Enzyme activity of topsoil layer on reclaimed and unreclaimed post-mining sites

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Abstract

Topsoil layer contains various components of soil organic matter such as branches, leaves, bark, or metabolites and residues of soil biota. Soil organic matter (SOM) in forest ecosystems consists mostly of lignin, cellulose, chitin and other hydrocarbons. These compounds are decomposed mainly by soil fungi which produce extracellular enzymes to decompose wide range of organic residues. These enzymes may alter nutrient cycling and change soil properties such as water retention capacity, ion exchange capacity, formation of soil microstructure, soil microbial respiration, etc. In this study, we studied enzyme activity in the topsoil layer of post-mining spoil heaps near Sokolov, Czech Republic. We investigated the effect of the following factors and their combinations on enzyme activity: i) dominant vegetation, ii) time of sampling, iii) reclamation, and iv) soil macrofauna.

We measured enzyme activity in plastic mesocosms with autochthonous litter deposited on reclaimed and unreclaimed post-mining sites. We used mesocosms accessible for macrofauna (mesh size > 2 mm) and mesocosms inaccessible for macrofauna (mesh size < 2 mm). Under laboratory conditions, we measured enzyme activity (laccase, oxidase, peroxidase, Mn-peroxidase, alkaline phosphatase, acid phosphatase, endoglucanase, xylanase, cellobiohydrolase, glucosidase, xylosidase and chitinase) using buffer extraction method followed by spectrophotometric assay. We did not find any statistically significant difference between mesocosms accessible and inaccessible for fauna. However, we found significant effect of time of sampling on enzyme activity. Our results showed significant difference between reclaimed and unreclaimed post-mining sites. Reclaimed sites showed significantly higher enzyme activity than unreclaimed sites.

Keywords: enzyme assay, microbial activity, litterbag, macrofauna, soil fauna

Introduction

Soil fungi are widely distributed decomposers of soil organic matter in forest soils (Lavelle et al., 1997; Tedersoo et al., 2014; Wardle and Lindahl, 2014). Cellulose is one of the most important polymeric substances in cell walls and most abundant polysaccharide on Earth (Allison and Vitousek, 2005; Valášková et al., 2007). Soil fungi produce large mycelial networks which grow throughout the litter layer and thus play important role in nutrient transport (Boddy, 1999). Extracellular enzymes such as cellulase released from mycelial networks are responsible for decomposition of plant cell walls and mineralization of complex organic compounds into simple molecules (sugars, amino acids, phosphates) which can serve as food resource for other soil biota (Bardgett and van der Putten, 2014; Pietsch et al., ...
2014; Wardle and Lindahl, 2014). These enzymes are initial agents for nutrient cycling. Activity of extracellular enzymes is widely used as an indicator of soil quality (Baldrian et al., 2008; Šnajdr et al., 2008; Voříšková et al., 2011). Higher activity of extracellular enzymes is associated with regions of high nutrient turnover and primary productivity as well (Bandick and Dick, 1999).

Interaction between soil fungi and other soil microorganisms is very important for enzyme production and its physiological activity. Competitive interactions between fungi and other soil biota (bacteria, actinomycetes) cause lysis of hyphal cells and thus release of nutrients (Wells and Boddy, 2002). In particular, fungifeeding soil fauna consume mycelia and thus release nutrients into the soil where they become accessible for other soil organisms (Bardgett and Cook, 1998). The enzyme activity of the upper soil layer is mainly influenced by selective grazing, selective digestion, selective distribution of microflora, fragmentation of litter, changes of pH and formation of soil aggregates (Lavelle, 2000; Caldwell, 2005; Heděnec et al., 2013; Frouz et al., 2015).

In this short contribution, we focused on the role of interactions between soil fauna and soil microbiota on reclaimed and unreclaimed post-mining sites near Sokolov city (Czech Republic). Open cast lignite mining causes massive destroying of native ecosystems and excavation of large amount of spoil material which is then deposited on spoil heaps (Helingerova et al., 2010). Mining areas and heaps cover over 6000 ha in the Sokolov lignite mining district. Spoil material has initially very low biological activity (Frouz et al., 2001; Helingerova et al., 2010). Soil formation connected with increased biological activity of spoil material colonized by microbes is the basic precondition for ecosystem restoration on spoil heaps (Bradshaw, 1997).

Our research was aimed at the following questions: (i) How does dominant vegetation affect enzyme activity in the topsoil layer? (ii) How is enzyme activity affected by time of sampling? (iii) How does enzyme activity differ in reclaimed and unreclaimed sites? And finally: (iv) How does soil macrofauna affect enzyme activity?

Materials and methods

STUDY SITES AND SAMPLING

For this experiment, we used a system of plastic mesocosms filled with 10 g of autochthonous litter. Mesocosms were deposited on reclaimed and unreclaimed post-mining sites in Velká podkrušnohorská spoil heap (Fig. 1). We chose two experimental sites. The first site (also known under code II) is a 20-year-old unreclaimed post-mining site colonized by Salix caprea, Populus tremula and Betula pendula. The second site (code V2) is a 30-year-old

![Fig. 1. Map of sampling sites on spoil heap.](image-url)
reclaimed post-mining site planted with *Alnus glutinosa* and *Alnus incana*. We used plastic cylindrical mesocosms (10 cm high, 5 cm in diameter, Fig. 2) closed at each end by a net with different mesh size. Mesocosms accessible to fauna were closed by net with mesh size more than 2 mm while inaccessible mesocosms were closed by net with mesh size less than 2 mm. Each mesocosm was filled with 10 g of dried autochthonous litter shaken from trees grown on examined experimental sites. We designed two fields on each site with 25 pairs of randomly distributed mesocosms in five groups (Fig. 2). Distance between pairs of mesocosms was one meter. The experiment was launched in October 2007 and sampling was conducted from February 2008 to September 2008 (Feb 2008, Apr 2008 and Sept 2008). In total, five accessible and inaccessible mesocosms were taken during one sampling campaign.

**ENZYME EXTRACTION AND ASSAY**

Samples from each mesocosm were purified, desalted and assayed under laboratory conditions (Šnajdr et al., 2008). Homogenized samples of autochthonous litter were extracted at 4°C for 2 h on an orbital shaker (100 rpm) with 100 mM phosphate buffer, pH 7 (16:1 w/w), filtered through Whatman 5 filter paper and desalted using PD-10 desalting columns (Pharmacia, Sweden) to remove inhibitory small-molecular-mass compounds (Baldrian et al., 2008). The desalted samples were kept at –18°C until enzyme activity assay (Šnajdr et al., 2008).

Laccase (IUBMB Enzyme Nomenclature, EC 1.10.3.2) activity was measured using monitoring the oxidation of ABTS (2,2-azinobis-3-ethylbenzothiazoline-6-sulfonic acid) in citrate–phosphate (100 mM citrate, 200 mM phosphate) buffer (pH 5.0) at 420 nm (Šnajdr et al., 2008).

Mn-peroxidase (MnP, EC 1.11.1.13) was measured using succinate–lactate buffer (100 mM, pH 4.5) (Šnajdr et al., 2008). The MBTH (3-methyl-2-benzothiazoline hydrazone) and DMAB (3-dimethylaminobenzoic acid) were oxidatively coupled by the enzymes, and the resulted purple indamine dye was detected spectrophotometrically at 595 nm. The results were corrected by comparison with samples without manganese (for MnP) — the addition of manganese sulfate was substituted by an equimolar amount of ethylenediaminetetraacetate (EDTA). One unit of enzyme activity was defined as the amount of enzyme forming 1 mmol of reaction product per min (Šnajdr et al., 2008).

Endoglucanase (EC 3.2.1.4) and xylanase (EC 3.2.1.8) were measured with azo-dyed carbohydrate substrates (carboxymethyl cellulose and birchwood xylan) according to manufacturer’s instructions (Megazyme, Ireland). The reaction mixture contained 0.2 mL of 2% dyed substrate in 200 mM sodium acetate buffer (pH 5.0), and 0.2 mL sample. The reaction mixture was incubated at 40°C for 60 min and the reaction was stopped by adding 1 mL of ethanol followed by 10 s vortexing and 10 min centrifugation (10 000 rpm) (Valášková et al., 2007). The amount of released dye was measured at 595 nm and the enzyme activity was calculated according to standard curves correlating the dye release with the release of reducing sugars. One unit of enzyme activity was defined as the amount of enzyme releasing 1 mmol of reducing sugars per min (Šnajdr et al., 2008).

Cellobiohydrolase (EC 3.2.1.91) was assayed in microplates using p-nitrophenyl-beta-D-cellobioside (PNPC). The reaction mixture contained 0.16 mL of 1.2 mM PNPC in 50 mM sodium acetate buffer (pH 5.0) and 0.04 mL sample. Reaction mixtures were incubated at 40°C for 90–120 min. The reaction was stopped by adding 0.1 mL of 0.5 M sodium carbonate, and absorbance was read at 400 nm (Valášková et al., 2007; Šnajdr et al., 2008).

Glucosidase (EC 3.2.1.21), xylosidase (EC 3.2.1.37) and chitinase (EC 3.2.1.52) were assayed using p-nitrophenyl-b-D-glucoside, p-nitrophenyl-b-D-xyloside and p-nitrophenyl-N-acetyl-b-D-glucosaminide, respectively, using the same method (Valášková et al., 2007).

Phosphatases (EC 3.1.3.1) were assayed using 2 g L–1 p-nitrophenylphosphate in 50 mM sodium acetate buffer (pH 5.0) as described previously. One unit of enzyme activity was defined as the amount of enzyme releasing 1 mmol of p-nitrophenol per min (Šnajdr et al., 2008).

All spectrophotometric measurements were done in a microplate reader (Sunrise, Tecan) on a UV–VIS spectrophotometer (Lambda 11, Perkin–Elmer) and expressed per g dry mass of autochthonous litter.
STATISTICAL ANALYSIS

All obtained data were subjected to Shapiro test to check data normality. Results of our study were analyzed using multivariate methods in statistical program Canoco 4.0 (Lepš and Hadincová, 1992). We used RDA multivariate method for statistical analysis of enzyme activity for different treatments. Time of sampling, accessibility for fauna and reclamation were used as explanatory variables. Monte-Carlo permutation test was used to check the statistical significance of particular environmental variables on the enzyme activity.

Results

Results after the application of multivariate data analysis (RDA) and Monte-Carlo permutation showed (Table) marginal effect of fauna-accessible mesocosms on enzyme activity (P = 0.8). The enzyme activity of fauna-inaccessible mesocosms was not statistically significantly different from fauna-accessible mesocosms (Figs 3, 4). Both inaccessible as well as accessible mesocosms placed in the reclaimed sites showed significantly higher (P = 0.002) enzyme activity than those mesocosms placed in unreclaimed sites (Figs 3, 4).

Figures 3 and 4 show activity of extracellular enzymes (laccase, oxidase, peroxidase, Mn-peroxidase, alkaline phosphatase, acid phosphatase, endoglucanase, endo-xylanase, cellobiohydrolase, glucosidase, xylodside and chitinase) as influenced by reclamation (Fig. 3) and accessibility of mesocosms for soil macrofauna (Figs 3, 4).

We found a statistically significant effect of the time of sampling on enzyme activity (Table). The activity of enzymes in mesocosms were statistically significantly different (P = 0.002) at the various sampling time (from Feb 2008 to Sep 2008). We found variable enzyme activity among individual samples. We found significant effect of reclamation on the activity of enzymes (laccase, oxidase, peroxidase, Mn-peroxidase, alkaline phosphatase, acid phosphatase, endoglucanase, endo-xylanase, cellobiohydrolase, glucosidase, xylodside and chitinase) in mesocosms deposited on reclaimed and unreclaimed sites.

The results subjected to multivariate data analysis (RDA) with the covariates reveal that the reclaimed area showed significantly higher (0.0012) enzyme activity than mesocosms distributed on unreclaimed site (Fig. 5).
Fig. 3. The effect of soil fauna on the enzymatic activity of litter in mesocosms deposited on reclaimed and unreclaimed post mining sites. Monte-Carlo permutation test ($F = 1.004$, $P$-value = 0.8) for the fauna, ($F = 5.456$, $P = 0.002$) for the restoration effect.

Fig. 4. The effect of fauna accessibility on the activity of extracellular enzymes. Explained 39.5% of data variability. Monte-Carlo permutation test ($F = 0.456$, $P = 0.852$).

Fig. 5. Effect of restoration on enzyme activity. Monte-Carlo permutation test with faunal and seasonal effects as covariates ($F = 6.589$, $P$-value = 0.0012). Alder — reclaimed area, Salix — unreclaimed area. Explained 48.5% data variability.
Discussion

Results of our study did not show any significant differences in enzyme activity between fauna-accessible and fauna-inaccessible mesocosms. We agree that small size of soil microorganisms enables their high dispersal activity in the topsoil layer (Barberán et al., 2014). Our results was supported by study of Baldrian et al. (2008) where the soil fauna did not show any significant effect on fungal biomass expressed as concentration of fungal phospholipid fatty acid (PLFAfungi). Results of our study also corresponded with study of Frouz et al. (2006) where the effects of soil macrofauna on basal soil respiration, microbial biomass and C:N ratio in topsoil layer were also not found to be statistically significant. According to Frouz et al. (2006) soil macrofauna significantly affected microbial biomass, basal soil respiration and C:N ratio in deeper soil horizon.

Biological activity in topsoil and litter layer is strongly governed by abiotic factors such as soil temperature, pH and humidity, and biotic factor such as interaction with aboveground biomass (Wardle et al., 2004; Rousk et al., 2010; Zhou et al., 2016). However Crowther et al. (2011) recorded species-specific impact of soil fauna on enzyme activity in laboratory attempts. Wang et al. (2009) found that the loss of biomass in litterbags deposited on Alpine meadows strongly correlated with abundance of oribatids. Frouz (1997) found that microbial activity of litter (expressed as basal soil respiration) significantly increased by adding of bibionid larvae (Bibionidae). These results were discussed in literature (Frouz and Makarova, 2001; Frouz and Simek, 2009) but these studies indicated only short-term effect of soil fauna on biological activity of litter layer.

In other part of our experiment we found that time of the year (season) significantly alters enzyme activity most likely due to climatic factors. Our results corresponded with study of Šnajdr et al. (2008) who found that enzyme activity differed in different seasons (from May 2007 to August 2008). Moreover, this study was supported by study of Baldrian et al. (2008) which was situated at the same study area and in which enzyme activity changed during the season as well. All samples indicated different enzyme activity in different seasons. They found the highest enzyme activity in October 2006. High enzyme activity was caused by high humidity (Wittmann et al., 2004). Effect of seasonality on biological activity of litter (loss of biomass, enzyme activity, decomposition rate and C:N ratio) in litter layer was studied in literature (Dilly and Munch, 1996; Fioretto et al., 2000; Wittmann et al., 2004).

In the last part of our study we focused on the effect of reclamation of post-mining sites on enzyme activity in the topsoil layer. Frouz and Nováková (2005) showed significantly higher microbial activity in reclaimed sites than in unclaimed sites but in successional higher stages, microbial activity among reclaimed and unclaimed sites was closely related. Frouz et al. (2007) recorded significantly higher abundance of soil meso and macrofauna in reclaimed post-mining sites. Reclaimed sites had higher abundance of endogenic earthworms which supported microbial activity in soil. According to Frouz and Nováková (2005), restoration of post-mining sites supported higher microbial biomass. Microbial biomass may be affected by the type of vegetation cover (Frouz and Nováková, 2005). Georgieva et al. (2005) found that the fungal and bacterial biomass in arable soil were significantly affected by vegetation cover. Sites planted by Viccia villosa showed significantly higher microbial biomass than sites planted by Secale cereale (Georgieva et al., 2005).

Conclusions

The results of our study did not show any significant effect of soil fauna on the enzyme activity in the litter layer. However, we found a statistically significant effect of time of sampling as well as significant effect of restoration of post-mining sites on enzyme activity in the topsoil layer.

References


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