First Record of Fungal Diversity in the Tropical and Warm-Temperate Middle Miocene Climate Optimum Forests of Eurasia

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The middle Miocene Climate Optimum (MMCO) was the warmest interval of the last 23 million years and is one of the best analogs for proposed future climate change scenarios. Fungi play a key role in the terrestrial carbon cycle as dominant decomposers of plant debris, and through their interactions with plants and other organisms as symbionts, parasites, and endobionts. Thus, their study in the fossil record, especially during the MMCO, is essential to better understand biodiversity changes and terrestrial carbon cycle dynamics in past analogous environments, as well as to model future ecological and climatic scenarios. The fossil record also offers a unique long-term, large-scale dataset to evaluate fungal assemblage dynamics across long temporal and spatial scales, providing a better understanding of how ecological factors influenced assemblage development through time. In this study, we assessed the fungal diversity and community composition recorded in two geological sections from the middle Miocene from the coal mines of Thailand and Slovakia. We used presence-absence data to quantify the fungal diversity of each locality. Spores and other fungal remains were identified to modern taxa whenever possible; laboratory codes and fossil names were used when this correlation was not possible. This study represents the first of its kind for Thailand, and it expands existing work from Slovakia. Our results indicate a total of 281 morphotaxa. This work will allow us to use modern ecological data to make inferences about ecosystem characteristics and community dynamics for the studied regions. It opens new horizons for the study of past fungal diversity based on modern fungal ecological analyses. It also sheds light on how global variations in fungal species richness and community composition were affected by different climatic conditions and under rapid increases of temperature in the past to make inferences for the near climactic future.

Keywords: MMCO, Slovakia, Thailand, fossil fungi, lignite, palynology
INTRODUCTION

Fungi play key roles in the forest environment and are essential drivers of nutrient cycling in peat-producing forested wetlands (Heitman et al., 2018). Very little is known about how forested wetland fungal communities will change in relation to forecasted warming scenarios. Predictions of this type require large-scale, long-term datasets which do not presently exist (Willis et al., 2018). While the needed time to produce such datasets is scarce, the paleosciences provide a lens with which we can build predictive models using fossil fungal assemblages. These assemblages occur in two major forms (Taylor et al., 2015): macrofossils, which may preserve significant portions of the fungal organism in exquisite detail, and microfossils, often fragmented or representative of a single phase of the life cycle of fungus, typically the spore, including the conidiophores, conidium, ascospore, and basidiospore, or other structures such as hyphae, conidiophores, rhizoids, etc.

Fungal microfossils are generally studied under the larger discipline of palynology. They are classified as non-pollen palynomorphs (NPPs), which have increasingly been used as paleoecological indicators in both Quaternary and deeper time studies (van Geel and Apte, 2006; Nuñez Otaño et al., 2021; Shumilovskikh et al., 2021; among others). Historically, fossil fungi in deep-time have been identified using an artificial classification scheme based on morphological descriptions that have little to no correlation with modern taxa (O’Keefe et al., 2021). This classification makes it impossible to apply the Coexistence Approach to fossil fungi (Utescher et al., 2014; Grimm and Potts, 2016), which is widely used with other palynomorphs to elucidate ancient climates. Recent advances in training and collaborations with modern mycologists are permitting the discipline to move past this limitation and to recognize members of modern families and genera in the fossil record (Nuñez Otaño et al., 2017, 2021; O’Keefe, 2017; Shumilovskikh et al., 2017; Pound et al., 2019; Bianchinotti et al., 2020; Loughlin et al., 2021).

The FiaWW (Fungi in a Warmer World) project is one such collaboration, bringing together palynologists, mycologists, and climate scientists. It has turned its lens upon the middle Miocene Climatic Optimum (MMCO), the warmest interval of the last 23 million years and an internationally recognized possible analog for the Intergovernmental Panel on Climate Change -Representative Concentration Pathway (IPCC-RCP) 4.5–8.5 warming scenarios, both in terms of intensity and rapidity of warming and in terms of latitudinal effects (Intergovernmental Panel on Climate Change [IPCC], 2014, 2021; Steinthorsdottir et al., 2021). Using fungal palynology, we aim to construct the first large-scale, long-term fungal diversity and worldwide distribution dataset of its kind to build a predictive model for fungal assemblage behavior under future warming scenarios.

As an initial test of the feasibility of the project, we examined low-rank coal samples from two regions, west-central Slovakia and northern Thailand, for fungal palynomorph content. Low-rank coals were chosen specifically for this initial test of concept to (1) control for depositional environment (O’Keefe et al., 2013; Dai et al., 2020), (2) minimize the effects of transport, as organic matter preserved in both systems is predominantly autochthonous (produced from standing biomass and preserved in situ) (O’Keefe et al., 2011; Belkin et al., 2013), and (3) minimize the impact of coalification on the fungal material (Dai et al., 2017, 2020). The two regions presently fall into warm-summer and humid continental climate (Slovakia), as well as tropical wet and dry climate (Thailand) classifications. They appear to have similar climatological settings at the onset of middle Miocene warming (Steinthorsdottir et al., 2021), and warmed unevenly, with temperate Slovakia experiencing relatively greater warming than tropical Thailand, as is anticipated for future warming scenarios (Intergovernmental Panel on Climate Change [IPCC], 2014, 2021). Of note, coals from all samples have similar organic and geochemical compositions and are interpreted as having been deposited in a mosaic of topogenous and ombrogenous mire settings (O’Keefe et al., 2011; Belkin et al., 2013). The topogenous regions are dominated by coniferous trees from the family Cupressaceae and aquatic to emergent herbaceous plants, including Haloragaceae, Cyperaceae, and Poaceae, while the ombrogenous regions are a mixture of hardwood and softwood trees and shrubs with an understory of ferns, grasses, and sedges (Songtham et al., 2005; Kučerová, 2010; Sepulchre et al., 2010; Hudáčková et al., 2020).

Preliminary studies of fungal remains preserved in the coals from Slovakia were carried out by O’Keefe and Hower in the period 2007–2011 (O’Keefe et al., 2011). These studies report only 23 morphotypes for the Slovakian material (O’Keefe et al., 2011). For Thailand, there are no previous studies about the fossil fungal assemblages during the middle Miocene. Here, we present a re-study of the Slovakian material and a first study from coals from Thailand to expand the fossil fungi record in both regions. This study was designed to assess fungal communities and paleoenvironments from the middle Miocene between Thailand and Slovakia. We also wanted to test if fungi were more abundant and diverse in the tropics than in the temperate regions during this period (Hyde et al., 2004; Tedersoo et al., 2014; Chaudhary et al., 2018; among others) or if its pattern of distribution differed from those from plants and other organisms (Rudolph et al., 2018; Větrovský et al., 2020). This study will provide a foundation for the main goal of the FiaWW project, which is MMCO climate reconstruction using fungi as reliable proxies.

MATERIALS AND METHODS

Study Area

We selected a total of 13 samples from the lower to middle Miocene, including the MMCO, five from Thailand and eight from Slovakia. These are part of previous collaborative research with the United States Geological Survey during the periods 2000–2005 and 2010–2015. Samples from Thailand are from multiple mines in the northern part of the country, while samples from Slovakia are from the west-central portion of the country (Figure 1 and Table 1).
FIGURE 1 | Maps of study areas indicate the regions in Slovakia (red) and Thailand (blue) where the coalfields are located. The gray points indicate the coal basins where the samples for this study were extracted. The regional maps indicating the coal fields and coal basins from Thailand and Slovakia were redrawn and modified from Machajová et al. (2000) and Ratanasthien (2011).

**Thailand Stratigraphy**

The regional structure of the geology of Thailand is complicated because the basin fill was generally deposited in variably subsiding rift basins and neotectonic overprints (Morley and Racey, 2011). Basin fills represent a mosaic of lowland environments developed atop basement rock and basin margin conglomeratic alluvial fans. Thus, rocks deposited in these environments range from fluvial sandstones with associated finer-grained overbank deposits to lacustrine mudstones and oil shales, and to mire-associated claystones and lignitic coal seams (Morley and Racey, 2011; Friederich et al., 2016). Considering that many of the basins in Thailand are both sources of coal and hydrocarbons and paleontologically important, significant data exists on intra-basin stratigraphy, biostratigraphy, magnetostratigraphy, and plant- and animal-based paleoecology (Benammi et al., 2002, 2010; Ratanasthien, 2002; Songtham et al., 2005; Ratanasthien et al., 2008; Thasod et al., 2008; Coster et al., 2010; Sepulchre et al., 2010; Friederich et al., 2016; Grohé et al., 2020; among others). This makes it possible to organize individual coal seams that occur within the basins by age. The lignites selected for this study were from the coalfields of Mae Moh, Nae Hong, Mae Lai, and Chiang Muan (Figure 1), and were deposited in the lower and middle Miocene (Figure 2; Tewalt et al., 2005; Belkin et al., 2013; World Coal Quality Inventory, 2019).

**Slovakia Stratigraphy**

Slovakian lignites examined for this study are from the Novaky and Handlova regions of the Upper Nitra Basin in the

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**TABLE 1 | Geological information of the samples from Thailand and Slovakia analyzed in this study.**

| Locality | Coalfield | (Member, where known) Coal | Age | Sample |
|----------|-----------|--------------------------|-----|--------|
| Thailand | Chiang Muan | Lower Main Bed | 12.4–14 | 453 |
| Thailand | Mae Lai | Hod Lignites | Middle Mocene | 454 |
| Thailand | Mae Moh | J Coal | 12.18–12.4 | 455 |
| Thailand | Mae Moh | Q Lignite | 13.13–13.3 | 456 |
| Thailand | Na Hong | Chiang Mai Lignites | Lower Mocene | 457 |
| Slovakia | Novaky | (Alive) Hlboky Slaj | Middle Mocene | 3 |
| Slovakia | Handlova | (Handlovak) Hlavny Slaj | Middle Mocene | 4 |
| Slovakia | Handlova | (Handlovak) Hlavny Slaj | Middle Mocene | 5 |
| Slovakia | Handlova | (Handlovak) Hlavny Slaj | Middle Mocene | 6 |
| Slovakia | Handlova | (?Handlovak) Podlozne Slaj | Middle Mocene | 7 |
| Slovakia | Novaky | (Novaky-Handlovak) Hlavny Slaj | Middle Mocene | 8 |
| Slovakia | Novaky | Novaky-Handlovak Hlavny Slaj | Middle Mocene | 9 |
| Slovakia | Novaky | Novaky-Handlovak Hlavny Slaj | Middle Mocene | 10 |
Carpathians (Figure 1) (Hók et al., 1995; Dill et al., 2008). Coals in this region were deposited during the middle Miocene (Figure 2; Sitár et al., 1987; Planderová, 1991). Like those in Thailand, the coals studied in Slovakia developed in a pull-apart basin produced during extensional tectonics (Hók et al., 1995; Dill et al., 2008). The Upper Nitra Basin is among the smaller basins bounded by normal faults in the western Carpathians (Hók et al., 1995), and has been a target of coal, oil and gas exploration and production. Interestingly, the stratigraphy within the two coal mining regions, which occur in the Kamanc and Nováky Formations, is not well understood (Machajová et al., 2002; Dill et al., 2008). Most authors agree with Hudaček et al. (1982) that the main seam (Hlavný slaj) lignites mined near Nováky and Handlová are probably the same seam, although little evidence is presented to support this, beyond the general stratigraphy of both mining areas. As in Thailand, the seams are developed in a mosaic of terrestrial siliciclastic environments in the Kamanc and Nováky Formations, ranging from fluvial-alluvial to lacustrine in internally-draining grabens with coarse-grained alluvial fans along the margins (Čechovič, 1950). The majority of sediment sources appear to be weathered extrusive igneous rocks from the adjacent proto-Carpathians (Čechovič, 1950; Machajová et al., 2000). The lowermost lignite in the region (Hlboký slaj and Novaký), as noted by Hudaček et al. (1982), is tentatively correlated with the “Alive” member sample SK-1 in our studied set; the majority of samples, however, come from the main seam (Hlavný slaj, Handlová produktivne súvrstvie; Novaký Fm.) or its underlier, the Podlozne slaj (Figure 2; Hudaček et al., 1982; Machajová et al., 2000, 2002). In the Handlova mining area, the main seam has two splits that are most greatly separated (by up to 50 m) in the eastern and northeastern portion of the area by a siliciclastic parting that tapers to nothing in the W and SW portion of the mining area (Machajová et al., 2000). The coal zone is overlain by dark gray fossil plant-bearing clays of the Serravallian-age (late Badenian) Koš Formation (Kučerová, 2010).

Paleoclimate

Plant-based reconstructions from coal basins in northern Thailand have indicated a transition from warm-temperate climates in the early Burgidalian (Chiang Mai lignites) to tropical climates by the late Burgidalian-early Langhian (Hod lignites) that largely persisted through the early middle and upper middle Miocene (Ratanasthien et al., 2008; Morley and Racey, 2011). It has been suggested that these changes coincide with increasing paleotopographic relief and intensification of the SE Asian monsoon (Morley, 2012; Steinthorsdottir et al., 2021).

In west-central Slovakia, Miocene paleobotanical reconstruction indicate that tropical to sub-tropical conditions gave way to highly seasonal warm-temperate to temperate conditions by the late Langhian, with few thermophilus plant taxa retained, even in coal-forming basins (Kvaček et al., 2006).
During this time, paleotopographic relief first increased, then decreased slightly in relation to tectonism associated with development of the western Alps and Carpathians (Kvaček, 1994; Steinthorsdottir et al., 2021).

**Palynology Methods**

We extracted fungal palynomorphs using HF-free protocols outlined for low-rank coal in O’Keefe and Eble (2012); Pound et al. (2021), and Riding (2021). These consisted of crushing the coal to −1 mm, disaggregation in non-phosphatic anionic detergent (Liquinox®), a treatment with hypochlorous acid to de-polymerize lignin, and a treatment with 5% KOH to produce water-soluble organic salts. The salts were then rinsed away before to subject the samples to heavy-density separation using LST® Heavy Liquid (a proprietary lithium-containing heteropolytungstate compound). After this, they were placed in 95% ethanol prior to mounting. We mounted individual coverslips of the palynological residues in polyvinyl

**FIGURE 3** | Fungal spores found in Thailand. Each photograph indicates: identification if any, the sample number, morphotype number (OPaL #), and England Finder coordinates (EF). (A) Arecophila sp., sample 457, OPaL 36, EF: N49-3; (B) cf. Canalisporium sp. 2, sample 455, OPaL 140, EF: P26-1; (C) cf. Asterina, samples 453, OPaL 125, EF: Q32-3; (D) Diporotheca, sample 457, OPaL 141, EF: Y35-2; (E) sample 453, OPaL 106, EF: L38-1, T38-3; (F) cf. Chaetomium, sample 457, OPaL 87, EF: N50-2; (G) cf. Cercophora, sample 455, OPaL 92, EF: F24-4; (H) sample 456, OPaL 33, EF: S26; (I) Thiolothecium propagules (Germling), sample 454, OPaL 102, EF: K37-2; (J) sample 453, OPaL 88, EF: G35-2, F35-4; (K) Oncopodiella aff. trigonella, sample 454, OPaL 96, EF: L26-4; (L) Megalohypha aff. aqua-dulces, sample 453, OPaL 97, EF: T37-4; (M) cf. Xylomyces sp., sample 456, OPaL 152, EF: W24-2; (N) Tetraploa, sample 454, OPaL 98, EF: AA34. Scale bar: 10 µm. More information of each specimen in Supplementary Table 1.
Fungal spores found in Slovakia. Each photograph indicates: identification if any, the sample number, morphotype number (OPaL #), and England Finder coordinates (EF).

(A) sample 10, OPaL 77, EF: Q49-2; (B) sample 10, OPaL 227, EF: T39-3; (C) sample 3, OPaL 243, EF: J69-1; (D) Jahnula, sample 4, OPaL 278, EF: F-33; (E) cf. Dictyosporium, sample 4, OPaL 274, EF: O41; (F) Glomeraceae (cf. Glomus), sample 10, OPaL 226, EF: X46-1; (G) cf. Asterina, sample 4, OPaL 229, EF: D34; (H) sample 6, OPaL 78, EF: N70-1; (I) sample 3, OPaL 250, EF: E85-4; (J) Ascotaiwania aff. limnetica, sample 10, OPaL 114, EF: N39-3; (K) Diporotheca, sample 5, OPaL 141, EF: M38; (L) cf. Tolstoma, sample 10, OPaL 209, EF: N40-4; (M) sample 10, OPaL 230, EF: W42-4; (N) sample 10, OPaL 195, EF: C46-2-4; (P) sample 3, OPaL 250, EF: W41-2; (Q) sample 4, OPaL 194, EF: V32-3. Scale bar: 10 µm. More information of each specimen in Supplementary Table 1.

alcohol (PVA). We examined the specimens on Leitz and Leica microscopes with variable magnifications from 10 to 1,000×. We obtained z-stack photomicrographs at 1,000× magnification, primarily with Leica ICC50W cameras and using the Leica Application Suite® (LAS) software. We then z-stacked the multiple focal planes using Helicon Focus® software. We imaged a minimum of one fungal palynomorph for each morphotype recorded, and more than five when possible. Fungal palynomorphs were consecutively numbered using the acronym for the laboratory name, OPaL # (O’Keefe Paleoecology Lab).

Fungal Identification

We separated the morphotaxa using the following criteria: shape, ornamentation, color, color variation, number and type of apertures, and/or attachment scars (Hughes, 1953; Pirozynski and Weresub, 1979; Kalugtkar and Jansonius, 2000; Gulis et al., 2005; Seifert et al., 2011; Guarro et al., 2012; Nuñez Otaño et al., 2021). When possible, we identified these taxa to family or genus using the above references and primary literature for each modern taxon (Supplementary Table 1). Ecological information was obtained using available literature about each taxon. FunGuild (a Python-based tool) was used for ecological guild data (Nguyen et al., 2016; Supplementary Table 2) and biogeographic distribution was complemented with Global Fungi database (2021; Větrovský et al., 2020).

Data Analysis

We used presence/absence data to record the diversity observed between Thailand and Slovakia and among samples. We did a boxplot diagram and a Kruskal–Wallis test to compare richness between countries. This test compares two groups without assuming them to follow a normal distribution (McDonald, 2014). The morphotypes were first separated into 10 categories based on shape variation (Seifert and Gams, 2011; Nuñez Otaño et al., 2021). The morphological categories identified were:
amerospore, didymospore, dictyospore, phragmospore, bulbil-like spore, epiphyllous structures, clathrate spore, helicospore, staurospore, and indeterminate fungal remains. We calculated the percentage of each morphological category in each sample and plotted them in bar plots to show morphological variation among the samples for both countries.

We used the Shannon–Wiener Index ($H'$) to calculate $\alpha$-diversity to examine variation among samples and between sites based on morphological variability. For the indices, we used the morphological categories as taxa and the number of specimens per group in each sample was used as the total number of individuals. We used principal component analysis (PCA) to compare morphology-environment variation (Nuñez Otaño et al., 2021) among samples within localities and between localities. Although the data collected were presence/absence, the analysis was applied to the morphological groups, and PCA was selected because it calculates Euclidean distances between samples (Legendre and Legendre, 2012). We also calculated all the pairwise similarities of our PCA using Euclidean distance to compare the samples. In addition, we completed a separate PCA that excluded amerospores to evaluate how the distances vary in the absence of this group. This is justifiable because the majority of amerospores do not possess sufficient spore traits to permit unambiguous identification in the absence of the rest of the fungus, thus obscures signals in the data in much the same way as widely-disseminated anemophilous pollen (Birks and Birks, 1980). The analyses were carried out in R (R Core Team, 2021).

**Pollen-Based Climatological Signals**

For a better interpretation of the fossil fungal assemblages found in our samples, we ran an additional analysis using CREST (Climate Reconstruction Software) (Chevalier et al., 2014) to reconstruct terrestrial paleoclimates of the study areas using published palynological records. CREST has been shown to provide suitable quantification of error in middle Miocene terrestrial climate reconstructions and is consistent with Co-existence Approach (Utescher et al., 2014) reconstructions from northwest Europe (Gibson et al., in review). We extracted taxa lists and nearest living relatives from the literature and used as the input for CREST, obtained from Nováky (Kučerová, 2010) and the Novohrad-Nőgrád Basin (Hudáčková et al., 2020) for Slovakia and Mae Moh (Songtham et al., 2005) and Chiang Muan (Sepulchre et al., 2010) for Thailand. CREST reconstructions were carried out in R (R Core Team, 2021).
variables were assessed in detail: Mean Annual Temperature (MAT), Mean Temperature of Warmest Quarter (MTWQ), Mean Temperature of Coldest Quarter (MTCQ), Mean Annual Precipitation (MAP), and Precipitation Seasonality (CoV x100). The CREST output provides 50 and 95% (2-σ) uncertainties as well as an optimum. Here we focus on the optima and 95% confidence interval for each variable. All analyses were carried out in R (R Core Team, 2021).

RESULTS

Fossil Fungi
A total of 637 fungal spore specimens were recorded in the samples from Thailand and Slovakia (Supplementary Table 1 and Figures 3–5). These were grouped into 281 morphotaxa, based on detailed morphological comparisons, of which 151 were found only in Thailand, 93 only in Slovakia, and 37 in both Thailand and Slovakia. The fungal diversity of the samples is mostly represented by Ascomycota, but we also found few spores from other phyla, such as Basidiomycota (OPAL 219) and Glomeromycota (OPAL 226) (Figures 3–5). Overall, richness between Thailand and Slovakia is different (Kruskal–Wallis test, $P = 0.02$), in which the samples from Thailand had a higher number of morphotaxa (mean: 48.8; SD: 16.8) than Slovakia (mean: 21.75; SD: 14.8) (Figure 6). Amerospores and phragmospores were present in all samples, except sample 7 (Figure 7). Reviewing the variation of abundances, amerospores were the dominant group in most of the samples from both countries (Figure 7). In Thailand, the dominance of ameromorphotypes varied from 32% (sample 457; Figure 7A) to 75% (sample 454; Figure 7A), while in Slovakia varied from 45.2% (sample 3; Figure 7B) to 100% in (sample 7; Figure 7B). However, sample 7 was the sample with the lower record of fungi spores, in which only 3 spores count for the 100% of amerospores present in this sample.

In the PCA of all morphologies, the two first axes explained 95% of the variation (Figure 8 and Supplementary Table 3). In Axis 1, amerospores drive the variation of the samples, especially sample 454 from Thailand, while in Axis 2, phragmospores drive this variation, especially sample 457 from Thailand (Figure 8). Most of the Slovakian samples were grouped close together, except for samples 3 and 10, while the samples from Thailand were more dispersed (Figure 8).

When amerospores were excluded from the PCA, there was a clear separation of the samples by country and within samples depending on the fungal community composition and based on morphotypes and fungal spores identified (Figure 9 and Supplementary Table 2). We also observed that the samples were grouped in a temporal scale, in which 457, from the lower Miocene from Thailand, was different from most of the samples (Figure 2 and Supplementary Table 3). Samples 453 (13.5–16 My) and 456 (13.3–13.1 My), from Thailand, clustered together in the PCA (Figure 9). These samples are temporally close, and their fungal composition suggest similar environments (Figure 2 and Supplementary Table 2). All samples from Slovakia were grouped together (Figure 9), and in the temporal scale these samples are in the same time interval (middle Miocene, Figure 2) and have also similar fungal composition (Supplementary Table 2). At the same time, the Slovakian samples were clustered with Thailand 455, located in the middle Miocene and post MMCO (Figures 2, 9).

Based on the shape variation of the spores, we found that the samples from Thailand had a higher morphological richness than Slovakia (Figure 6). Among the samples from Thailand, α diversity was higher in samples 457 ($H': 1.56$; lower Miocene) and 456 ($H': 1.53$; middle Miocene). Among those from Slovakia, α diversity was higher in samples 3 ($H': 1.36$; MMCO) and 5 ($H': 1.42$; middle Miocene). The sample with the lowest diversity was sample 7 from Slovakia ($H': 0.0$; middle Miocene). Sample 457 had the highest morphological variation, containing nine of the 10 morphological groups; it contained no staurospores (Figure 7A). Sample 7 contained only three amerospores (Figure 7B).

Samples 457 (Thailand) and 5 (Slovakia) were the only samples in which amerospores were not dominant (Figure 7). In sample 457, phragmospores were the dominant morphology (39.6%) followed by amerospores (32.1%, Figure 7A). In sample 5, didymospores were the most abundant (31.3%), followed by phragmospores (25%) and amerospores (21.3%) (Figure 7B). For Thailand, some of the spores identified in 457 were: Catenularia aff. catenulata and Phaeosclera-type (Figure 5 and Supplementary Tables 2, 3). Sample 454 followed sample 457 in...
morphological variation with spores from eight morphological groups (Figure 7A). In this sample, amerospores represented 75% of the variation, while the other morphologies accounted for the remainder (Figure 7A). Some of the spores identified in 454 were cf. Ascobolaceae, cf. Catenularia sp., Oncopodiella aff. trigonella and Tetraploa (Figures 3, 5 and Supplementary Table 2). Samples 453 and 456 have representatives of five and six morphological groups, respectively (Figure 7A). Spores
found only in 453 include *Megalohypha* aff. *aqua-dulces*, while in 456 we found cf. *Berkleasmium* (Figures 3, 5 and Supplementary Tables 2, 3). From Thailand, sample 455 had the lowest morphological variability with only four morphological groups and contained identifiable spores of cf. *Arthrinium, Ascothaiwania* aff. *limnetica*, and *Monosporites* sp. (Figures 3, 5, 7A and Supplementary Table 2).

In Slovakia, sample 4 had the highest morphological variations with six groups, and we observed spores of *Jahnula* and cf. *Cercophora* (Figures 4, 7B and Supplementary Table 2). Samples 3, 5, and 10, each had five morphological groups (Figure 7B). Samples 6 and 9 each had 4 groups (Figure 7B), while sample 8 had only three morphological groups (Figure 7B). Spores unique to these samples include *Jahnula* in sample 4 and cf. *Monotosporella* in sample 8 (Figures 4, 5 and Supplementary Table 2).

Among the samples from Thailand, spores identified in more than one sample were *Arecophila* sp., cf. *Canalisporium*, and *Phragmocephala* (Figures 3, 5 and Supplementary Table 2). In Slovakia, we found cf. *Dictyosporium*, Glomeraceae, cf. *Delitschia*, and cf. *Tulostoma* in several samples (Figures 4, 5 and Supplementary Table 2). Several spores also occurred in both localities, including cf. *Rosellinia*, cf. *Canalisporium*, Thyriothecium propagules, cf. *Asterina*, and *Diporotheca* (Figures 3–5 and Supplementary Table 2).

### Pollen-Based Miocene Climate of Thailand and Slovakia

For the Thailand sites (Table 2), CREST reconstructs a mean annual temperature of 26.3°C (19.2–27.9°C) for Mae Moh with a cooler MAT reconstruction for Chiang Muan of 18.7°C (12.7–27.3°C) (Table 2 and Supplementary Figure 1). Summer temperatures at both sites are comparable with the MTWQ reconstructed at 26.9°C (25.9–29.8°C) for Mae Moh and 26.4°C (20.0–30.8°C) for Chiang Muan (Table 2 and
Supplementary Figure 2). For winter temperatures, MTCQ is reconstructed at 24.6°C (16.3–28.1°C) for Mae Moh but considerably cooler at 10.9°C (3.8–24.6°C) for Chiang Muan (Table 2). While MAP at Mae Moh [2,332 mm (796–4,323 mm)] is approximately double Chiang Muan [967 mm (455–3,299 mm)] (Table 2), but precipitation seasonality is very similar, reconstructed at 56.4 (20.5–112.8%) for Mae Moh and 61.5 (17.1–114.5%) for Chiang Muan (Table 2).

For the Slovakia sites (Table 2), CREST reconstructs a MAT of 10.5°C (6.2–19.7°C) for both Nováky and the Novohrad-Nógrád Basin (Supplementary Figure 1 and Table 2). Summer and winter temperatures at the two sites are comparable with a MTWQ reconstructed at 19.0°C (14.5–27.3°C) for Nováky and 18.0°C (14.1–26.9°C) for the Novohrad-Nógrád Basin (Supplementary Figure 2 and Table 2), and a MTCQ reconstructed at 3.2°C (−5.7–12.1°C) and 3.8°C (−5.1–12.1°C) respectively (Table 2). The MAP for both sites is the same at 967 mm (398–2,105 mm) (Table 2), and precipitation seasonality for Nováky [967 mm (455–2,105 mm)] is lower than for the Novohrad-Nógrád Basin [20.5 (10.3–87.1%)] (Table 2).

**DISCUSSION**

This study tried to describe fossil fungal communities from temperate and tropical regions following a trait-based approach to fungal community. Our results showed, after PCA analyses using morphotaxa, that middle Miocene fungal communities from Thailand and Slovakia may have had similar ecological characteristics and/or responses to those past environments (Figures 8, 9). Spore morphology may be related to ecological requirements due to selective pressures, affecting release pathways, transport, survival rate and deposition (Nguyen et al., 2016; Nuñez Otaño et al., 2021). Similarities between
TABLE 2 | Palaeoclimate reconstruction for Nováky, Novohrad-Nógrád Basin, Mae Moh, and Chiang Muan.

| Locality       | MAP (°C) | Optima | 50% | 95% | MTWQ (°C) | Optima | 50% | 95% | MTCQ (°C) | Optima | 50% | 95% | Precipitation (mm) | Precipitation seasonality (CoV) |
|----------------|----------|--------|-----|-----|-----------|--------|-----|-----|-----------|--------|-----|-----|-------------------|-------------------------------|
| Slovakia       | 10.5     | 8.9–13.2 | 6.9–19.7 | 19.0 | 17.0–21.1 | 14.5–27.3 | 3.2 | 0.6–9.6 | 6.2–17.2 | 967    | 729–1318 | 729–1318 | 238–2105 | 17.1 | 12.0–27.3 |
| Novohrad-Nógrád Basin | 10.5 | 8.9–13.2 | 6.9–17.2 | 19.0 | 16.5–20.5 | 14.1–28.9 | 3.2 | 0.6–5.6 | 6.2–12.2 | 967    | 729–1318 | 729–1318 | 449–1205 | 20.5 | 10.3–37.4 |
| Thailand       | 26.3     | 24.6–26.8 | 19.2–27.9 | 26.9 | 25.9–27.3 | 22.9–29.8 | 24.6 | 22.8–25.8 | 16.9–28.1 | 2332   | 1139–2729 | 796–4303 | 150–1356 | 9.4  | 27.9–82.0 |
| Mae Moh        | 18.7     | 16.5–21.9 | 12.7–27.3 | 26.4 | 24.4–27.3 | 20.7–30.8 | 10.9 | 8.6–16.3 | 3.9–24.6 | 967    | 729–1479 | 729–1479 | 455–5699 | 61.5 | 42.7–78.6 |
| Chiang Muan    | 26.3     | 24.6–26.8 | 19.2–27.9 | 26.9 | 25.9–27.3 | 22.9–29.8 | 24.6 | 22.8–25.8 | 16.9–28.1 | 2332   | 1139–2729 | 796–4303 | 150–1356 | 9.4  | 27.9–82.0 |

Optima, 50 and 95% confidence intervals were generated using CREST (Climate REconstruction SofTware) (Chevalier et al., 2014). MAT, mean annual temperature; MTWQ, mean temperature of warmest quarter; MTCQ, mean temperature of coldest quarter; MAP, mean annual precipitation.

Most of our samples presented amerospores as an important component of the total fungal richness along with phragmospores (Figure 7). Amerospores or rounder spores may indicate a humid environment with structurally closed vegetation and are more related to saprophytic fungi as litter specialist, having better air dispersal pathways and reaching longer distances (Roper et al., 2008; Calhim et al., 2018; Crandall et al., 2020). Otherwise, larger and more allantoid spores could be found frequently on wood and herbs, probably in sites with drier vegetation types (Calhim et al., 2018; Crandall et al., 2020) having more affinity to water dispersal (Magyar et al., 2016). The presence of aero-aquatic fungi recognized throughout spore shapes, such as stauropores, helicospores and bulbil-like spores, allow us to add biotic and abiotic characteristics from freshwater habitats to our paleoenvironmental reconstructions in Thailand and Slovakia in the Miocene (Ingold, 1975; Nuñez Otaño et al., 2021). These results suggest a freshwater setting with high rainfall rates for our study sites and coincided with the CREST paleoclimatic reconstructions based on paleobotanical records (Table 2). These results also suggest the presence of a more structured forest with herbaceous layers, aquatic plants, and hypoxic conditions (Tetraploa), among others (Supplementary Table 2 and Supplementary Figures 1, 2; Schoenlein-Crusius and Piccolo Grandi, 2003; Zhao et al., 2007; Chauvet et al., 2016; Magyar et al., 2021; Nuñez Otaño et al., 2021). Modern epiphyllous fungi have been found in warm, humid subtropical to tropical regions around the world with high annual rainfall and air moisture and in paleontological research works these findings are correlated with warm temperate to subtropical and moist mixed angiosperm forest (Dilcher, 1965; Selkirk, 1975; Johnson and Sutton, 2000; Reynolds and Gilbert, 2005; Limaye et al., 2007; Hofmann, 2010; Piepenbring et al., 2011; Bannister et al., 2016; Bianchinotti et al., 2020).

Recent mycological researchers are trying to describe fungal spore diversity related to fungal community structure and vegetation patterns (Pringle et al., 2015; Glassman et al., 2016; Magyar et al., 2016, 2021; Crandall et al., 2020; Nuñez Otaño et al., 2021; among others). In this study our results show that spore morphology can give us more information about paleofungal assemblages than previously thought. And after adding identified fungal species, we found a clearer pattern of fungal distribution related with local biotic and abiotic characteristics (see section “Paleoenvironments of Thailand and Slovakia Based on Fungal Composition”).

Paleoenvironments of Thailand and Slovakia Based on Fungal Composition

Based on fungal composition and morphological variation, the environments for the Burdigalian of Thailand and Langhian and Serravallian of Thailand and Slovakia were:
In the Burdigalian of Na Hong Thailand (sample 457), the fungal community is composed by clathrate spores, bulbil-like spores, helicospores, and epiphyllous structures (Figure 7A), usually associated to local humid and warm conditions and freshwater habitats (Ingold, 1975; Zhao et al., 2007). Meanwhile, the presence of genera, such as *Catenuariella* aff. *catenulata*, *Arcetaphila* sp. and *Diporotheca* (Supplementary Tables 2, 3), suggests a temperate to tropical environment with freshwater habitats (Umali et al., 1999; Hawksworth and Lücking, 2017; Réblova et al., 2021). Previous paleobotanical and paleoecological studies indicated similar paleoenvironments for the lower Miocene of Na Hong with warm-temperate forest and *Taxodium* swamp with fern-sedge communities (Songtham et al., 2005; Sepulchre et al., 2010).

In the Langhian MMCO, the fungal composition of Mae Lai Thailand (sample 454) indicates tropical to subtropical environments with freshwater habitats and local humid conditions, based on the presence of helicospores, bulbil-like spores, stauropores, and epiphyllous structures such as Thyriothecium propagules (Figure 7A), and the presence of *Tetratlopa*, *Oncomodiella* aff. *trigonella*, and cf. *Catenuariella* sp. (Supplementary Tables 2, 3; Field and Webster, 1985; Bärlocher, 2009; Chauvet et al., 2016). In Slovakia, during the same period (sample 3), the presence of bulbil-like spores, and spores of cf. *Canalisporium*, cf. *Delitschia*, and *Savoryella* aff. *fusiformis* indicates a tropical to subtropical environment with low energy wetlands (i.e., swamp or lake, Figure 7B). The CREST models for Slovakia indicate similar paleoclimatic conditions with warm-temperate climates MAT of 10.5°C (6.2–19.7°C at 95% confidence), and high precipitation MAP of 967 mm (398–2,105 mm) (Supplementary Table 2 and Supplementary Figures 1, 2). Thus, our fungal records and paleoclimatic reconstructions support previous paleobotanical works that indicated that during the early and middle Miocene Thailand (Mae Lai) was a paratropical forest with warm and humid conditions (Yabe, 2002; Kunimatsu et al., 2005), and Slovakia had warmer and wetter conditions that present day (Kvaček et al., 2006; Hudáčková et al., 2020).

During the Langhian and post MMCO, the fungal composition of Thailand (Chiang Muan sample 453) suggests a tropical to subtropical environment and freshwater habitats with the presence of *Megalohypha* aff. *aqua-dulcis*, cf. *Asterina*, cf. *Xylomyces* sp., cf. *Dictyosporium* and cf. *Sordaria* (Figure 7A and Supplementary Tables 2, 3) (Goh et al., 1997; Hyde and Goh, 1997; Ferrer et al., 2007; Hosagoudar and Riju, 2013; Hofmann and Piepenbring, 2014; Shumilovskikh et al., 2017; Yang et al., 2018). The CREST models indicated similar climatic conditions for Chiang Muan with a MAT of 18.7°C (12.7–27.3°C at 95% confidence) and a MAP of 967 mm (455–3,299 mm at 95% confidence), suggesting a subtropical and modestly humid climate (Supplementary Table 2 and Supplementary Figures 1, 2). Studies in this region have indicated tropical woodlands and peat swamp forests with fern-sedge communities (Songtham et al., 2005; Sepulchre et al., 2010). For Slovakia (sample 7), we were not able to get paleoecological inferences for the same period because the sample was fungal palynomorph-poor and contained only three amerospore types (Figure 7B and Supplementary Table 1). Based on previous published organic petrography and geochemistry, sample 7 likely represents an area of root-dominated peat produced by plants in an ombrogenous setting (O’Keefe et al., 2011). Previous paleo reconstructions for the late Langhian of Slovakia have indicated environments with highly seasonal warm-temperate to temperate conditions by the late Langhian for Nováky and the Novohrad-Nógrád Basin (Kvaček et al., 2006; Hudáčková et al., 2020).

In the early Serravallian from Thailand (sample 456 from Mae Moh), the presence of dicots, bulbil-like spores, helicospores and didymosporic, as well as the record of cf. *Canalisporium*, *Phragmoechta*, and cf. *Berkleasniun* (Supplementary Table 2 and Figures 3, 5, 7A) suggest a tropical environment and humid local conditions, with forests and freshwater habitats (Holubová-Jechová, 1987; Yip, 1988; Chen and Tzean, 2010; Zhao et al., 2012; Hüseyin et al., 2014; Su et al., 2015). These results concur with the paleoclimatic reconstructions for Mae Moh that indicate tropical and wet environments during the Serravallian [MAT: 26.3°C (19.2–27.9°C at 95% confidence) and a MTCQ of 24.6°C (16.3–28.1°C at 95% confidence); MAP: 2,332 mm (796–4,323 mm at 95% confidence)] (Supplementary Table 2 and Supplementary Figures 1, 2). Other studies have indicated a warm tropical peat swamp with fern-sedge vegetation, and in the uplands a subtropical warm temperate mixed forest with *Pinus* and grasses (Songtham et al., 2005; Sepulchre et al., 2010).

Some of the fungi identified in Slovakian samples that were deposited in the Serravallian, in Hlavný slaj (samples 4–6 and 8–10, Figure 2) were cf. *Monotospora*, *Jahnula*, cf. *Tulostoma*, cf. *Dictyosporium*, and Thyriothecium propagules, among others (Figures 4, 5 and Supplementary Table 2). The modern ecology of these spores indicates tropical to temperate environments with humid conditions and freshwater habitats (Goh et al., 1999; Sri-indrasutdi et al., 2010; Cortez et al., 2011; Sadowski et al., 2012; Jeppson et al., 2017; Yang et al., 2018). The Paleoclimatic reconstructions for Slovakia using CREST suggest a warm-temperate climate with high precipitation [MAT: 10.5°C (6.2–19.7°C)]; MAP: 967 mm (398–2,105 mm)] (Supplementary Table 2 and Supplementary Figures 1, 2). Preliminary studies for the Serravallian of Slovakia have indicated warm-temperate climates persisted through this time in the upper Horna Nitra Basin (Hudáčková et al., 2020). Additionally, samples 4, 6, and 10 are from the base and roof of the seams, where ombrogenous conditions had yet to develop or were giving way as peat production ceased (Hudáček et al., 1982; Machajová et al., 2000, 2002; Dai et al., 2020).

The fungal composition of sample 455 from Mae Moh, Thailand includes *Arcetaphila* sp., cf. *Arthrinium*, cf. *Canalisporium*, *Savoryella* aff. *fusiformis*, cf. *Rosellinia*, and *Ascolatiauina* aff. *limnetica* (Figures 3, 5 and Supplementary Table 2). The modern ecology of these genera suggests a tropical to temperate environment with freshwater habitats similar to the paleoenvironments of the Serravallian samples from Slovakia (Umali et al., 1999; Petrini, 2003, 2013; Songtham et al., 2005; Zhao et al., 2007; Sepulchre et al., 2010; Chauvet et al., 2016). This also explains the clustering of this sample with the Slovakian samples in the PCA
Paleoenvironments and Paleoclimate During the Middle Miocene

The composition and diversity of fungal communities seems to be influenced by climate, affecting ecosystem functioning (Větrovský et al., 2020). This has been observed in several studies of modern fungal biogeographic distributions (Tedersoo et al., 2014; Bahram et al., 2016; Daws et al., 2020). Our results seem to reflect similar fungal community composition under similar environments in different countries (Figures 8, 9), showing an apparent correlation with the paleoclimate. The paleoclimate reconstructions made for the middle Miocene of Slovakia and Thailand, using paleobotanical records indicate that Slovakia was seasonal warm temperate (Nováky and the Novohrad-Nógrád Basin), while Thailand was tropical with increasing seasonality during the late Langhian and early Serravallian (Chiang Muan), and ever warm tropical later in the Serravalian (Mae Moh) (Table 2 and Supplementary Figures 1, 2). These results complement our paleoenvironmental reconstructions for the Miocene of Thailand and Slovakia based on the overall fungal community composition using a spore traits-based approach, in which the environments from both countries vary from temperate, subtropical to tropical with local humid conditions. In the fossil record, it appears that the fungal community composition is tightly linked with local floral diversity and microhabitat conditions, but this could be driven by environmental factors at a regional scale, as observed in modern ecosystems (Crandall et al., 2020).

Even though preliminary, the results of this study reflect the presence of uncommon taxa in each country, however, we need to deepen our samples analysis to understand those species distribution and presence at a regional scale in the fossil record. Despite the age differences among the samples, amerospores were common, which could suggest that they either share better dispersal pathways, i.e., air, favoring long dispersal distances or have a wider host range (Halbwachs and Bäsßler, 2015; Magyar et al., 2016; Calhim et al., 2018). Applying current knowledge about plant-fungal interactions to fossil taxa will help to identify regional fungal distribution and to elucidate ecosystem-level changes of fungal communities through time (Amend et al., 2010; Talbot et al., 2014; Davison et al., 2015).

CONCLUSION

The FiaWW project is just beginning its exploration of Burdigalian-Serravallian global fungal assemblage distributions. Our assessment of differences at fungal community level between temperate and tropical regions in the middle Miocene hints at greater species richness in Thailand ($p < 0.02$) and variations in the diversity in both countries within the samples.

This study has resulted in the identification of 281 unique fungal morphotaxa, of which 29 have been identified. More fungi need to be identified to the family or genus level from a larger geographical area and with better age control, especially in Europe. Likewise, more identifications are needed to explore fossil fungal distributions.

This study shows that morphological spore traits could be used in the fossil record as first instance, and when identifications are scarce, to start working at fungal community levels, to describe patterns of fungal diversity and paleoenvironments in the fossil record, and to compare different assemblages at large spatial and temporal scales.

Based on fossil fungal composition and paleoclimatic reconstructions, the paleoenvironments for Thailand and Slovakia were similar during the middle Miocene, with tropical to subtropical environments during the MMCO, but more seasonal warm temperate and tropical with increasing seasonality post MMCO.

Identifying fossil fungi based on its modern analogs and following current nomenclature rules is helpful to redefine fungi as reliable proxies to complement paleontological research works.

These results will be integrated in a larger study with three working hypotheses that underpin the FiaWW project: (1) Overall fungal diversity was higher during the MMCO; (2) MMCO fungal assemblages globally showed a greater species richness and abundance of forest-associated ascomycetes; and (3) Fungal diversity during the MMCO was greatest in areas of warm-unseasonal climates.

DATA AVAILABILITY STATEMENT

The original contributions presented in the study are included in the article/Supplementary Material, further inquiries can be directed to the corresponding authors.

AUTHOR CONTRIBUTIONS

JO'K and MP conceived the idea for and designed the study. HB supplied samples and background information. IR, TS, CF, SJ, and LT primarily generated data with supporting data generated by MG and JO'K. NN, IR, and JO'K made taxonomic identifications and determined ecological tolerances. IR, NN, JO'K, MG, and MP completed data analyses. IR and NN drafted figures. IR, NN, and JO'K equally led initial writing with contributions from MG, MP, and SW. IR, NN, and JO'K led manuscript editing with supporting editing by all other authors. All authors contributed to the article and approved the submitted version.

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SUPPLEMENTARY MATERIAL

The Supplementary Material for this article can be found online at: https://www.frontiersin.org/articles/10.3389/ffgc.2021.768405/full#supplementary-material
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