alterations and a greater water deficit. The greater runoff contribution and increased nutrient levels modify the main chemical parameters of water (pH, electrical conductivity, and nutrients), therefore causing a substitution of testate amoeba communities (Figure 3), as has been experimentally observed in N- and P-fertilized *Sphagnum* peatlands or sedge tundra [36,74] and in surface soils subjected to agricultural use [75]. Historically, population density and intensive agricultural activities have been higher in lowlands near coastal areas of Portugal (http://www.ine.pt; accessed on 12 February 2021) and in western coastal areas of Galicia (http://www.ine.es; accessed on 12 February 2021). In addition, peat formation and peatland restoration are fragile because the climate in these areas is milder due to the oceanic influence, and summer drought periods are more intense (https://www.meteogalicia.gal/; accessed on 12 February 2021) [76].

Koenig et al. [11] studied testate amoeba taxa in *Sphagnum* peatlands in relation to altitudinal gradient in the northern Swiss Alps (580–1890 m) and found diverse ecolog-
tical preferences related to micro-scale rather than meso- or macro-scale environmental conditions. In addition, Mitchell et al. [26] demonstrated a close relationship between the diversity of testate amoebae and the lithology of *Sphagnum* peatlands at different altitudes in the Italian Alps (1000–2000 m), and Krashevska et al. [77,78] found that testate amoeba communities from litter and soil surfaces in a tropical mountain forest at different altitudes (1000–3000 m) in Ecuador were intimately linked to soil composition and typology with respect to altitudinal patterns. Analogously, Bonnet [79–84] extensively established that testate amoebae showed preferences in terms of the geochemical properties inherent to different soil types.

However, environmental variables can follow pronounced geographical gradients and indirectly influence biogeochemical cycles at micro-scale [11]. Testate amoebae are very sensitive to water availability, as defined by the depth of the water table in *Sphagnum* peatlands [85–87], but also to the degree of moisture of the moss layer [88]. Therefore, longer water deficit periods in lowlands can be more intense in those microhabitats most exposed to the atmosphere in *Sphagnum* peatlands, such as hummocks, thus differentially determining the structure of testate amoeba communities [11,53].

The diversity and structure of testate amoeba communities follow a gradient defined by human disturbance in western Iberian peatlands: mixotrophic and filose taxa are linked to optimal ecological peatland conditions, while lobose taxa are linked to a certain degree of human disturbance. Therefore, high L:F ratios are associated with minerotrophic conditions (Figure 4). The results obtained may seem contradictory with the previous assumptions that lobose and filose species are ecological K- and r-strategists, respectively [27,40], and that a high L:F ratio is characteristic of more stable or climax habitats [40]. However, we interpret that these findings are a consequence of drought stress according to the disturbance degree of *Sphagnum* peatlands, since lobose taxa are more resistant to drought than filose ones [27,89]; this could also explain the higher proportion of lobose taxa in blanket bog microhabitats (i.e., X_CL, X_PC), similar to those found in the Suído peatlands (Figure 3).

![Figure 4](image-url)

*Figure 4.* Changes in diversity of testate amoeba communities along the environmental gradient defined by the human disturbance of peatland ecosystems.
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Mixotrophic taxa, especially present in ombrotrophic typologies such as raised and blanket bogs, are linked to the human disturbances of peatland ecosystems, with high diversities and low values for pH, electrical conductivity, nutrients, and proportion of anthropized area (Figure 3). These taxa, which host photosynthetic symbionts (e.g., Amphiotrema wrightianum, Archerella flavum, Hyalosphenia papilio, and Heleopera sphagni), have an adaptive advantage in oligotrophic and ombrotrophic conditions [15,16,74,90,91], although H. sphagni also tolerates minerotrophic conditions (Figure 3). These taxa are also characteristic of moist to wet microhabitats and are very rare or absent in dry conditions [36,68,74]. A. wrightianum and A. flavum are almost exclusively found on Sphagnum moss in peatland ecosystems [36] and have not been extensively recorded in the Iberian Peninsula, with only two previous records of A. flavum in Sphagnum minerotrophic peatlands in Gándaras de Budiño and Plans de Bohaví (Central Pyrenees) [62–67].

In addition, highly conserved Sphagnum peatlands have a higher proportion of taxa characteristic of humid ombrotrophic conditions, such as mixotrophs and other species like Centropyxis aculeata and large Nebela species [69], such as Nebela flabellulum, a typical species of European Atlantic raised bogs [71]. Higher runoff contributions and P inputs are associated with taxa characteristic of minerotrophic peatlands, such as Centropyxis aerophila, Heleopera petricola, Heleopera rosea, Nebela collaris, and Quadrulella symmetrica, as well as with the replacement of Assulina seminulum by Assulina muscorum under drought stress and in more minerotrophic conditions. Sphagnum peatlands with greater degrees of alteration, higher runoff contributions, high pH and DIN values, and greater proportions of anthropized area are dominated by Centropyxis aerophila, resistant to desiccation and tolerant to high levels of N in peatlands [26,36,69,74] (Figure 3). Corythion dubium, Assulina muscorum, Nebela tincta, and Trinema spp., all of them highly tolerant to desiccation and acidic conditions, are commonly found on a wide diversity of mosses outside of peatlands. Nevertheless, they have also been found to be ubiquitous in peatlands as indicators of drought and are dominant in peatlands with deep water tables and in cutover peatlands [26,36,69]. Cyphoderia ampulla, Alabasta longicollis, Nebela barbata, and Paduangiella tubulata are taxa linked to Sphagnum in aquatic or wet soil environments [64,92,93], as well as to other soil mosses such as Microcorycia radiata [94], all indicative of para-peaty ecosystems or highly transformed peatlands (i.e., CO_ALC, CO_PL, AG_G5) (Figure 3).

5. Conclusions

The diversity of testate amoeba is closely linked to their micro-scale ecological niches, defined by the complex biogeochemical interactions of multiple variables along environmental gradients as a function of the degree of spatio-temporal water saturation. Human use of peatland ecosystems, mainly related to agriculture and water availability, disturbs the hydrological regime and nutrient levels, resulting in an imbalance of the conditions that determine the microhabitats to which testate amoebae are highly sensitive. Water chemistry and testate amoeba community structures change along the geographical gradient defined by the magnitude of human impacts on western Iberian peatlands. Therefore, the diversity of testate amoeba is a good indicator of the ecological optimum and can be applied in ecological studies to estimate the resilience and recovery of peatland ecosystems, as well as at an environmental management level to assess the ecological conservation status of peatlands, especially considering the key role that these ecosystems play in supporting a very unique biodiversity and their geochemical role as carbon sinks in the context of climate change, as well as their fragility due to the climatic conditions at the southern limit of their distribution in southern Europe.

Supplementary Materials: The following are available online at https://www.mdpi.com/article/10.3390/d13060269/s1, Table S1: List of testate amoeba taxa from Sphagnum samples in the thirty-seven western Iberian peatlands studied. Table S2: Diversity and mean relative abundances of testate amoebae greater than 10% in at least one peatland ecosystem from Sphagnum samples of the thirty-seven peatlands studied in the western Iberian Peninsula. Table S3: Mean water chemistry data, functional groups and diversity of testate amoeba of the studied Iberian peatlands. Table S4:
Spearman’s correlation coefficients for the environmental variables and testate functional groups of the western Iberian peatlands. Student’s t-test was used to indicate statistically significant correlations (p-value < 0.05 * (orange); <0.1 ** (yellow)).

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