Biological activities of soils under rubber trees (Hevea brasiliensis) and interactions with trunk phloem necrosis

Anan Wongcharoen

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Biological activities of soils under rubber trees (*Hevea brasiliensis*) and interactions with trunk phloem necrosis

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RESUME

La qualité du sol influence fortement la productivité d’Hevea brasiliensis qui est un des plus grands producteurs de latex naturel. La nécrose de l’écorce de l’Hévéa (Trunk Phloem Necrosis : TPN) est la maladie de l’hévéa économiquement la plus importante or, à l’heure actuelle, son origine et les causes de son apparition sont mal connues bien que l’apparition de la TPN soit souvent liée à des sylvicultures réalisées dans des conditions extrêmes. Cette étude vise à préciser d’une part les relations entre les activités biologiques du sol et la présence de TPN, d’autre part à déterminer l’influence de différentes types de management du sol sur l’apparition de cette maladie.

Une étude a été menée sur une parcelle d’hévéa très infestée par la TPN. Des échantillons de sols et de faune ont été collectés sous les arbres malades (B), entre un arbre malade et un arbre sain (BH), sous un arbre sain (H) et entre deux arbres sains (HH). L’analyse de la faune du sol a montré que la présence de termites était fortement liée aux arbres malades.

Par ailleurs, la nécrose de l’hévéa semble interagir avec l’activité biologique du sol particulièrement pendant la saison des pluies. Ainsi, l’étude des activités enzymatiques du sol montre que les polysaccharidases (cellulase, xylanase et surtout amylase) sont plus faibles sous les arbres affectés par la TPN alors que les activités liées au cycle de l’azote y sont plus élevées.

Les communautés fongiques cultivables des différents types d’échantillons (B, BH, H, HH) ont été déterminées par culture sur milieu solide de Sabouraud. Bien que la densité fongique ne soit pas significativement différente entre les différentes zones, il apparaît que les espèces Paecilomyces lilacinus et Trichoderma asperellum sont majeures dans les sols prélevés à proximité des arbres malades (B et BH). Un dendrogramme de similarité réalisé à partir des séquences de 28S rDNA fongique montre que les communautés fongiques présentes sous les arbres sains sont très différentes de celles détectées dans les zones comportant des arbres infestés par la TPN. Ce résultat original doit faire l’objet d’études complémentaires afin de déterminer en suivant l’évolution de la maladie sur la parcelle, si les modifications de communautés fauniques ou fongiques observées entre les 4 zones (B, BH, H, HH) sont antérieures ou postérieures à l’apparition de la maladie et donc si elles pourraient constituer un élément de diagnostic de l’état sanitaire de l’exploitation afin de prévenir l’apparition de la TPN.
Une seconde étude s’est intéressée à l’influence de différents amendements organiques sur le fonctionnement biologique du sol sous hévéa ce qui devrait permettre en comparant les informations données par l’étude sur la plantation infestée de prévenir l’apparition de TPN. Les deux amendements comparés, apport de fertilisant organique et couverture de *Pueraria* + amendement organique, entraînent une augmentation de la croissance des arbres ainsi que de la production de latex, en comparaison avec une plantation sans fertilisation. Ces deux types d’amendements semblent aussi prévenir l’apparition de TPN.

Les échantillons de sols ont ici été prélevés dans les interlignes afin d’avoir la plus grande homogénéité possible entre les échantillons. Comme précédemment la macrofaune et les activités enzymatiques du sol ont été déterminées.

La densité de macrofaune du sol est significativement plus faible dans les parcelles sans amendement organique. Le groupe le plus abondant est constitué par les fourmis dont les populations sont beaucoup plus denses dans les zones ayant reçu un amendement organique.

L’analyse en composante principale réalisée à partir des activités enzymatiques montre que la plupart des enzymes présentent des activités plus faibles dans les sols des parcelles sans amendement organique. Les sols des parcelles recouvertes de *Pueraria* ont les plus fortes activités xylanases alors que l’activité FDase est plus élevée dans les sols des parcelles n’ayant reçu que l’apport organique.

L’apport d’amendement organique en améliorant efficacement la teneur en matière organique du sol, non seulement permet un meilleur rendement en latex mais aussi en améliorant le fonctionnement biologique du sol prévient l’apparition de TPN.
ABSTRACT

Soil quality greatly sustains productivity of *Hevea brasiliensis*, which is the important source of natural latex production. The soil quality may have an influence in incidence of trunk phloem necrosis (TPN), which is the currently one of economically important disease of *H. brasiliensis*. This study aims to define the impact of TPN on soil biological activities and to investigate the impact of different soil management practices on soil functions. The TPN disease seems to have effects on soil biology, especially, in the rainy season. Principle component analysis showing associations between healthy and TPN trees, and soil enzyme activities revealed that polysaccharidase (cellulase, xylanase and, particularly, amylase) activities were lower under soil affected by TPN whereas *N*-acetyl-glucosaminidase activity was higher. Termite density was associated with soil under TPN areas. The method of culturable-dependant was used in order to analyze fungal populations. Although fungal density in soil under trees affected by TPN was not significantly different from soil under healthy trees, the fungi *Paecilomyces lilacinus* and *Trichoderma asperellum* were abundant in soil with TPN. The similarity dendrogram of 28S rDNA-DGGE (DNA was extracted directly from soil) revealed that soil fungal community in healthy tree zones was different from TPN tree zones. Soil biological activities in soils with different managements including without external organic input, external organic input and *Pueraria* cover crop + external organic input were investigated in order to find out the proper soil management for *H. brasiliensis*. Principal component analysis showing relationship between different soil managements and soil enzyme activites revealed that most enzyme activites were low in soil without organic management. Xylanase activity was the highest in soil with *Pueraria* cover crop + external organic input whereas the highest FDA hydrolysis activity was associated with soil under external organic input. In investigating macrofauna showed that macrofauna density was the lowest in soil without organic amendment. Ant group was higher abundance in soil with organic amendments than soil without organic amendment. The productivity of *Hevea* tree including tree growth and latex yield was the highest in plantation with *Pueraria* cover crop +organic input, which was also the highest in organic matter and organic carbon contents in soil. This study observed that the soil with the lowest fertility, which indicated in soil without external organic input, seems to associate with high TPN incidence.
THESIS OUTLINE AND OBJECTIVE

The main objectives of this study are i) to investigate impact of trunk phloem necrosis (TPN) disease of *H. brasiliensis* on soil biological activities and ii) to examine the influence of soil management with different organic practices on soil biological activities in *H. brasiliensis* plantation.

Soil biological activities in this study consist of macrofauna community, microbial community and enzyme activities.

In the chapter III.1, discriminations of soil biological activities between TPN affected trees and healthy trees were reported. This chapter reveals impact of TPN affected trees on soil biological activities. Some soil physical and chemical properties were also assessed to compare quality of soil between TPN affected trees and healthy trees. To understand the role of TPN disease on soil biology is an essential component of natural rubber ecosystem.

In the chapter III.2, impact of different soil management practices including soils without external organic input, with external organic input and with *Pueraria* cover crop + external organic input on soil physical and chemical properties, and soil biological activities was revealed. The selecting organic source is needed to ensure short-term productivity and to construct long-term soil quality.

The prospective outputs of this research are i) to obtain potentially biological indicators for impact of TPN affected tree on soil ecosystem. These indicators are useful to early detection of declining soil quality and ii) to obtain the database of soil managements on soil properties. This database contributes to provide the proper soil management practice, which is beneficial to improve soil quality. In addition, increasing soil fertility enhances plant productivity and TPN protection.
I - REVIEW PART
I.1 *Hevea brasiliensis* and trunk phloem necrosis

I.1.1 *H. brasiliensis* as economically important tree

Para rubber tree (*Hevea brasiliensis* Müll. Arg.) had been first found in Amazon basin. This tree is a member of family Euphorbiaceae which can produce natural latex (Purseglove, 1987). The latex is tapped by excision of external tissue of trunk (known as tapping) which contains laticifers (Rudall, 1987). This tree species is a perennial and humid tropical tree, the optimal conditions for growth of *H. brasiliensis* that are: the temperature is in range of 22-30 °C, relative humidity does not exceed 70 to 80 %, and annual rainfall is in between 1500 and 3000 mm. Furthermore, soil properties are also influence the growth and productivity of *H. brasiliensis*. Low latex yield was exhibited in soil under condition of acidic (pH range 4.37-4.54), sandy (75.6-82.6%). Additionally, the higher soil fertility strongly associated with higher latex yield (Akpan *et al*., 2007). On the commercial plantation, *H. brasiliensis* take 5-10 years to reach maturity and the mature trees are 20-30 meters tall.

The *H. brasiliensis* is an economically important tree because its latex is the primary source of natural rubber. All of trees in family Euphorbiaceae, the natural rubber from *H. brasiliensis* is the most elasticity, resilience and toughness. These latex properties are efficient to use as raw materials of many products including transportation (e.g., tires and tire products), industrial (e.g., transmission and elevator belts, hoses and tubes, industrial lining, and bridge bearings), consumer (e.g., sport goods, erasers, foot wear and other apparel), and hygienic and medical (e.g., condoms, catheters and surgical gloves) sectors. The latex can be collected around 30 years or over depend on plantation management and tapping system. After the termination of latex production, the wood of this tree can be used in manufacture of furniture such as small boards, matches, packing boxes, compressed wood textiles and round arches, and also used for fuel.

There has been reported that the natural rubber production is the most in Asia, Africa and Latin American, respectively. During 2000-2006, the most natural rubber in the world was produced from Thailand (FAO, 2009). The *H. brasiliensis* is greatly important in socio-economic fabric of the Southern Thailand and nowadays, it rapidly extends to the Northeastern Thailand (which the soil quality is lower). However, currently, the main problem of latex production in Thailand is trunk phloem necrosis disease. This disease directly affects quantity and quality of latex.
I.1.2 Trunk phloem necrosis disease

The *H. brasiliensis* can be threatened by several pests and disease (which cause by biotic and abiotic factors), trunk phloem necrosis (TPN) is one of the most devastating in *H. brasiliensis* plantation worldwide (especially, industrial scale) because TPN disease is irreversible syndrome and can obstruct latex production. The TPN disease was first found in commercial plantation in Ivory Coast in 1980s by Nandris et al. (1991a). The first stage of this disease begins in the internal bark of trunk, this stage is difficult to detect by naked eye, and the phloem tissue containing the laticifers is necrotic. The investigation of phloem section of TPN affected tree by transmission electron microscopy revealed that cell walls, the middle lamella, plasmodesmata and membrane structures were disorganized (Nicole et al., 1991). When accumulated necrosis was increased, the external bark is cracked, the cracking spread from collar towards the tapping panel. Furthermore, the high severity of TPN disease cause of incision drying up and discontinuing latex production (Nandris et al., 1991a) (Figure 1).

The epidemiology pattern of TPN in the *H. brasiliensis* plantation seems to be contagious disease causing by biotic pathogen. Since the TPN was discovered, numerous of conventional and molecular etiological methods were used to diagnose the causal agents, however, no biotic causal agents involving fungi, bacteria, mycoplasma, virus and viroids were detected (Nandris et al., 1991b; Pellegrin et al., 2007). While, pathogen transmissions had been also investigated consisting bud grafting, bark implantation, tapping knife disinfection and spatio-temporal modeling of TPN dynamics, but no evidence revealed that the TPN disease can spread in plantation by transmission of pathogen (Pellegrin et al., 2004; Peyrard et al., 2006).

Because verification of TPN etiology cannot detect the biotic pathogens by present techniques, Nandris et al. (2004; 2005; 2006) who is the first breakthrough on TPN disease conducted that this disease may be caused by a combination of exogenous and endogenous stresses. The environmental stresses including soil compaction and soil fertility seem to be the major exogenous stresses for TPN disease apparition because soil compaction directly affects plant health by reducing soil porosity, soil water availability and plant root development. Meanwhile, low soil fertility directly influence plant physiological processes, this may reduce the resistance of TPN disease. However, endogenous stresses are also component of casual agent. Incompatibility of rootstock/scion can affect plant physiological strength. The tapping systems may, especially, high frequent tapping and over deep tapping
may also involve with plant physiological dysfunction. Disruption of cyanide metabolism may cause of cell death in TPN affected tree (Cherstin et al., 2004). Hence, productivity of *H. brasiliensis* depends on soil quality level. The good soil quality may also induce resistance of *H. brasiliensis* to TPN disease.

**Figure 1** Symptom of trunk phloem necrosis of rubber tree (*Hevea brasiliensis*)
I.2 Soil quality

I.2.1 Soil quality components

Soils are fundamental resource for plants and soil organisms. Most nutrients and carbon are held in the soil surface, especially, 5-10 cm of the soil profile, and hence this soil layer is important to regulate soil functions and processes (Sparling et al., 2000). Soil quality is potential of soil to perform functions that are essential to plant, animal and human health. The definition of soil quality usually refers to three soil properties including physical chemical and biological properties (Table 1), these soil properties considerably influence plant growth development, plant available nutrients and reducing soil environmental stresses effect on plant (Elliott et al., 1996).

Soil physical properties as refer to soil structural ability are the greatest important to soil quality because soil structure is the primitive factor of soil productivity. Soil textures consist of the relation of sand, silt and clay proportions are primarily important to plant development, soil organism activities and soil biogeochemistry cycling (Chiarini et al., 1998; Silver et al., 2000; Plante et al., 2006). The soil textures also have effect on others soil physical processes. Increasing clay content within soil texture greatly influences soil aggregate stability (Schlecht-Pietsch et al., 1994; Amezketa, 1999). Soil aggregation has positively associated with soil water holding-capacity, soil aeration, and soil infiltration (Mapa and Gunsena, 1995). Soil porosity property is an important part of soil physical quality because porosity levels can indicate the soil compaction which is measured by soil bulk density (Abu-Hamdeh, 2003). Increasing soil compaction levels can decrease quality of soil in respect to reduce infiltration, slow drainage, reduce aeration, restrict plant root development and increase topsoil erosion (Pabin et al., 1998; Lipiec et al., 2003; Neves et al., 2003; Mandal and Tripathi, 2009). High soil compaction also adversely affects soil biological activities such as soil macrofauna community structure (Radford et al., 2001) and microbial activities (Dick et al., 1988; Torbert and Wood, 1992; Lee et al., 1996; De Neve and Hofman, 2000; Jordan et al., 2003). Hence, the good soil structure is fundamental to soil biological and chemical processes.

Soil chemical properties closely interact with soil physical and biological processes. Soil pH is a principal component of soil chemical properties because it strongly influences
available nutrients and activity of soil organisms (Skinner and Todd, 1998; Léonard et al., 2004; Fuentes et al., 2006). Moreover, pH levels in soil also affect soil physical process, especially, clay dispersion and flocculation (Amezketa, 1999). While, organic matter containing in soil is main source to supply energy for soil organisms. The organic matter decomposition processes result in releasing plant available nutrients which can be absorbed by plant (Chander et al., 1998; Warren and Zou, 2002; Faterrigo et al., 2006). Furthermore, soil organic matter can also enhance soil physical properties including soil aggregate stability, water holding capacity and soil bulk density. (Amezketa, 1999; Martens, 2000; Loveland and Webb, 2003; Tripathy and Singh, 2004). The cation exchange capacity (CEC) which is determination of cation absorbable and exchangeable capacity of soil particles is important for soil nutrient retention. The levels of CEC depend on soil texture characteristics and organic matter levels (Kaiser et al., 2008). The CEC has benefit on soil chemical stabilization by binding nutrient-soil particle together, it is applied to be a tool for investigation of soil fertility an nutrient retention capacity (Yagi et al., 2003; Suganya and Sivasamy, 2006; Vagen et al., 2006). Additionally, salinity of soil which can be detected by electrical conductivity (EC) is a main constituent for disturbance of plant growth, soil microbial community and activities, and soil-water balance (Frankenberger and Bingham, 1982; Sinha et al., 1986; Ramoliya et al., 2004; Anjum et al., 2005; Wong et al., 2008). Hence, soil chemical mechanisms are a main factor for plant growth development and soil biological processes.

Soil biological properties have significantly influenced soil fertility and linked to soil physical and chemical properties. The biological indicators in respect of soil quality mostly assess the size and diversity of microbial biomass, soil respiration, available C and N, enzyme activity and macrofaunal biomass. Soil microbes and their activities have benefit on soil structure including soil erosion resistance and water infiltration, these properties positively affect plant growth (Gasperi-Mago and Troeh, 1979; Elliott et al., 1996). Moreover, the soil microorganisms play the major role on soil nutrient cycling because they can degrade organic materials and convert the organic matter to available form for plant growth (McLatchey and Reddy, 1998; Ingram et al., 2005; Schultz and Urban, 2008). Furthermore, soil respiration is used to assess competency of soil to sustain crop growth. Soil respiration is determined by measuring carbon dioxide production in soil surface which results from plant root, microbial decomposition of organic matter and faunal breathing. The rate of soil respiration is mostly regulated by organic matter levels, soil moisture, soil
temperature, and microbial activity (Buchmann, 2000; Epron et al., 2004; Tang et al., 2006). In addition, soil invertebrates are essential organisms to high-quality of soil because they can comminute and distribute material residues, and improve nutrient availability (Anderson and Ingram, 1993; Schädler and Brandl, 2005). The role of macrofaunal community can also enhance soil structure, soil stability, water infiltration and activities of soil microorganisms (Lavelle, 1988; Léonard et al., 2004; Frouz et al., 2006b).

Hence, numerous soil managements are created to improve the quality of soil for sustainable agriculture. At present, soil organic management is concentrated because it can reduce polluted chemical in soil and maintain long-term soil quality. However, efficiency of soil organic management depends on quality of organic matter. The different organic practices regulate soil fertility levels.

**Table 1** Soil properties for characterizing soil quality

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<tr>
<th>Physical</th>
<th>Chemical</th>
<th>Biological</th>
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<tr>
<td>Texture</td>
<td>pH</td>
<td>Microbial biomass</td>
</tr>
<tr>
<td>Infiltration</td>
<td>Salinity</td>
<td>Soil respiration</td>
</tr>
<tr>
<td>Water-holding capacity</td>
<td>Electrical conductivity</td>
<td>Labile organic carbon</td>
</tr>
<tr>
<td>Aggregate stability</td>
<td>Cation exchange capacity</td>
<td>Labile organic nitrogen</td>
</tr>
<tr>
<td>Soil bulk density</td>
<td>Organic matter</td>
<td>Key invertebrate</td>
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<tr>
<td></td>
<td>Extractable, N, P, K</td>
<td>Earthworms</td>
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<tr>
<td></td>
<td>Total organic C</td>
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*Source*: (Elliott et al., 1996)

**1.2.2 Soil quality improvement**

The goal of agriculturist is obtaining the good production which depends on quality of soil. Soil properties are important determinant for plant growth, crop productivity and sustainable agriculture (Warkentin, 1995). Several soil managements are created to maintain and improve soil health in all physical, chemical and biological properties. Additional, organic matters in the soil are the most important for soil quality improvement because organic carbon and nitrogen are the main source for plants and soil organisms. Furthermore, organic input can provide the short-term productivity and simultaneously, building long-term
soil quality. For commercial cropping system in Thailand, there are two main agricultural practices to improve soil efficiency namely organic amendments and cover cropping.

**Organic amendments**

Organic amendments are addition of organic matter sources into soil for modifying soil properties. Sources of organic materials include compost, crop residues, animal manure and green manure. Soil organic matter levels strongly affect soil physical, chemical and biological processes. The final product of organic matter decomposition is humus (or stable soil organic matter) which has benefit on soil physical properties. Soil organic matter levels can improve soil structure including aggregate stability, water infiltration and water holding capacity (Franzluebbers, 2002; Loveland and Webb, 2003; Lado et al., 2004; Gajic et al., 2006). The increasing ability of soil infiltration and permeability can also reduce excess salinity accumulation in soil (Miller et al., 2005). Organic matter contributes to soil fertility because its decomposition processes have released available nutrients for plant growth such as nitrogen, phosphorus and potassium (Blair and Boland, 1978; Pinamonti, 1998; Ingram et al., 2005; Lupwayi et al., 2007; Mukuralinda et al., 2009). In addition, organic matter can increase cation exchange capacity (CEC). This effect contribute to soil chemical stabilization by binding humic acid-mineral nutrients (Oorts et al., 2003; Kaiser et al., 2008). Furthermore, soil organic matter contents also enhance soil microbial community. Increasing microbial activities contribute to soil structure and directly affect soil fertility (Wander et al., 1994; Nelson and Mele, 2006; Fliessbach et al., 2007; Hamer et al., 2008). Additionally, quality of organic materials strongly influence soil macrofaunal, the activity of macrofaunas enhance soil physical properties such as soil hydraulic capacity. Increasing macrofaunal biomass and diversity can contribute to soil microbial activity (Lal, 1988b; Mando, 1997; Mboukou-Kimbatsa et al., 1998; Frouz et al., 2006b; Frouz, 2008). Furthermore, organic matter can reduce soil polluted chemicals because humic acid can absorb chemical pollutants such as pesticides and herbicides (Senesi et al., 1995; Beyer and Blume, 1996; Cells et al., 1997).

**Cover crops**

Cover cropping or green manuring is agricultural practice by growing beneficial plants that can improve soil quality in agroecosystem. Several cover crops including grasses, small grains and legumes are useful to improve soil physical, chemical and biological properties. The leguminous plants are commonly used as cover crop to improve soil fertility
because they can provide nitrogen into soil by nitrogen fixation process which arises from rhizobial bacteria (Frioni et al., 1998). Some cover crop species were efficient in recycling nutrients (Rosolem et al., 2002; da Silva and Rosolem, 2003; Franchini et al., 2004). When cover crops are plowed down or terminated, they release nutrients by mechanisms of decomposition and this also increase soil fertility. Because cover cropping is adding energy sources of living soil organisms, diversity and abundance of soil organisms are enhanced. Increasing activity of soil organisms positively influence soil quality (Boyer et al., 1999; Cederbaum et al., 2004; Ingels et al., 2005; Blanchart et al., 2006; Dinesh et al., 2009). Furthermore, increasing organic matter levels by cover crop residues can improve soil physical properties such as soil structure, and water and nutrient holding capacity (Patrick et al., 1957; Robertson et al., 1991; Arevalo et al., 1998; Muñoz-Carpena et al., 2008). In addition, cover crop can protect damaging soil productivity by reducing or preventing soil surface from erosion (Greene et al., 1994; Martinez-Raya et al., 2006; Zuazo et al., 2006; Zuazo and Pleguezuelo, 2008). The networks of cover crop roots are the suitable habitat for soil macrofaunas and also increase soil porosity (Carof et al., 2007). Increasing soil porosity involves water infiltration and soil water storage (Joyce et al., 2002). Cover crops can protect soil water evaporation and increase soil moisture (Hopkinson, 1971), this is important for activity of soil organisms. Additionally, cover crop can protect desired plant from pests and weeds. The growth of weeds and the germination of weed seeds are suppressed by thick cover crops (Bradshaw and Lanini, 1995; Linares et al., 2008). Some cover crops have ability in destroying plant pests (know as trap crops) or attracting beneficial insects and predators of plant pests, these plant species can be used as biological control of plant pests (Bugg and Waddington, 1994; Rea et al., 2002; Shelton and Badenes-Perez, 2006; Youn and Jung, 2008). Several cover crops can destroy plant disease cycle or reduce population of plant pathogens, and release toxic chemicals affecting phytopathogens in soil including bacteria, fungi and nematode (Crow et al., 2001; Hartz et al., 2005; Blanchart et al., 2006; Timper et al., 2006). Furthermore, cover crops can increase biomass of beneficial invertebrates and microorganisms in soil, which theirs activities can also reduce soil borne pathogens (Boyer et al., 1999; Abawi and Widmer, 2000).

I.3 The role of soil macrofauna
Soil macrofaunal community plays important role on soil quality because its activities influence soil processes. Soil macrofaunal activities enhance soil physical properties including soil porosity and aggregate stability. These soil properties positively affect water infiltration, drainage, aeration and root growth development (Lavelle and Spain, 2001). Furthermore, soil chemical processes are also regulated by soil macrofaunal activity. The soil macrofaunas greatly associate with distribution and protection of organic matter, increasing available nutrients (Lavelle, 1988; Huhta et al., 1998; Scullion and Malik, 2000; Warren and Zou, 2002). The soil macrofaunas are also important in neutralizing soil pH (Frouz et al., 2006a). However, all of macrofaunal taxonomic groups (Anderson and Ingram, 1993), earthworms, termites and ants have been considered as soil ecosystem engineers because they strongly influence soil properties and provide the energy source for microorganism (Jouquet et al., 2006). These soil invertebrates are important in soil physical properties including soil porosity, drainage, aeration and reducing bulk density (Lobry de Bruyn and Conacher, 1990; Mando et al., 1996; Lavelle, 1997; Mando, 1997; VandenBygaart et al., 2000). In addition, these macrofaunal species contribute to soil fertility because they play important role in nutrient turnover and soil nutrient cycling (Lobry de Bruyn and Conacher, 1990; Lavelle et al., 1993; Mora et al., 2005).

The earthworm burrows within soil can increase soil aeration, water infiltration (Pitkänen and Nuutinen, 1998). These soil properties can stimulate soil microbial activities. For instance, the nitrifying bacteria were higher in lining of earthworm burrow than in the soil outside burrow (Parkin and Berry, 1999). In addition, microorganisms living in guts of earthworms and termites can rapidly degrade components of organic matter such as cellulose and lignin (Rouland et al., 1988; Rouland et al., 1991; Zhang et al., 1993; Rouland-Lefevre et al., 2002; Hyodo et al., 2003). This result can increase carbon source and mineral nutrients in soil. Likewise, the activities of earthworms, termites and ants can stimulate microbial activity (Scheu, 1987; 1990; Binet et al., 1998; Holt, 1998; Daube and Wolters, 2000; Scullion and Malik, 2000; Chaou et al., 2003; Desjardins et al., 2003; Ndiaye et al., 2004; Stadler et al., 2006). Increasing microbial activities can contribute to soil structure and soil fertility.

In addition, the prominent ability of termites is clay transportation. Termites can transport clay particles from subsurface horizons to termiria (Fall et al., 2001; Jouquet et al., 2002; Jouquet et al., 2007). The stabilization of soil organic matter was influenced by binding mineral clay-organic matter (Wattel-Koek et al., 2001). Likewise, soil clay can protect organic molecules by slow mineralization rate (Bosatta and Agren, 1997; Yang et al., 2006). Some ants and termites are distributer of organic matter within ecosystem by
transporting organic matter over long distance (Wood, 1988; Anderson and Ingram, 1993). However, combining function of soil macrofauna and soil microorganism is important to maintain and to increase soil health.

I.4. Soil microorganisms

I.4.1. Importance of soil microorganisms

Soil microorganisms are greatly important in soil food web and natural equilibrium. Microorganisms in soil are significantly relative to healthy soil and healthy plant because they are a considerable component of soil physical and chemical processes. The soil microbial communities can improve soil structure for plant growth by enhancing soil aggregate stability. Bacterial polysaccharides and, fungal hypha and metabolic products play important role in binding soil particles together (Robertson et al., 1991; Hu et al., 1995; de Caire et al., 1997; Caesar-TonThat and Cochran, 2000). Soil aggregation is necessary for soil quality to improve infiltration rate, water holding capacity and plant root development. Thus, decreasing microbial biomass and theirs activities are a result in reducing soil aggregation (Neves et al., 2003; An et al., 2009). Numerous species of microorganisms including protozoa, bacteria, fungi and nematode are contained in soil. However, bacteria and fungi seem to be the most important in soil nutrient cycling because they are the first organisms degrading organic materials as theirs energy source. Soil fertility is enhanced by increasing microbial biomass. Soil microbes strongly influence soil biogeochemical because they obviously express their ability in organic matter decomposition, nutrient mineralization and nutrient cycling (Barbhuiya et al., 2004; Ingram et al., 2005; Devi and Yadava, 2006; Faterrigo et al., 2006; Fosu et al., 2007; Lucas et al., 2007). Furthermore, some soil microorganisms have potential to degrade or detoxify chemical pollutants and pesticides in soil (Reed et al., 1989; Rhine et al., 2003; Trabue et al., 2006; Zhang et al., 2008). However, structure of microbial community depend on many factors such as climate, moisture, topography, plant growth, and quantity and quality of substrates (Schinner, 1982; Alvarez et al., 1995; Smith and Goodman, 1999; Norris et al., 2002; Barbhuiya et al., 2004; Tilak et al., 2005; Devi and Yadava, 2006; Waldrop et al., 2006). Additionally, soil chemical (e.g., pH and salinity) and physical condition (e.g., texture and soil-water potential) are also influence soil microbial efficiency (Zhang and Zak, 1998; Emmerling et al., 2001; Gleeson et al., 2008; Wong et al., 2008; Sawada et al., 2009). Hence, soil microbial community composition
structure can respond to environmental change and it has been used for monitoring impact of agricultural practices and ecological stresses on soil health (Steenwerth et al., 2002; Potthoff et al., 2006).

I.4.2. Analysis of soil microbial community structure

The reservoir of soil microorganisms is the pore structure within and between soil particles. The community of soil microorganisms can use as useful indicator of soil quality and ecological stresses because of theirs adaptation. Because soil microbes are greatly important in agroecosystem, the microbial community structure assessment with accurate and reliable methodology is necessary. The methods for studying soil microbial diversity can be categorized into two major groups: biochemical-based and molecular-based methods (Kirk et al., 2004).

Plate count is one of common methods that used for analyzing soil bacterial and fungal diversity. This method is a biochemical-based method to enumerate the viable soil microorganisms. The viable count has benefit to provide information on the active, heterotrophic component of population. This method has been used to evaluate the response of soil microorganisms to polluted soil (Thompson et al., 1998; Piotrowska-Seget et al., 2005; Oliveira and Pampulha, 2006) and agricultural practices (Buyer and Kaufman, 1997; Willison et al., 1997; Leckie et al., 2004; Zablotowicz et al., 2007; Zheng et al., 2007). However, many bacterial and fungal species in soil cannot be cultured in the present methods under laboratory condition (Torsvik et al., 1990; Torsvik et al., 1998; van Elsas et al., 2000). The molecular-based methods have been developed to study microbial diversity in natural environments because these methods can detect non-culturable microorganisms. The denaturing gradient gel electrophoresis (DGGE) which is one of PCR-based approaches is mostly used to evaluate microbial community of environmental sample. Because the PCR products of environmental sample consisting the similar DNA sizes cannot be separated by conventional separation using agarose gel electrophoresis, these DNA can separate and migrate through a polyacrylamide gel by DGGE technique. Furthermore, the DGGE can rapidly analyze the large number of samples and is also reliable and reproducible (Lerner et al., 2006; Nakatsu, 2007; Liang et al., 2008). This approach has been diversely used to analyzed soil microbial community such as impact of perturbed agricultural soil (Jensen et al., 1998; Øvreås et al., 1998), agricultural practices (Nakatsu et al., 2000; Vepsäläinen et al., 2004; Stark et al., 2007), fumigants (Ibekwe et al., 2001), land usage (Bossio et al., 2005),
thermal gradient (Norris et al., 2002) and plant root exudation (Yang and Crowley, 2000) on soil microbial diversity. However, the accuracy of microbial community investigation can be increased by using a combination of culture-dependent and culture-independent methods (Ellis et al., 2003; Edenborn and Sexstone, 2007; Fang et al., 2007). Furthermore, using soil microbial community structure combining soil enzyme activity are usually used to assess impact of soil environment change on soil health arising from polluted chemical (Akmal et al., 2005; Martínez-Iñigo et al., 2009) and soil managements (Sun et al., 2004; Stark et al., 2007; Stark et al., 2008) because large numbers of soil samples are rapidly investigated and it reveal the real impact.

I.5. Enzyme activity in soil

All biochemical transformations in soil are related to the presence of enzymes which are produced by microorganisms (de Caire et al., 2000; Stark et al., 2008), soil animals and plants (Gramss et al., 1999). Nevertheless, the most of enzymes in soil are originated from microbes because they have larger biomass, higher metabolic activity and larger amount of extracellular than plants and animals. Enzymatic mechanism contributes to soil health, which is important for plant productivity. Soil enzymes are the great importance for agriculture because they are involved in soil fertility (Dick et al., 1988; Bandick and Dick, 1999; Hu and Cao, 2007). Organic materials are degraded and transformed into available forms by mechanism of soil enzyme. Soil enzymes play specific role in soil cycling of nutrients such as nitrogen, carbon, phosphorus and sulphur (Gianfreda et al., 2005; Acosta-Martínez et al., 2007; Chen et al., 2008; Sardans et al., 2008). The evaluation of integrative activity of several soil enzymes is greatly efficient to predict the quality of soil because soil enzyme activity is closely related to soil physical and chemical properties, and soil microbial community (Decker et al., 1999; Andersson et al., 2004; Roldán et al., 2005; Acosta-Martínez et al., 2007; Iovieno et al., 2009). Furthermore, some toxic chemicals including pesticides and heavy metals are degraded or detoxified by soil enzyme (Gibson and Burns, 1977; Burns and Edwards, 1979; Niemi et al., 2009). However, the activity of enzyme depends on soil texture, temperature, pH, moisture, heavy-metal contamination, quality and quantity of substrates (Sinsabaugh and Linkins, 1987; Virzo De Santo et al., 1993; Rutigliano et al., 1996; Fioretto et al., 2000; Kourtev et al., 2002; Hinojosa et al., 2004; Sardans and Peñuelas, 2005; Acosta-Martínez et al., 2007).
The activity of soil enzymes is more sensitive than other soil properties (Ndiaye et al., 2000) and rapidly responds to soil environmental change due to agricultural practices and cropping systems (Bandick and Dick, 1999; Kandeler et al., 1999; Klose et al., 1999; Acosta-Martínez and Tabatabai, 2001; Badiane et al., 2001; Hinojosa et al., 2004; Hu and Cao, 2007; Iovieno et al., 2009; Jin et al., 2009). In biochemical analysis, using soil enzyme activity to assess soil quality seems to be elementary whereas obtained results are correlated to other soil properties (Klose et al., 1999; Ndiaye et al., 2000). Hence, several soil enzyme activities are useful to early detection of soil quality change and they can provide information on soil managements or soil ecological stress.

I.6 Soil ecological stress

Soil ecological stress is a result from soil environmental stresses. The stresses of soil environment including abiotic and biotic factors can disturb soil physiological, chemical and biological functions. However, capacity of soil ecosystem to resistant environment stress was limited by thresholds of its resistance. When the strength of these stresses was increasingly, soil ecological changes are caused and declining soil quality is resulted. In addition, the environmental stress is considered that as the great damage of plants because it affects plant physiological functions and plant cell death is consequent.

The environmental stress can classify to several categories arising from both natural and artificial factors such as physical stress, wildfire, pollution, radiation, climatic stress and biological stress. The biological stress can be resulted from soil managements and diseases and hence, this stress may significantly affect on soil agricultural ecosystem.

Tillage

Tillage is an agricultural practice which directly affect on soil physical and chemical properties (Hulugalle et al., 1997). Tillage practice can obstruct process of soil aggregate and is primary cause of soil ecological deterioration over long term (Horn et al., 2003; Williams, 2004). Soil physical properties which affect soil organisms such as water content, soil temperature, soil aeration and mixing of crop residue within soil matrix were change (Kladivko, 2001; Clermont-Dauphin et al., 2004). Soil organic matter in 0-10 cm deep layers which benefit soil microbes and macrofauna was decreased by ploughing (Clermont-Dauphin et al., 2004).
The conventional tillage have reduced soil macrofaunal abundance (Neave and Fox, 1998), especially, earthworms which greatly influence on soil properties such as soil nutrient cycling, soil fertility and soil aggregate (Lavelle, 1988; Lavelle and Spain, 2001). Because the stability of soil aggregates is deteriorated by conventional tillage, the soil compaction is consequence (Shakir and Dindal, 1997). Moreover, the furrows within soil which originate by soil organisms have been also destroyed.

Agricultural soil under reduced tillage and no tillage have associated with higher microbial biomass carbon and microbial biomass nitrogen which are important in increasing soil organic matter content, N mineralization in soil and nutrient supply to crops (Wright et al., 2005). Hence, tillage conservation can maintain soil physical, chemical and biological properties (Reeves, 1997; Amezketa, 1999; Shukla et al., 2003).

**Monocropping**

Monocropping system or continuous cultivation is the agricultural practice which planting the same crop year after year, without practicing crop rotation or resting soil. Monocropping system may have negative impacts on long-term productivity due to decreasing soil quality. Some plant monocultures decreased soil aggregate stability, soil organic matter, available nutrients and cation exchange capacity, while acid saturation was increased (Dalal and Mayer, 1987; Ewel et al., 1991; Amezketa, 1999; Buschiazzo et al., 1999). The lower soil microbial diversity and biomass were detected in the longer monoculture (Holt and Mayer, 1998; Kaur et al., 2000; Yao et al., 2006; Wu and Wang, 2007), the lowering of soil microbes may associate with reducing soil enzyme involving nutrient cycling (Gajda and Martyniuk, 2005). Soil macrofaunal community including biomass, density and diversity was also decreased in long-term continuous planting (Lal, 1988a; Lavelle, 1988; Dangerfield, 1990; Decaëns et al., 1994; Schmidt et al., 2001; Giller et al., 2005; Sileshi and Mafongoya, 2006). These biological effects can cause of deterioration of soil physical and chemical properties in the future, and which consequently led to decline of plant productivity.

**Soil salinity and sodicity**

Soil salinity and sodicity are soil which contains excessive accumulation of salts. Excess salts in soil cause deterioration of soil quality and agricultural productivity. Although soluble salinity positively associates with soil aggregate stability, high salinity can reduce plant growth, plant yield and quality of plant products (Verma and Neue, 1984; Abdul-Halim
et al., 1988; Marosz, 2004; Ramoliya et al., 2004; Ramoliya et al., 2006; Dai et al., 2009). While, high sodium concentrations or high sodicity can disrupt soil physical processes including decreasing infiltration, decreasing hydraulic conductivity and surface crusting (Frenkel et al., 1978; Agassi et al., 1981; Painuli and Abrol, 1986; Mamedov et al., 2001; Levy et al., 2005). However, both salinity and sodicity have adversely affected soil microbial community and biogeochemical processes (Frankenberger and Bingham, 1982; Rietz and Haynes, 2003; Sardinha et al., 2003; Tripathi et al., 2006; Wichern et al., 2006; Muhammad et al., 2008; Wong et al., 2008).
II. MATERIALS AND METHODS
II.1 Experimental site description

The experimental site was the rubber (*H. brasiliensis*) plantation in Buri Rum province, northeastern Thailand (15° 16’ 18” N and 103° 0’ 6” E). This region is tropical climate. The rainy season is from May to October and the dry season is November through March. During the 5 year period 2001-2006, the mean annual temperature for this area was 26 °C and annual precipitation was 1207 mm (which is very low for rubber plantation). Three experimental sites were selected and the description of each site was showed in table 7. All the sites were sampled during the rainy season (July to October 2006) and dry season (December to February 2008).

For chapter III.1, the *H. brasiliensis* plantation with high incidence of TPN disease (20-25% of trees affected) was selected. The *H. brasiliensis* was planted in 1997. Before planting *H. brasiliensis*, the sugarcane (*Saccharum* sp.) was planted in this area. This plantation characterized by low input farming.

For chapter III.2, three *Hevea* plantations were selected by depending different soil managements as follows: no organic amendment (S1) and amending with external organic input (S2) and amending with *Pueraria* cover crop + organic input (S3). The description of each site showed in table 7.

II.2 Macrofauna sampling

In each sampling zones, macrofauna in soil and litter were collected to analyze density and diversity. The macrofauna was defined as the invertebrates with body length greater than 2 millimeters. Litter macrofauna was sampled in an area of 2 x 2 meters. The litter was rapidly put into plastic bag and then macrofauna was carefully collected by hand. Soil macrofauna was performed according to the Tropical Soil Biology and Fertility (TSBF) method (Anderson and Ingram, 1993). The block of soil (25 x 25 x 30 cm) within middle of litter macrofaunal sampling area was taken and macrofauna was collected by hand at three depths (0-10 cm, 10-20 cm and 20-30 cm). The collected macrofauna was preserved in 70% alcohol, except earthworm was kept in 4% formaldehyde. The invertebrates were determined taxonomic groups under standard laboratory.
II.3 Soil sampling and preparation

In each sampling zone, a block of soil (12 x 12 x 10 cm) adjoining zone of litter macrofaunal sampling was taken using hand trowel. The soil block was separated into three layers (0-2 cm, 2-5 cm and 5-10 cm). In each depth, sample soil was air-dried and ground to pass through a 2 mm sieve. Plant residues and roots were removed.

II.4 Soil analysis

Prepared soil sample was used to analyze some soil properties. Soil pH was measured by pH meter using soil : water ratio of 1:1. Organic carbon was measured by oxidizing the soil with potassium dichromate and concentrated sulphuric acid and the remaining concentration of dichromate and ferrous ions determined by titration. The total N was measured by micro kjeldahl method. The organic matter was determined by Walkley and Black method using wet oxidation. Cations exchange capacity (CEC) was determined using double acid (Boric acid and sulphuric acid). Soil texture was performed by hydrometer method.

II.5 Soil enzyme activity

The soil sample with purified water (1 g/150 µl) were incubated at 25 °C for 48 h to reactive the microflora. The activity of β-glucosidase (EC 3.2.1.21), N- acetyl- β-D-glucosaminidase (EC 3.2.1.30), amylase (EC 3.2.1.1), cellulase (EC 3.2.1.4) and xylanase (EC 3.2.1.8) were assayed by a modified procedure described by Mora (2005). The modified Berthelot reaction (Kandeler, 1996) was used to measure urease (EC 3.5.1.5) activity. Total microbial activity potential was measured through Fluorescein diacetate (FDA) hydrolysis assay, which hydrolyze colorless fluorescein diacetate and release a colored end product fluorescein (Green et al., 2006). The enzyme assays were carried out in triplicate to measure the activities and Control was simultaneously performed in each sample

Heterosidases

β-glucosidase and N-acetyl-β-D-glucosaminidase activities were assayed using p-nitrophenyl (PNP)-β-D-glucopyranoside and p-nitrophenyl (PNP) N-acetyl-β-D-
glucosaminide (Sigma) as substrates. Soil (200 mg) was incubated at 37 °C for 2 hours with 200 µl of 1% (w/v) substrate and 100 µl of citrate phosphate buffer at pH 5.8 (McIlvaine, 1921). After incubation, the sample was centrifuged at 13000 rpm for 5 min and the supernatant (250 µl) was placed into spectrophotometer tube (size 5 ml). The reaction was stopped by adding 3 ml of 2% Na₂CO₃ (w/v). The absorbance was measured at 420 nm and the enzyme activity was expressed as µg PNP g⁻¹ h⁻¹ by comparison with standard curve.

Polysaccharidases

The amylase, cellulase and xylanase activities were determined using starch, carboxymethyl-cellulose (CMC) and xylan as substrates. Soil (200 mg) was incubated with 200 µl of 1% (w/v) substrate and 100 µl of citrate phosphate buffer at pH 5.8 at 37 °C for 4 hours with constant agitation. After incubation, sample was centrifuged at 13000 rpm for 5 min and 250 µl of supernatant was placed into glass tube. The reducing sugar was measured using the method described by (Somogyi, 1945) and (Nelson, 1944). 500 µl of Somogyi solution (diluted with purified water, 1:1) was added and boiled for 20 min. The sample solution was immediately cooled using laboratory water. Afterwards, 250 µl of Nelson solution was added and gently vortexed. Purified water (4 ml) was added and the absorbance was measured at 650 nm. The result was expressed as µg glucose g⁻¹ h⁻¹ by comparison with standard curve.

Urease

The urease activity was measured by determination of released ammonium. Soil sample (200 mg) was mixed with 400 µl of 100 mM borate buffer (pH 10.0) and 50 µl of 720 mM urea. After incubation at 37 °C for 4 hours with constant agitation, the reaction was stopped with 3 ml of acidified 2 M KCl and 1.5 ml of the mixture was centrifuged at 13000 rpm for 5 minutes. The supernatant (1 ml) was mixed with 5 ml of a 1:1:1 (v/v/v) mixture of sodium salicylate/ sodium nitroprussiate reagent, 0.3 M NaOH and distilled water. Afterwards, 2 ml of 39.1 mM sodium dichloroisocyanurate and 9 ml of distilled water were added. After agitation in the dark for 30 minutes, the absorbance was measured at 660 nm and the enzyme activity was express as µg N-NH₄⁺ g⁻¹ h⁻¹ by comparison with standard curve.

Fluorescein diacetate (FDA) hydrolysis

Soil (200 mg) was mixed with 25 µl of 4.8 mM FDA solution and 1.5 ml of a citrate phosphate buffer at pH 5.8. The sample was incubated at 37 °C for 2 h with constant
agitation. After incubation, 25 µl of substrate was added to control. The suspension was centrifuged at 13000 rpm at 4 °C for 5 min and 750 µl of supernatant was taken. The activity was stopped by adding 750 µl of acetone. The amount of FDA hydrolyzed was measured as absorbance at 490 nm and the activity was expressed as µg fluorescein g⁻¹ h⁻¹ by comparison with standard curve.

II.6 Soil fungal community

III.6.1 Soil fungal enumeration and isolation

Enumeration of soil fungi were conducted according to a method adapted from Diouf et al. (2005) using plating dilution method. Soil (5g) was agitated in 50 ml of dispersing solution for 30 min and allowing soil precipitation for 15 min. The fungal suspensions were decimal diluted in physiological solution (9g/l of Nacl). Each of fungal dilution with 200 µl was cultured in Sabouraud medium (added 0.005% w/v of Chloramphemicol – Sigma). The assay was performed in triplicate. The numbers of fungal colonies were determined after incubation at 27 °C for 5 days. The differential colony morphotypes were isolated and purified on Sabouraud medium. The mycelia of selected fungi were conserved at -20 °C for molecular identification.

III.6.2 Identification of culturable fungi

The selected fungi were identified the taxonomic position by using molecular biology technique. DNA was extracted from mycelium according to Di Battista method (Di Battista, 1997). The nucleotide fragment of 650 bp (ITS1-5.8S-ITS2 region) was amplified by PCR technique using primer set ITS1 (5’TCCGTAGGTAACCTGCGG 3’) and ITS4 (5’ TCCTCCGCTTATGGATATGC 3’) (White et al., 1990). PCR amplification was performed using Ready-To-Go Taq poloymerase (Pharmacia). The condition of PCR cycles were initial denaturation at 94 °C for 5 min, 29 cycles of 94 °C for 45s, 60 °C for 45s, 72 °C for 1 min and final elongation step at 72 °C for 30s. PCR amplification was performed with a thermal cycle (GenAmp PCR System 2400; Perkin-Elmer). All PCR products were investigated using 5 µl of product by electrophoresis in 2% (w/v) agarose gel in 0.5X TAE buffer (Biorad) and ethidium bromide (0.5 µg/ml) staining. DNA sequencing was performed by Genome Express
The DNA sequences were compared with other DNA sequences using GenBank BLAST searches.

### III.6.3 Genetic diversity of soil fungi

Fungal DNA was extracted from soil samples as described by Porteus (Porteous et al., 1997). The DNA region of 28S rDNA was amplified using primer set 403f (5' GTGAAATTGTTGAAAGGGAA 3') and 662r (5' GACTCCTTGGTGCCGTGTT 3') with 40 bases GC clamp (Sigler and Turcob, 2002). PCR amplification was performed using Ready-To-Go Taq polymerase (Pharmacia). Cycling condition were initial denaturation at 94 °C for 5 min, 35 cycles of 94 °C for 30s, 50 °C for 1 min, 72 °C for 2 min and final elongation step at 72 °C for 10 min. PCR amplification was performed with a thermal cycle (GenAmp PCR System 2400; Perkin-Elmer). The PCR products were investigated as described above and PCR amplicons were determined for different position by DGGE (Denaturing Gradient Gel Electrophoresis) using DcodeTM Universal Mutation Detection System (Biorad, Richmond, CA). For DGGE, 20 µl of product was loaded onto 8% (w/v) polyacrylamide gel containing linear gradient of the denaturants urea and formamide increasing from 35%-60% (100% denaturant contains 7 M urea and 40% (v/v) formamidine). Gel electrophoresis was carried out at initial 20 V for 10 min then 75 V for 16 h. Gels were run in 0.5X TAE at constant temperature of 60 °C. Gels were stained in an ethidium bromide solution and photographed under UV transillumination using GelDoc 2000 system (Biorad).

### II.7 Analysis of plant growth

In each sampling zone, four trees surrounding the sampling zone, perimeter in each tree at height of 150 cm from base was measured for assessment of tree growth.

### II.8 Statistical analysis

The averages of tested parameters in different zones were compared and statistic difference was measured by ANOVA-Tukey test (\( P<0.05 \)) using R program version 2.7.0. The relationship between enzyme activities and sampling zones was determined by principal component analysis (PCA) using ADE-4 software (Thioulouse et al., 1996). A permutation test was used to check the significance of the groups suggested by PCA.
For 28S rDNA diversity, the DGGE banding patterns were analyzed based on the band appearance and disappearance using image analysis software Quantity One (Bio-rad). Banding data of DGGE gel were used to evaluate genetic similarity of fungi among different four sampling zones. The genetic similarity of fungal 28S rDNA was measured using Dice coefficients and similarity dendrogram was created using UPGMA (Unweighted Pair Group Method of Arithmetic means) through the program, Quantity One version 4.2.1.
III - RESULTS
CHAPTER III.1
TPN AND SOIL ENVIRONMENT
Abstract

Trunk phloem necrosis (TPN) is currently one of the most economically important diseases of the *Hevea brasiliensis*. Investigation for etiology TPN revealed that no biotic pathogen was detected, however, a multidisciplinary study has recently suggested that the disease may be caused by a combination of exogenous and endogenous stresses. These stresses can affect plant physiology and may also have an incidence on soil biology and soil biochemistry. In this study, macrofauna diversity, soil enzyme activities, soil microbial community and soil chemical properties were compared for the soils under trees affected by TPN and under healthy trees. The impact of TPN on soil biological activities seems to be expressed in rainy season because maybe the trees were continuously stressed by tapping. The macrofauna analysis revealed that termite density in both litter and soil was the highest in areas with TPN affected trees. Earthworm and ant abundances were high in soil under healthy trees. Coleoptera abundance and diversity were low in areas with TPN affected trees. The principal component analysis revealed that polysaccharidase (cellulase, xylanase and, particularly, amylase) activities were lower in soil under trees affected by TPN whereas N-acetyl-glucosaminidase activity was higher. Investigating fungal density in soil showed that *Paecilomyces lilacinus* and *Trichoderma asperellum* were dominated in soil with TPN. The similar dendrogram of 28S rDNA-DGGE banding patterns for examining fungal community composition showed that the community of fungi under TPN was clearly divided from under healthy tree. These findings suggested that termite density and N-acetyl-glucosaminidase activity are useful as indicators of TPN, while lower values of polysaccharidase activities in TPN areas are the consequences of this disease. The impact of TPN on soil ecosystem was discussed.
Hevea brasiliensis (well know as para rubber tree) is an important source of natural. Trunk phloem necrosis had become a serious problem in H. brasiliensis plantation system because this disease prevents latex production. TPN was first described in the 1980s in Ivory Cote by (Nandris et al., 1991a). The first visible symptoms are the necrosis of the bark at the collar with thin cracks that extend upward towards the tapping panel. When the inner phloem, which contains lacticifers, becomes affected, the latex production is affected and the tapping panel and incision dry up (Nicole et al., 1991). On-going and completed studies to identify a biotic causal agent have failed to confirm any biotic causal agents such as fungi, bacteria, mycoplasma, virus or viroids (Nandris et al., 1991b; Pellegrin et al., 2004; Pellegrin et al., 2007). In addition, several experiments (bud grafting, bark implantation, tapping knife disinfection) and the spatio-temporal modeling of TPN dynamics have shown no evidence of pathogen (Peyrard et al., 2006).

The first breakthrough on TPN was reported by (Nandris et al., 2004; 2005; Nandris et al., 2006) who concluded that this disease may be caused by a combination of exogenous and endogenous stresses. In particular, soil compaction which can induce water stress affecting the health of the trees and low soil fertility were thought to increase the risk of the disease. Furthermore, the non-optimal vascular interface between the rootstock and scion, combined with a major physiological dysfunction (cell disruption) can lead to the release of the native cyanide in the inner bark, thus causing the death of the cells and the spread of the necrosis in the phloem (Cherstin et al., 2004). All these stresses may disrupt physiological metabolisms of rubber trees and consequently affect the soil biological microenvironment around the tree. However, the real impact of TPN on soil ecology is unknow.

Soil is a basic resource for agricultural production systems. Soil quality is fundamental to the success of crop production systems. Soil quality is associated with soil biological activity. Soil macrofauna, considered to be ecological engineers or ecosystem engineers, play an important role in soil formation, affecting organic matter decomposition and nutrient cycling. Increasing soil macrofauna diversity tends to increase the density of soil microorganisms as macrofauna can modify the conditions for microorganisms which are important sources of enzymes in soil (Mora et al., 2005). Enzymes are the main mediators of soil biological process: organic matter degradation, mineralization and nutrient recycling. In this study, cellulase, xylanase and amylase were selected because they are very important in
carbon-cycle. β-glucosidase releases low molecular weight sugars, which are important as energy sources for soil microorganisms. Urease and N-acetyl-glucosaminidase play an important role in nitrogen-cycle. Fluorescein diacetate (FDA) hydrolysis assay was used to measure enzyme activity of decomposer microorganisms, which are important in forest ecosystems for mineralization and turnover of organic matter. Both the soil microorganisms and macrofauna can modify the physical and chemical properties of the soil and can be used to monitor soil quality in agricultural land. The soil biochemical state has been suggested as sensitive indicator of soil ecological stress recovery processes in agro-ecosystems. Numerous studies have been published on the potential use of enzyme activities as indicators of soil productivity. Enzyme activities in soil have been used to investigate pesticide pollution (Sannino and Gianfreda, 2001), the impact of cultivation (Vepsäläinen et al., 2001; Acosta-Martínez et al., 2007) and metal pollution (Epelde et al., 2008). Soil enzyme activities have also been proposed as indicators for suppressive defense against soil pathogens (Naseby et al., 1998; Bruggen and Semenov, 2000; Rasmussen et al., 2002; Leon et al., 2006).

The relationship between plant and microorganisms are an important part of forest soil ecosystem. Soil microbes are dominant for soil quality because they are greatly important in soil functioning and fertility (Elliott et al., 1996). Soil microbes can be used as sensitive indicator of disturbing soil ecology (Dick, 1992; Dilly and Blume, 1998). The microorganisms play role in soil nutrient cycling, because their extracellular enzymes have degraded organic matters as their energy source (Pavarina et al., 1999; Vidal et al., 1999; Tribak et al., 2002; Khaid et al., 2006). The capacity of microbes in decomposition depends on type and abundance of organic materials (Saliha et al., 2005). Some microbes have ability to convert toxic chemical (such as cyanide) in soil to essential nutrient for plant (Maier-Greiner et al., 1991). The microbes also influence soil physical property, especially, the stability of soil aggregation (Harris et al., 1964; TonThat and Cochran, 2000), which benefit for water movement in soil and plant root growth. Soil microbial diversity is influenced by different factors including soil management system (Lucasa et al., 2007), climate change (Freya et al., 2008) and soil pollution (Kandeler et al., 2000), and thus, soil microbial community structure has been noted as important for understanding the relationship between environmental factors and ecosystem functioning.

The effect of different factors on composition of soil microbial community can be investigated by both cultivation-dependent and cultivation-independent. However, the methods under cultivation-dependent still have limitation because numerous microorganisms in soil cannot culture in synthetic media under laboratory conditions (Borneman et al., 1996;
To study these microbes, cultivation-independent methods have been invented for examination of environmental samples. The molecular biology techniques are powerful tools to detect the microbial populations present in soil both culturable and unculturable microorganisms. The denaturing gradient gel electrophoresis (DGGE) has been used widely for investigation of soil microbial community. This technique involves amplifying target-DNA region and PCR products with similar length are separated in polyacrylamide gel containing gradient of denaturing conditions (Kirk et al., 2004; Nakatsu, 2007). The DGGE technique is reliable, reproducible and rapid to study soil microbial diversity (Duineveld et al., 2001; Maarit-Niemi et al., 2001). This technique is used for studying soil microbial community on agricultural practices (Nakatsu et al., 2000; Hagn et al., 2003; Siddique et al., 2005; Sekiguchi et al., 2008), their degrading substrates (van der Wal et al., 2006; Cosgrove et al., 2007; Das et al., 2007) and plant genotypes (Ikeda et al., 2006).

The biological indicators in soil under TPN affected tree merit study. The determination of macrofauna diversity and soil enzyme activities may be useful for obtaining a greater understanding of the impact of this disease on soil ecosystem functions. The main objectives of this study were (i) to investigate soil biological indicators in *H. brasiliensis* plantation, which had high incidence of TPN and (ii) to determine the differences in such indicators between the soils under trees affected by TPN and under healthy trees.
III.1.2 EXPERIMENTAL DESIGN

To determine the impact of TPN, TPN affected trees and healthy trees were selected using the annual tree health survey which carried out by D. Nandris. Ten plots were chosen and in each plot, samples were taken from four different sampling zones: near collar of affected tree (B), midway between affected trees and healthy trees (BH), near collar of healthy trees (H) and midway between healthy trees (HH) (Figure 2).

For enzyme assay and microbial culture, the soil samples were mixed using three identical sampling zones with depth. For genetically microbial diversity, the soil samples were mixed using three identical sampling zones with 0-5 cm and 5-10 cm depths.

**Figure 2** Sampling scheme in the rubber tree plantation: healthy tree (O), tree affected by TPN (●). B is near the collar of tree affected by TPN, BH is midway between affected tree and healthy tree, H is near the collar of a healthy tree, HH is midway between healthy rubber trees.
III.1.3 RELATIONSHIP BETWEEN TPN AND DIFFERENT SOIL PROPERTIES

Soil properties in rainy and dry seasons were showed in table 2.

*Soil pH*

In rainy and dry seasons, no significant difference in soil pH was observed for soil under midway between healthy trees (HH), midway between TPN affected tree and healthy trees (BH), near collar of healthy tree (H) and near collar of TPN affected tree (B).

For each zone, soil are more acidic in rainy than in dry season

*Organic carbon*

In rainy season, organic carbon was the highest in soil under midway between healthy trees (HH) and was significantly higher ($P<0.05$) than under near collar of TPN affected tree (B).

In dry season, no significant difference was observed from whole samples.

*Total nitrogen*

No treatment was significantly different in soil nitrogen in rainy season.

For dry season, the soil in midway between healthy trees (HH) was significantly higher ($P<0.05$) than in near collar of trees (B and H).

*C:N ratio*

In rainy and dry seasons, no significant difference of the C:N ratio was observed between sampling zones.

*Organic matter (OM)*

The values of organic matter were the highest in soil under midway between healthy trees (HH) in both seasons. In rainy season, the OM content in soil was significantly lower ($P<0.05$) under near collar of TPN affected tree (B) than under midway between healthy trees (HH). No significant different in soil organic matter was observed in dry season.

*Cation exchange capacity (CEC)*
The evaluation of cation exchange capacity of soil in rainy and dry seasons showed that the CEC was significantly higher ($P<0.05$) in the soil under midway between TPN affected tree and healthy trees (BH) than under near collar of healthy tree (H).

**Soil moisture**

In rainy season, soil moisture was clearly higher than in dry season. For each season, the moisture under near collars of trees (B and H) was significantly lower ($P<0.05$) than under midway between trees (BH and HH).

**Clay content**

In rainy season, soil clay content was higher in healthy tree areas (H and HH) than in TPN areas (B and BH) and was the highest in midway between healthy trees (HH). However, no significant different was observed in both rainy and dry season.
Table 2  Soil properties in *H. brasiliensis* plantation under near collar of TPN affected tree (B), midway between TPN affected tree and healthy trees (BH), near collar of healthy tree (H) and midway between healthy trees (HH) in rainy and dry seasons.

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<th>B</th>
<th>BH</th>
<th>H</th>
<th>HH</th>
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<tr>
<td><strong>Rainy season</strong></td>
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<tr>
<td>pH</td>
<td>4.29 ± 0.30 a</td>
<td>4.58 ± 0.25 a</td>
<td>4.34 ± 0.34 a</td>
<td>4.50 ± 0.37 a</td>
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<td>OC (%)</td>
<td>0.51 ± 0.11 a</td>
<td>0.76 ± 0.22 ab</td>
<td>0.55 ± 0.14 ab</td>
<td>0.78 ± 0.27 bc</td>
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<td>Total N (%)</td>
<td>0.07 ± 0.01 a</td>
<td>0.09 ± 0.03 a</td>
<td>0.08 ± 0.02 a</td>
<td>0.08 ± 0.02 a</td>
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<tr>
<td>C:N ratio</td>
<td>7.02 ± 1.59 a</td>
<td>8.76 ± 2.00 a</td>
<td>7.09 ± 2.85 a</td>
<td>9.95 ± 3.20 a</td>
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<tr>
<td>OM (%)</td>
<td>0.88 ± 0.19 a</td>
<td>1.30 ± 0.39 ab</td>
<td>0.94 ± 0.24 ab</td>
<td>1.34 ± 0.48 b</td>
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<td>CEC (%)</td>
<td>3.83 ± 0.26 a</td>
<td>5.21 ± 1.46 b</td>
<td>3.83 ± 0.39 a</td>
<td>4.43 ± 0.56 ab</td>
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<td>Moisture (%)</td>
<td>11.63 ± 0.91 a</td>
<td>17.24 ± 2.21 b</td>
<td>10.62 ± 2.94 a</td>
<td>16.22 ± 2.30 b</td>
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<tr>
<td>Clay (%)</td>
<td>8.15 ± 2.26 a</td>
<td>8.43 ± 2.47 a</td>
<td>9.05 ± 1.79 a</td>
<td>9.54 ± 1.36 a</td>
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<td><strong>Dry season</strong></td>
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<tr>
<td>pH</td>
<td>5.02 ± 0.26 a</td>
<td>5.08 ± 0.54 a</td>
<td>5.06 ± 0.32 a</td>
<td>5.02 ± 0.37 a</td>
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<tr>
<td>OC (%)</td>
<td>0.53 ± 0.06 a</td>
<td>0.57 ± 0.28 a</td>
<td>0.52 ± 0.06 a</td>
<td>0.71 ± 0.20 a</td>
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<tr>
<td>Total N (%)</td>
<td>0.07 ± 0.02 a</td>
<td>0.08 ± 0.02 ab</td>
<td>0.06 ± 0.01 a</td>
<td>0.11 ± 0.03 b</td>
</tr>
<tr>
<td>C:N ratio</td>
<td>8.17 ± 1.59 a</td>
<td>7.44 ± 3.29 a</td>
<td>9.14 ± 1.41 a</td>
<td>7.68 ± 4.18 a</td>
</tr>
<tr>
<td>OM (%)</td>
<td>0.91 ± 0.10 a</td>
<td>0.99 ± 0.48 a</td>
<td>0.90 ± 0.11 a</td>
<td>1.23 ± 0.35 a</td>
</tr>
<tr>
<td>CEC (%)</td>
<td>4.13 ± 0.52 ab</td>
<td>4.94 ± 1.08 bc</td>
<td>3.85 ± 0.48 a</td>
<td>5.20 ± 0.86 c</td>
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<tr>
<td>Moisture (%)</td>
<td>1.71 ± 0.93 a</td>
<td>3.18 ± 0.60 b</td>
<td>1.57 ± 0.98 a</td>
<td>3.99 ± 0.65 b</td>
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<tr>
<td>Clay (%)</td>
<td>9.17 ± 1.64 a</td>
<td>8.45 ± 1.75 a</td>
<td>8.76 ± 2.23 a</td>
<td>9.97 ± 1.31 a</td>
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</table>

Means within the same column followed by contrasting letters are significantly different among treatment using TukeyHSD test (*P*<0.05) (+: standard deviation, n=9)
**Multivariate analysis**

Principal component analysis on soil variables with the results obtained in rainy and dry seasons were constructed (Figure 3). In rainy season, PCA showed axis 1 and axis 2 for 56.6% and 17.8% of the variance, respectively. In dry season, PCA showed axis 1 and axis 2 for 50.7% and 28.2% of the variance, respectively. Testing the significance of groupings (sites) was carried out on 10,000 permutations and was significantly discriminated ($P<0.00$) in rainy seasons. In both seasons, the PCA revealed that the soil under near collar of trees (B and H) was associated with the low values of soil variables. The high values of organic matter, organic carbon and CEC were associated with soil under midway between trees (BH and HH). The total N was associated with soil under midway between trees (BH and HH) in, particularly, dry season. The C:N ratio was associated with soil under midway between trees (BH and HH) in, particularly, rainy season.
Figure 3 Principal component analysis showed the relationship between soil variable and sampling zones with near collar of TPN affected tree (B), near collar of healthy tree (H), midway between TPN affected tree and healthy tree (BH) and midway between healthy trees (HH) during rainy (A) and dry (B) season.
III.1.4 TPN AND SOIL MACROFAUNA

III.1.4.1 Macrofauna density

Litter macrofauna

The litter macrofauna was sampled in rainy and dry seasons including Coleoptera, termites, earthworms, Chilopoda, Diplopoda, Arachnida, ants and other invertebrate (Figure 4). In each season, the macrofauna was the most abundant in area under between tree affected by TPN and healthy trees (BH), followed by under between healthy trees (HH), near collar of TPN affected tree (B) and near collar of healthy tree (H) but densities were significantly different only in dry season.

During the rainy season, ants are the major group in the 4 sampling zones whereas “other group” was the dominate in the dry season. Densities of Arachnida and Coleoptera were quite the same in each zone and for each season. They were relatively stable groups. Termite density was the highest in HH zone during dry season. In rainy season, termites are only present near TPN zones (B, BH). Earthworm was collected only during rainy season between tree affected by TPN and healthy trees (BH). The density of “others” increased more than ten folds in dry season in comparison to rainy season. This important increase is due to an important density of Orthoptera, which does not present in rainy season.

![Litter macrofauna](image)

**Figure 4** Density of macroinvertebrates in litter (individual m⁻²) in the rainy and dry season
Soil macrofauna

There was great difference in the macrofauna density between rainy and dry season. In dry season, macrofauna was found only in TPN areas (B and BN). The density near collar of TPN tree (B) was significantly higher than between affected TPN and healthy trees (BH). Coleoptera and Chilopoda were only found under between TPN affected tree and healthy trees (BH). However, these invertebrates were present in low numbers. Furthermore, termite and ant were only found under near collar of TPN affected tree (B).

The soil macrofauna in rainy season included Coleoptera, termites, earthworms, Chilopoda, ants and other invertebrate (Figure 5). The macrofauna was the most abundant in soil near collar of TPN affected tree (B). The coleoptera density was the highest under between healthy trees (HH) while no Coleoptera was observed in area under near collar of TPN affected tree (B). Termite density was higher under TPN disease (B and BH) than under healthy tree (H and HH). The earthworm population was lower in areas near collar of trees (B and H) than areas midway between trees (BH and HH). The Chilopoda was only found under between healthy trees (HH). Abundance of ant was the highest in soil under between healthy trees (HH). In area near collar of TPN affected tree (B), only termite group was observed.

![Soil macrofauna](image)

**Figure 5** Density of macroinvertebrates in soil (individual m\(^{-2}\)) in the rainy and dry season
III.1.4.2 Macrofauna diversity

Diversities of litter and soil macrofauna were showed in table 3.

**Litter macrofauna**

In dry and rainy seasons, the diversity of Coleoptera was the highest under between healthy trees (HH) (included Carabidae, Elateridae, Scarabaeidae, Staphylinidae, Tenebrionidae, Dysticidae and larva). In rainy season, the termite diversity was the highest under near collar of TPN affected tree (B) (included two species collected *Ancistrotermes* sp. and *Macrotermes* sp.) while *Macrotermes* sp. was only found in litter under BH and HH zones. In contrary, in dry season, *Ancistrotermes* sp. and *Macrotermes* sp. were found in all sampling zones.

For the two sampling period, ant diversity was the highest under between healthy trees (HH) but the species richness of ant was the highest in rainy season.

**Soil macrofauna**

In rainy season, the diversity of Coleoptera was the highest under between healthy trees (HH) (included Scarabaeidae, Tenebrionidae and larva) while soil under between TPN affected tree and healthy trees (BH), and near collar of healthy tree (H) have only one Coleoptera species (included larva and Carabidae, respectively). The termite *Ancistrotermes* sp. and *Macrotermes* sp. were found in all sampling zones. Ant diversity was the highest under between healthy trees (HH) (*Morphospecies* sp. 2 and larva) while ant *Camponotus* sp. 3 and *Odontoponera transvers* were observed under between TPN affected tree and healthy trees (BH), and near collar of healthy tree (H), respectively.

In dry season, Coleoptera Tenebrionidae and Chilopoda were only found under between TPN affected tree and healthy trees (BH). However, these invertebrates were present in low numbers. Furthermore, termite (*Macrotermes* sp.) and ant (*Calyptomyrmex* sp.) were only found under near collar of TPN affected tree (B).
Table 3  Diversity of macrofauna in soil and litter under near collar of affected tree (B), midway between affected trees and healthy trees (BH), near collar of healthy trees (H) and midway between healthy trees (HH) during rainy and dry seasons

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<th>Litter</th>
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<td><strong>Coleoptera</strong></td>
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<td>Carabidae sp.</td>
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<td>Cerambycidae sp.</td>
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<td>Coccinellidae sp.</td>
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<td>Elateridae sp.</td>
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<td>Scarabaeidae sp.</td>
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<td>Staphylinidae sp.</td>
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<td>Tenebrionidae sp.</td>
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<td>Dytiscidae sp.</td>
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<td><strong>Termites</strong></td>
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<td>Ancistrotermes sp.</td>
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<td>Macrotermes sp.</td>
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<td><strong>Ants</strong></td>
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<td>Odontoponera transversa</td>
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<td>Pachycondyla sp.</td>
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<td>Pheidole sp. 2</td>
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<td>Morphospecies sp. 2</td>
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<td>Myrmecina sp.</td>
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<td>Calyptomyrmex sp.</td>
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<td>Paratrechina longicornis</td>
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<td>Anoploepis gracilipes</td>
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<td>Oecophylla longinoda</td>
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<td>Camponotus sp. 1</td>
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<td><em>Camponotus sp. 6</em></td>
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<td><em>Larva</em></td>
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(Table 3 continue)
Diversity index

The specific data obtained on the major group were used to calculate several diversity indexes: specific richness, Shannon index and diversity Simpson index (Table 4).

The specific richness considered that only the number of species or morphospecies was higher under near collar of TPN tree (B) and between healthy trees (HH). However, the analysis of the Shannon and Simpson diversity indexes showed that these two zones were clearly different. The zone under midway between healthy trees (HH) was high in both indexes (1.10 and 0.90, respectively), this indicated that the macrofauna community was composed by species with equivalent population. For the zone under near collar of TPN tree (B), both indexes were low (0.54 and 0.67, respectively). This demonstrated that there are few species dominated by a great number of individuals whereas other species were represented by few individuals. Thus, the macrofauna community was equilibrated in area of midway between healthy trees (HH) whereas it was incompletely equilibrated in area of near collar of TPN tree (B). Thus, the macrofauna community was equilibrated in area of midway between healthy trees (HH) whereas it was incompletely equilibrated in area of near collar of TPN tree (B).

On the other hand, the similarity index of Jacquard was constructed (Table 5) and showed that the macrofauna composition in condition of identical sampling zones (H and B; 0.79, and BH and HH; 0.80) was higher similar than in different sampling zones (H and HH; 0.71, and BH and H; 0.62). This indicated that the influence of conditions of sampling zones were important more than the presence of TPN disease.
**Table 4** Specific richness, Shannon index and Simpson index of macrofauna

<table>
<thead>
<tr>
<th></th>
<th>Specific richness</th>
<th>Shannon</th>
<th>Simpson</th>
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<tbody>
<tr>
<td>B</td>
<td>20</td>
<td>0.54c</td>
<td>0.67c</td>
</tr>
<tr>
<td>BH</td>
<td>16</td>
<td>0.93b</td>
<td>0.83b</td>
</tr>
<tr>
<td>H</td>
<td>15</td>
<td>0.90b</td>
<td>0.83b</td>
</tr>
<tr>
<td>HH</td>
<td>21</td>
<td>1.10a</td>
<td>0.90a</td>
</tr>
</tbody>
</table>

**Table 5** Jacquard similarity index of macrofauna

<table>
<thead>
<tr>
<th></th>
<th>B</th>
<th>BH</th>
<th>H</th>
<th>HH</th>
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</thead>
<tbody>
<tr>
<td>B</td>
<td>1</td>
<td>0.77</td>
<td>0.79</td>
<td>0.78</td>
</tr>
<tr>
<td>BH</td>
<td>1</td>
<td>0.62</td>
<td>0.80</td>
<td></td>
</tr>
<tr>
<td>H</td>
<td>1</td>
<td>0.71</td>
<td></td>
<td></td>
</tr>
<tr>
<td>HH</td>
<td></td>
<td>1</td>
<td></td>
<td></td>
</tr>
</tbody>
</table>
Conclusion

The macrofauna diversity has used as biological indicator of soil disturbance because it reflects the impact of environmental change (Nahmani et al., 2006; Velasquez et al., 2007; Yankelevich et al., 2007). In rainy season, most invertebrates were associated with areas between trees (BH and HH) more than areas near collars of trees (B and H). The areas between trees in this plantation were accumulated with organic materials, especially, litter. The higher food source influences macrofaunal population and diversity. In rainy season, the difference in Coleoptera abundance between BH and HH was not obviously different but the diversity was the highest in area between healthy trees (HH). Likewise, the earthworm population was mostly found in soil under between healthy trees (HH). For the tropical forest, the quality of litter has been showed to affect macrofaunal population (Zou, 1993; Mboukou-Kimbatsa et al., 1998; Gonzalez and Zou, 1999). The quality of plant materials depends on chemical component, which is the important factor to regulate the macrofaunal abundance and diversity. As mentioned above, the TPN affected tree may contain tannins more than healthy tree. The exceeding tannins can adversely affect some invertebrates. Tannins are binding protein agents, which are able to bind digestive enzymes to limit assimilation in insects. This has effect on survival and growth of macrofauna. Consequently, macrofaunal abundance and diversity was low (Loranger-Merciris et al., 2007). However, (Ayres et al., 1997) suggested that the same tannin had different effect on different herbivore insects. In this plantation, Coleoptera genus Cerambycidae, Elateridae, Scarabaeidae and Dytiscidae were not found in both soil and litter under TPN affected trees (B and BH). These Coleoptera genera may be affected by tannins from trees affected by TPN. On the other hand, some insect species in Carabidae and Tenebrionidae were detected in all sampling zones. These macrofaunas may resist polyphenols.

The termite density was higher under TPN affected trees (B and BH). Severe TPN causes the bark to slough off at the collar. As only healthy bark containing latex and cyanide is able to block boring insects, the unprotected wood can be attacked by termites to become part of their habitat. Consequently, the zones with the highest termite population were strongly associated with areas under the affected trees.

In dry season, soil macrofaunal diversity in dry season was lower than in rainy season. The Coleoptera was detected only in soil under between TPN affected tree and health trees (BH) but its diversity was very low, only Tenebrionidae was found. This may be due to climate condition. The climate is considered to be the most important factor in the regulation of macrofaunal community. In drought circumstance, soil was arid and hard because of
decreasing soil water levels. Soil moisture, which is greatly associated with seasonal variation, significantly affects soil macrofauna (Sroka and Finch, 2006). In this season, no earthworm was found in all sampling zones because the earthworms maybe migrate to deeper soil layers. In during winter and summer, the earthworms can reach a depth of 40-45 cm and was in quiescent stage (Reddy and Pasha, 1993) whereas macrofauna was collected in a maximum depth of 30 cm in this study.
III.1.5  TPN AND SOIL ENZYME ACTIVITY

III.1.5.1  Rainy season

These results have been published in Forest Pathology
Biological activity of soils under rubber trees (Hevea brasiliensis) affected by trunk phloem necrosis

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Summary

Trunk phloem necrosis (TPN) is currently one of the most economically important diseases of the rubber tree (Hevea brasiliensis). Investigations of the aetiology of the disorder have failed to identify any biotic causal agents but a multidisciplinary study has recently suggested that the disease may be caused by a combination of exogenous and endogenous stresses. These stresses can affect plant physiology and may also have an impact on soil biology and soil biochemistry. In this study, macrofauna diversity and soil enzyme activities were compared for the soils under trees affected by TPN and under healthy trees. Principal component analysis revealed associations between TPN and macrofauna diversity and soil enzyme activities. Groups of ants and termites were associated with the soil under healthy and trees affected by TPN respectively. Polysaccharidase (cellulase, xylanase and, particularly, amylase) activities were lower in the soil under trees affected by TPN whereas N-acetylglucosaminidase activity was higher. These findings suggested that termite density and N-acetylglucosaminidase activity are useful indicators of TPN, while lower values of polysaccharidases activities are the consequence of this disease. The impact of TPN on soil ecosystem was discussed.

1 Introduction

The rubber tree (Hevea brasiliensis) is an important source of natural rubber. Rubber tree bark necrosis [Trunk phloem necrosis (TPN)] has become a serious problem in rubber tree plantations because this disease prevents latex production. TPN was first described in the 1980s in Ivory Coast by Nandris et al. (1991a). The first visible symptoms are necrosis of the bark at the collar with thin cracks that extend upward towards the tapping panel. When the inner phloem, which contains the laticifers, becomes affected, the latex production is affected and the ‘panel cut’ becomes dry (Nicole et al. 1991). On-going and completed studies to identify a pathogen have failed to confirm any causal agents such as fungi and fungus-like organisms (e.g., Phytophthora, Pythium, Fusarium), bacteria, phytoplasma, viruses or viroids (Nandris et al. 1991b; Pellegrin et al. 2007). In addition, several experiments (bud grafting, bark implantation, tapping knife disinfection) and the spatio-temporal modelling of TPN dynamics have shown no evidence of pathogen transmission (Peyrand et al. 2006).

The first breakthrough on TPN was reported by Nandris et al. (2004, 2005, 2006) who concluded that this disease may be caused by a combination of exogenous and endogenous...
stresses. In particular, soil compaction which can induce water stress affecting the health of the trees and low soil fertility were thought to increase the risk of the syndrome. Furthermore, the non-optimal vascular interface between the rootstock and the scion, combined with a major physiological dysfunction (cell disruption) can lead to the release of native cyanide in the inner bark, thus causing cell death and spread of necrosis in the phloem (Cherstín et al. 2004). All these stresses may disrupt physiological metabolisms of rubber trees and consequently affect the soil biological microenvironment around the tree. However, the real impact of TPN on soil ecology is unknown.

Soil is a basic resource for agricultural production systems. Soil quality is fundamental to the success of crop production systems and is associated with soil biological activity. Soil macrofauna, considered to be ecosystem engineers (sensu Jones et al. 1994), play an important role in soil formation, affecting organic matter decomposition and nutrient cycling. Increasing soil macrofauna diversity tends to increase the density of soil microorganisms as macrofauna can modify the conditions for microorganisms which are important sources of enzymes in soil (Mora et al. 2005). Enzymes are the main mediators of soil biological processes: organic matter degradation, mineralization and nutrient cycling. Both the soil microorganisms and macrofauna can modify the physical and chemical properties of the soil and can be used to monitor soil quality in agricultural land. The soil biochemical state has been suggested as sensitive indicator of soil ecological stress recovery processes in agro-ecosystems. Numerous studies have been published on the potential use of enzyme activities as indicators of soil productivity. For example, enzyme activities in soil have been used to investigate pesticide pollution (Sannino and Gianfreda 2001), the impact of cultivation (Vepsäläinen et al. 2001; Acosta-Martínez et al. 2007) and metal pollution (Ezelde et al. 2008, Li et al. 2009). Soil enzyme activities have also been proposed as indicators for suppressive defence against soil pathogens (Naseby et al. 1998; Bruggen and Semenov 2000; Rasmussen et al. 2002; Leon et al. 2006).

Biological indicators in soil under rubber trees affected by TPN merit study. The determination of macrofauna diversity and soil enzyme activities may be useful for obtaining a greater understanding of the relationship between this disease and soil ecosystem functions. The main objectives of this study were (i) to investigate soil biological indicators in a rubber tree plantation which had high incidence of TPN and (ii) to determine the differences in such indicators between the soils under trees affected by TPN and under healthy trees.

2 Materials and methods

2.1 Field site description

The study was conducted in the 9-year-old rubber plantation in Buri Ram province, northeastern Thailand (15°16′18″N and 103°05′6″E). The climate is tropical with the rainy season occurring from May to October. During the 5-year period 2001–2006, the mean annual rainfall was 1207 mm (which is very low for rubber plantations) and the mean annual temperature was 26°C. The soil is sandy loam with 67, 24 and 9% of sand, silt and clay, respectively. The rubber tree plantation was characterized by low input farming and a high incidence of TPN (20–25% of trees affected). The study site was sampled during the rainy season (July to October 2006).

2.2 Experimental design

Trees affected by TPN and healthy trees were selected using the annual tree health survey carried out by D. Nandris. Ten plots were chosen and in each plot, four different
sampling zones were selected: next to affected trees (B), half-way between affected trees and healthy trees (BH), next to healthy trees (H) and half-way between healthy trees (HH) (Fig. 1).

2.3 Sampling design

2.3.1 Sampling of macrofauna

The tropical soil biology and fertility method was used to sample soil macrofauna (Anderson and Ingram 1993). A block of soil (25 \times 25 \times 30 \text{ cm}) was taken and macrofauna collected at three depths (0–10, 10–20 and 20–30 cm). The macrofauna were hand sorted to extract invertebrates with a body length greater than 2 mm. Invertebrates were preserved in 4% formalin and separated into nine broad taxa (orders or families).

2.3.2 Sampling of soil and preparation

For sampling the soils in each zone, a block (12 \times 12 \times 10 \text{ cm}) was taken using hand trowel. The block was separated into three layers (0–2, 2–5 and 5–10 cm). The samples were air-dried, ground and sieved to 2 mm. Plant residues and roots were removed. Three identical sampling zones with depth, samples were mixed for enzyme activity analysis.

2.4 Soil enzyme activity

To evaluate soil enzyme activity, the soil samples were incubated with purified water (1 g/150 \mu l) at 25°C for 48 h to reactivate the microflora. Enzyme assays were carried out in triplicate to measure activities. β-Glucosidase, N-acetyl-glucosaminidase, amylase, cellulase and xylanase activities were assayed using a modification of the procedure described by Morea et al. (2005). The modified Berthelot reaction (Kandeler et al. 1999) was used to measure urease (EC 3.5.1.5) activity. Total microbial activity potential was measured using fluorescein diacetate (FDA) hydrolysis in which colourless FDA is hydrolysed releasing the coloured end product fluorescein (Green et al. 2006).

β-Glucosidase and N-acetyl-glucosaminidase activities were assayed using p-nitrophenyl β-D-glucopyranoside and p-nitrophenyl N-acetyl glucosaminide (Sigma, St. Louis, MO, USA) as substrates. Soil (200 mg) was incubated at 37°C for 2 h with 200 \mu l of 1\% (w/v) substrate and 100 \mu l of citrate phosphate buffer at pH 5.8. The reaction was stopped with 2% Na2CO3 (w/v). After centrifugation at 12000 g for 5 min, absorbance was measured at 420 nm. Enzyme activities were expressed as μg PNP released g\(^{-1}\) soil h\(^{-1}\).

![Sampling scheme in the rubber tree plantation: healthy tree (⊙), tree affected by TPN (●). B is near the collar of tree affected by TPN, BH is midway between affected tree and healthy tree, H is near the collar of a healthy tree, HH is midway between healthy rubber trees.](image)
Activities of amylase, cellulase and xylanase were determined using starch, carboxymethyl-cellulose and xylan as substrates, respectively. Soil (200 mg) was incubated with 200 µl of 1% (w/v) substrate and 100 µl of citrate phosphate buffer at pH 5.8 at 37°C for 4 h. After incubation, reducing sugars were measured using the method of Somogyi (1945) and Nelson (1944). Absorbance was measured at 650 nm and the results were expressed as μg glucose released g⁻¹ soil h⁻¹.

For urease activity, 200 mg of soil was mixed with 400 µl of 100 mM borate buffer (pH 10.0) and 50 µl of 720 mM urea. After incubation at 37°C for 4 h, the reaction was stopped with 3 ml acidified 2 M KCl and 1.5 ml of the mixture was centrifuged at 13000 rpm for 5 min. One millilitre of the supernatant was mixed with 5 ml of a 1:1:1 (v/v/v) mixture of sodium salicylate/sodium nitroprussiate reagent (1.7%, w/v and 0.12%, w/v, respectively), 0.3 M NaOH and distilled water. Finally, 2 ml of 39.1 mM of sodium dichloroisocyanurate and 2 ml of distilled water were added. After agitation in the dark for 30 min, absorbance was measured at 660 nm and enzyme activity expressed as μg N-NH₄⁺ g⁻¹ soil h⁻¹.

For FDA hydrolysis, 200 mg of soil was incubated for 2 h at 37°C, with 25 µl of 4.8 mM FDA solution and 1.5 ml of a citrate phosphate buffer at pH 5.8. After incubation, the suspension was centrifuged at 13000 rpm at room temperature for 5 min and 750 µl of the supernatant was taken. The activity was stopped by adding 750 µl of acetone. The amount of FDA hydrolysed was measured as absorbance at 490 nm and the activities were expressed as μg fluorescein g⁻¹ h⁻¹.

2.5 Statistical analysis

The relationship between macrofauna diversity and enzyme activities and the different sampling zones was determined by principal component analysis (PCA) using ADE-4 software (Thiouloise et al. 1997). A permutation test was used to check the significance of the groups suggested by PCA. ANOVA was used to apply the Tukey test (p < 0.05).

3 Results

3.1 Macrofauna

The soil macrofauna density was higher near the collars of the affected trees (B); it was not significantly different in between the trees (BH and HH). Near the collars of the healthy trees (H), the macrofauna density was clearly lower than in the other zones (Fig. 2). Termites were the largest taxonomic group near the collars of the trees and were the only macrofauna group near the affected trees (B) and in between healthy and affected trees (BH). The macrofauna diversity was higher in both zones between trees than near the collars of the trees (Fig. 3). Although the macrofauna density was low between healthy trees (HH), the diversity was the highest.

3.2 Enzyme activities

3.2.1 Polysaccharidases

Cellulase, xylanase and amylase activities were lowest in soil sampled near the collars of affected trees (B) and lower between affected and healthy trees (BH) than near healthy trees (H and HH) (Table 1). Xylanase and amylase activities near the collars of the affected trees (B) were significantly lower (p < 0.05) than near healthy trees. The soil under healthy trees had the highest amylase activity, significantly higher than under affected trees (Table 1).
Fig. 2. Density of soil macrofauna in the four zones (B, BH, H and HH).

Fig. 3. Macrofauna diversity: proportions of each taxonomic group in sampling zones, number of species in the taxon.

3.2.2 Heterosidases

β-Glucosidase, N-acetyl-glucosaminidase hydrolysis activities were not significantly different between soil sampled near affected trees or healthy trees.
3.2.3 Urease

The highest values of urease were between healthy and affected trees (BH) and were significantly higher than for the soil close to both affected and healthy trees (B and H). There was no significant difference in activity for the samples taken from between trees (BH and HH) (Table 1).

3.2.4 FDA hydrolysis

Fluorescein diacetate hydrolysis activities of the soil samples collected in the various zones were not significantly different (Table 1).

3.3 Multivariate analysis

Principal component analysis showed that the soil under healthy trees (H) was associated with the highest amylase activity and the highest density of ants along axis 1 (Fig. 4). For the soil under affected trees (B), all the enzyme activities tested were low while the termite density was the highest. Soil samples with high cellulase, xylanase and amylase activities were associated with healthy trees (H and HH). Multivariate analysis revealed that the highest N-acetyl-glucosaminidase activity was associated with the soil between affected and healthy trees (BH). β-Glucosidase, urease and FDA hydrolysis activities were higher in sampling areas between trees (HH and BH).

4 Discussion

Macrofauna diversity was used as a biological indicator of soil disturbance because it reflects the impact of ecological stresses (VELASQUEZ et al. 2007). Termite density was highest near affected trees (B and BH). Severe TPN causes the bark to slough off at the collar. As only healthy bark containing latex and cyanide is able to block boring insects, the unprotected wood can be attacked by termites to become part of their habitat. Consequently, zones with the highest termite populations were strongly associated with the soil under the affected trees.

Principal component analysis revealed that most of the enzyme activities were low near the collars of the rubber trees (B and H), as these zones did not accumulate leaf litter that is a major source of soil organic matter. Plant litter is an importance source of nutrients for soil microorganisms (BARDETT and SHINE 1999; RUAN et al. 2005; ABRAHAM and CHUDEK 2008) and litter input can increase soil enzyme activity (DORNBUSH 2007). High urease, β-glucosidase and FDA hydrolysis activities were found between the trees (HH and BH).

**Table 1.** Soil enzyme activities (µg of product g⁻¹ soil h⁻¹) in four different zones; near collar of tree affected by TPN (B), midway between tree affected by TPN and healthy tree (BH), near collar of healthy tree (H) and midway between healthy trees (HH).

<table>
<thead>
<tr>
<th></th>
<th>Cellulase</th>
<th>Xylanase</th>
<th>Amylase</th>
<th>β-Glucosidase</th>
<th>N-Acetyl-glucosaminidase</th>
<th>Urease</th>
<th>FDA hydrolysis</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>0.3 ± 0.5 a</td>
<td>0.8 ± 1.6 a</td>
<td>2.3 ± 2.2 a</td>
<td>11.1 ± 8.2 a</td>
<td>1.1 ± 0.8 a</td>
<td>0.8 ± 0.3 ab</td>
<td>3.1 ± 1.0 a</td>
</tr>
<tr>
<td>BHF</td>
<td>1.0 ± 0.8 ac</td>
<td>2.8 ± 0.8 b</td>
<td>6.7 ± 1.6 a</td>
<td>29.8 ± 28.6 a</td>
<td>1.2 ± 0.7 a</td>
<td>2.0 ± 0.9 c</td>
<td>4.7 ± 2.6 a</td>
</tr>
<tr>
<td>H</td>
<td>0.7 ± 0.7 ac</td>
<td>3.2 ± 1.2 b</td>
<td>22.9 ± 7.8 c</td>
<td>8.6 ± 5.8 a</td>
<td>0.9 ± 0.8 a</td>
<td>0.7 ± 0.5 a</td>
<td>3.4 ± 1.0 a</td>
</tr>
<tr>
<td>HHF</td>
<td>1.5 ± 1.0 bc</td>
<td>3.8 ± 1.1 b</td>
<td>12.9 ± 2.9 b</td>
<td>32.2 ± 32.6 a</td>
<td>1.3 ± 0.6 a</td>
<td>1.7 ± 1.1 bc</td>
<td>5.3 ± 2.3 a</td>
</tr>
</tbody>
</table>

Means within the same row followed by different letters are significantly different using Tukey HSD test (p < 0.05) (±: standard deviation, n = 9).
These zones accumulated abundant litter and thus the environmental conditions (such as soil microorganisms, humidity, etc.) producing these enzymes were similar.

Cellulase, xylanase and amylase activities were higher near healthy rubber trees (H and HH) than affected trees (B and BH) (Fig. 2). Thus, healthy trees were strongly associated
with polysaccharidases which can break down carbohydrates and release glucose as a source of available energy. These enzyme activities were rather low near affected trees. This effect may result from the difference in chemical composition between the leaves from healthy and affected trees which could have an impact on the quality of the litter and consequently on enzyme activities. Numerous studies have shown that litter quality influences various enzyme activities, increasing or decreasing the litter decomposition in the soil (Fioretto et al. 2000; Kourtev et al. 2002; Gusewell and Freeman 2005). In the rubber tree plantation in this study, dead plant materials or plant components (such as bark, branches, litter and latex) were the main source of organic matter for decomposition. The accumulation of defensive chemicals in the leaves or plant components of affected trees may affect soil enzyme activities. Some papers have reported that TPN may be caused by multiple exogenous and endogenous stresses (Nandris et al. 2004, 2005, 2006). Tapping bark for latex (too frequently, badly carried out or using chemicals for latex stimulation) contributes to the cumulative stress. Rubber trees can respond to such wounds or stresses by producing defensive chemicals including polyphenol as tannins (Eyles et al. 2004). Tannins can inhibit the activity of extracellular microbial enzymes (Scalbert 1991) including polysaccharidases such as cellulase and xylanase (Gamble et al. 2000; Barahona et al. 2006). Furthermore, high levels of tannins may influence soil mineralization and humus formation, decreasing soil fertility (Bradley et al. 2000; Driese and Whitham 2000; Kraus et al. 2004; Nierop et al. 2006; Wurzburger and Hendrick 2007).

Principal component analysis showed that N-acetyl-glucosaminidase activity was associated with the area between health and affected trees (BH). N-Acetyl-glucosaminidase is one of the enzymes involved in chitin degradation (Ekenler and Tabatabai 2002). Chitinase has been found in plant tissue such as seed, bark, stems, leaves and roots (Hernes and Hedges 2004; Kosola et al. 2004; Kraus et al. 2004; Santos et al. 2004) but are also produced by microorganisms. In plant, this enzyme is rarely detectable in healthy plants but can be produced in reaction to damage or stress caused by various biotic or abiotic factors (Maugh et al. 1988). Hevea brasiliensis can produce chitinases (Bokma et al. 2002) with antifungal activity against several pathogenic fungi (Parijs et al. 1991; Giordani et al. 2002; Kanekwiroon et al. 2008). Chitinase activity is high in laticifers producing latex but low in healthy tissues such as leaves, stems or roots (Martin 1991). However, it would be important to determine the origin (trees or microorganisms) of N-acetyl-glucosaminidase activity detected in the BH to understand why it is produced only in this zone.

This study showed that TPN of H. brasiliensis has strong long-term effects on the soil ecosystem and soil fertility. PCA showed that the soil near affected rubber trees has lower polysaccharidase activities. This may be the result of secondary metabolites produced by affected trees. As little is known about the relationship between TPN and defensive chemicals and the impact of these agents on soil enzymes in rubber plantations, research is required into the secondary metabolites of trees affected by TPN.

5 Conclusion

The PCA carried out in this study showed for the first time that some biological activity indicators can help to distinguish between healthy trees (H and HH) and trees affected by TPN (B and BH). This distinction seems to be useful for observing the impact of TPN on the ecology of rubber tree plantations. Termite density was strongly associated with TPN (B and BH). High cellulase, xylanase and amylase activities were associated with healthy trees and were lower near diseased trees. PCA also indicated that N-acetyl-glucosaminidase activity was associated with the zone between healthy and affected trees (BH). It is suggested, therefore, that termite density, polysaccharidase activities and
N-acetyl-glucosaminidase activity can be used as indicators of the impact of TPN on soil ecology. However, the production of secondary metabolites as a result of TPN must be investigated. Further research needs to be carried out to assess the biological activities during the dry season in order to confirm this hypothesis.

Acknowledgements

This research was performed under Franco-Thai cooperation programme in higher education and research. This work was supported financially by the Institute for Research and Development (IRD), France government and Khon Kaen University, Thailand government and by French Institute for Rubber-Michelin- Soechinco-SIPH.

References


Conclusion

The principal component analysis carried out in this study showed for the first time that some biological activity indicators can help to distinguish between healthy trees (H and HH) and trees affected by TPN (B and BH). This distinction seems to be useful for observing the impact of TPN on the ecology of rubber tree plantations. Termite density was strongly associated with TPN (B and BH). High cellulase, xylanase and amylase activities were associated with healthy trees and were lower near diseased trees. PCA also indicated that \( N \)-acetyl-glucosaminidase activity was associated with the zone between healthy and affected trees (BH). It would seem that termite density, polysaccharidase activities and \( N \)-acetyl-glucosaminidase activity can be used as an indicator of the impact of TPN on soil ecology. However, the production of secondary metabolites as a result of TPN must be investigated. Further research needs to be carried out to assess the biological activities during the dry season in order to confirm this hypothesis.
III.1.5.2 Dry season

The expression of soil enzyme activities in dry season was showed in table 6. Enzyme activities were higher in soil under midway between trees (BH and HH) than under near collar of trees (B and H).

Polysaccharidases

Cellulase and xylanase activities under TPN trees (B and BH) were higher than under healthy trees (H and HH) and were the highest in midway between TPN and healthy trees (BH). Amylase activity was the highest in midway between healthy trees (HH) and was the lowest in near collar of TPN affected tree (B). However, no statistically significant difference in polysaccharidase activities was observed.

Heterosidases

β-glucosidase activity was higher in soil under midway between trees (BH and HH) than under near the collar of tree (B and H). This enzyme activity in soil under midway between healthy trees (HH) was the highest and soil under near collar of TPN tree (B) was the lowest. However, no significant difference was observed.

The assay of N-acetyl-glucosaminidase activity showed that the soils under midway between trees (HH and BH) were significantly higher \((P<0.05)\) than under near collar of tree (B and H). Soil under midway between healthy trees (HH) was higher N-acetyl-glucosaminidase activity than under midway between TPN and healthy trees (BH).

Urease

Soil under midway between healthy trees (HH) showed the highest urease activity and it was significantly higher \((P<0.05)\) than other zones.

Fluorescein diacetate (FDA) hydrolysis

Soil under midway between TPN and healthy trees (BH) was the highest FDA hydrolysis activity and was significantly higher \((P<0.05)\) than under near collar of trees (B and H).
Table 6  Soil enzyme activities (µg of product g⁻¹ soil h⁻¹) in *H. brasiliensis* plantation under four different zones; near collar of tree affected by TPN (B), midway between tree affected by TPN and healthy tree (BH), near collar of healthy tree (H) and midway between healthy trees (HH) during dry season

<table>
<thead>
<tr>
<th></th>
<th>B</th>
<th>BH</th>
<th>H</th>
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</tr>
</thead>
<tbody>
<tr>
<td>Cellulase</td>
<td>2.1±2.2 a</td>
<td>2.4±1.6 a</td>
<td>2.0±0.6 a</td>
<td>1.8±1.5 a</td>
</tr>
<tr>
<td>Xylanase</td>
<td>0.9±1.5 a</td>
<td>1.2±1.2 a</td>
<td>0.0±0.1 a</td>
<td>0.8±1.0 a</td>
</tr>
<tr>
<td>Amylase</td>
<td>2.1±2.7 a</td>
<td>4.1±1.4 a</td>
<td>3.9±5.9 a</td>
<td>5.6±1.7 a</td>
</tr>
<tr>
<td>β-glucosidase</td>
<td>10.0±7.6 a</td>
<td>25.1±26.4 a</td>
<td>14.4±19.2 a</td>
<td>34.0±27.0 a</td>
</tr>
<tr>
<td>N-acetyl-glucosaminidase</td>
<td>3.4±1.3 a</td>
<td>6.9±2.0 b a</td>
<td>2.6±1.5 a</td>
<td>7.2±2.0 b</td>
</tr>
<tr>
<td>Urease</td>
<td>1.2±0.5 a</td>
<td>2.0±0.6 a</td>
<td>1.5±0.3 a</td>
<td>3.2±1.4 b</td>
</tr>
<tr>
<td>FDA hydrolysis</td>
<td>0.8±0.2 ab</td>
<td>1.3±0.4 c</td>
<td>0.7±0.3 a</td>
<td>1.2±0.5 bc</td>
</tr>
</tbody>
</table>

Means within the same column followed by contrasting letters are significantly different among treatment using TukeyHSD test (*P*<0.05) (+: standard deviation, n=9)
Multivariate analysis

PCA showed axis 1 and axis 2 for 55.3% and 17.5% of the variance, respectively (Figure 6). Testing the significance of groupings (zones) was carried out on 10,000 permutations and was significantly discriminated (P<0.002). The PCA revealed that the soil under near collar of healthy tree (H) was associated with the low expression of all tested enzymes. The high values of urease, β-glucosidase, amylase, N-acetyl-glucosaminidase and FDA hydrolysis activities were associated with soil under midway between trees (BH and HH). The soil under midway between TPN affected tree and healthy trees (BH) had the highest expression of xylanase and cellulase.

Figure 6  Principal component analysis showed the relationship between soil enzyme activities and sampling zones with near collar of TPN affected tree (B), near collar of healthy tree (H), midway between TPN affected tree and healthy tree (BH) and midway between healthy trees (HH) in dry season.
Conclusion

In contrast with rainy season, in dry season, PCA showed that xylanase and cellulase activities in soil with TPN affected trees (BH and B) were higher than with healthy trees (HH and H). This indicated that soil enzyme activities may be not greatly affected by tannins. At the time of taken samples, the *H. brasiliensis* shed leaves and is not collected latex, the trees are not stressed by tapping and thus, the tannins may be not stimulated. In addition, the concentration of soil tannins in dry season may be not enough to inhibit enzyme activities because tannins, the water soluble polyphenols, are readily leached and dispersed into soil by rainfall (Hättenschwiler and Vitousek, 2000; Teklay, 2004).
III.1.6 TPN AND FUNGAL COMMUNITY

These results have been submitted to “Environmental microbiology”.

The effect of trunk phloem necrosis on the structure of the fungal community in soil under rubber trees (*Hevea brasiliensis*)

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Summary

Trunk Phloem Necrosis (TPN) is an economically important disease of the rubber tree (*Hevea brasiliensis*) because it obstructs the latex flow. TPN is considered to be a physiological plant disorder caused by a combination of exogenous and endogenous stresses. Environmental stresses can affect the soil ecosystem and may also include plant stress. This study of the fungal community composition was carried out on soil in rubber plantations with a high incidence of trees affected by TPN to obtain better understanding of the impact of TPN on soil fungi. The analysis of culturable soil fungal populations revealed that the fungal density near trees affected by TPN was not significantly different from that near healthy trees. The analysis of the fungal diversity using a molecular approach (fungal 28S DNA-DGGE) with DNA extracted directly from the soil showed that fungal diversity near trees affected by TPN tended to be higher than near healthy trees. The structure of the soil fungal community was determined from a similarity dendrogram of 28S rDNA-DGGE banding patterns. There were two distinct clusters: near trees affected by TPN and near healthy trees. This suggested that there was a significant correlation between certain fungi and trees affected by TPN. The effect of TPN on soil fungi is discussed.

Keywords: Rubber tree; Trunk phloem necrosis; Plant physiological disorder; fungal community.

Introduction

The rubber tree (*Hevea brasiliensis*) is an important source of industrial rubber. However, rubber tree trunk phloem necrosis (TPN) has become a major problem as it prevents latex production, which has had a serious effect on the economy. TPN is considered to be a plant disease caused by a physiological disorder as some etiologists have reported that it was not possible to detect biotic pathogens using either conventional (Peyrard *et al.*, 2006) or molecular methods (Pellegrin *et al.*, 2007). Exogenous and endogenous stresses may cause this syndrome (Nandris *et al.*, 2004) and stressed trees may also affect the soil ecosystem. Interaction between plants and microorganisms is an important part of a forest soil ecosystem. Soil quality depends to a considerable extent on soil microbes which are of fundamental importance for soil functioning and fertility (Elliott *et al.*, 1996). Soil microbiological parameters can, therefore, be used as indicators of disturbance to soil ecology (Dick, 1992; Dilly and Blume, 1998).
Fungi are heterotrophic organisms which play a role in soil nutrient cycling, because their extracellular enzymes degrade the organic matter that is the source of nutrients for them as well as for plants (Khaid et al., 2006; Pavrina et al., 1999; Tribak et al., 2002; Vidal et al., 1999). However, the decomposition capacity of fungi depends on the type and abundance of organic materials (Saliha et al., 2005). Some fungi are able to convert toxic chemicals (such as cyanide) in soil into essential nutrients for plants (Maier-Greiner et al., 1991). Fungi can also regulate soil physical properties as the stability of soil aggregates is affected by fungal hyphae (Harris et al., 1964; TonThat and Cochran, 2000): this can assist water movement in soil and encourage root growth. Soil fungal diversity is influenced by various factors including the soil management system (Lucasa et al., 2007), climate change (Freya et al., 2008) and soil pollution (Kandeler et al., 2000). The structure of the soil fungal community is, therefore, considered to be important for understanding the relationship between environmental factors and ecosystem functioning.

The effect of various factors on the composition of the soil microbial community can be studied using culture dependent and culture independent techniques. However, culture dependent methods still have limitations because many microorganisms in the soil cannot be cultured in synthetic media in laboratory conditions (Borneman et al., 1996; Torsvik et al., 1998; van Elsas et al., 2000). Culture independent methods have been developed to study these microorganisms in their environment. Molecular biology techniques are effective for detecting both culturable and non-culturable microorganisms in the microbial populations present in soil. Denaturing gradient gel electrophoresis (DGGE) has been used widely to study the soil microbial community. This technique involves amplifying the target-DNA region and PCR products of similar length are separated in polyacrylamide gel containing a gradient of denaturing conditions (Kirk et al., 2004; Nakatsu, 2007). The DGGE technique is a reliable, reproducible and rapid method for studying soil microbial diversity (Duineveld et al., 2001; Maarit-Niemi et al., 2001). This technique is used for studying the soil microbial communities associated with agricultural practices (Hagn et al., 2003; Nakatsu et al., 2000; Sekiguchi et al., 2008; Siddique et al., 2005), degradation substrates (Cosgrove et al., 2007; Das et al., 2007; van der Wal et al., 2006) and plant genotypes (Ikeda et al., 2006)

There is currently no information on how the soil fungal community responds to TPN. However, an examination of this process would be useful to show how trees affected by TPN influence soil fungi as this may have a long-term effect on soil ecology. This study was designed to provide information on the fungal community structure in soil under rubber trees affected by TPN, using both culture dependent and molecular approaches.
2. Methods and materials

2.1 Field site description

The field site was a 9 year-old rubber plantation in Buri Ram province, north east Thailand (15° 16’ 18” N and 103° 0’ 6” E). The plantation was planted with trees of variety PRIM600 and had a high incidence of TPN (20-25% of trees affected). The climate is tropical with a mean annual temperature of 26°C and mean annual rainfall 1207 mm (very low for a rubber plantation). The soil was sandy loam (67% sand, 24% silt and 9% clay) with low organic matter content (0.88-1.34%).

2.2 Soil sampling design and preparation

The soil samples were collected during the rainy season (July to October 2006). Trees affected by TPN and healthy trees were selected using the annual tree health survey carried out by D. Nandris. Ten plots were chosen and samples were taken from four different zones in each plot: near to trees affected by TPN (B), half-way between affected trees and healthy trees (BH), near to healthy trees (H) and half-way between healthy trees (HH). In each zone, a block of soil (12x12x10 cm) was taken and separated into three layers (0-2 cm, 2-5 cm, and 5-10 cm). The soil was air-dried, ground and sieved to 2 mm. Plant residues and roots were removed. The samples were prepared for fungal analysis by mixing soil from the same depth in three identical zones.

2.3 Soil fungal enumeration and isolation

The soil fungi were counted using a plate dilution method adapted from Diouf et al. (2005). The soil (5g) was agitated in 50 ml of dispersing solution (6g/l of sodium pyrophosphate and 1.2 g/l of bactopeptone) for 30 minutes and allowed to precipitate for 15 minutes. The fungal suspensions were diluted to 10% in a physiological solution (9g/1 of NaCl). 200 µl of each diluted fungal sample was cultured on Sabouraud medium (with 0.005% w/v of Chloramphenicol – Sigma). The assay was performed in triplicate. The size of fungal colonies was determined after incubation at 27°C for 5 days. The distinct colony morphotypes were isolated and purified on Sabouraud medium. The mycelia of selected fungi were conserved at -20 °C for molecular identification.

2.4 Identification of culturable fungi

The taxonomic position of selected fungi was identified by molecular biology. DNA was extracted from the mycelium using the method described by Di Battista (Di Battista,
1997). A 650 bp nucleotide fragment (ITS1-5.8S-ITS2 region) was amplified by PCR using primer sets ITS1 (5’TCCGTAGGTGAACCTGCGG 3’) and ITS4 (5’ TCCTCCGCTTATTGATATGC 3’) (White et al., 1990). PCR amplification was performed using Ready-To-Go Taq polymerase (Pharmacia). The PCR cycles were initial denaturation at 94°C for 5 minutes, 29 cycles at 94°C for 45 seconds, 60°C for 45 seconds, 72°C for 1 minute and a final elongation step at 72°C for 30 seconds using GenAmp PCR System 2400 (Perkin-Elmer). All PCR products were fingerprinted by electrophoresis using 5 µl of product in 2% (w/v) agarose gel with a 0.5X TAE buffer (Biorad) and ethidium bromide (0.5 µg/ml) staining. DNA sequencing was performed using Genome Express (Grenoble, France). The DNA sequences were compared with other DNA sequences using GenBank BLAST searches.

2.5 Genetic diversity of soil fungi

Fungal DNA was extracted from soil samples as described by Porteus (Porteous et al., 1997). The 28S rDNA region was amplified using primer sets 403f (5’ GTGAAATTGTTGAAAGGGAA 3’) and 662r (5’ GACTCCTTGTTCCGTGTT 3’) with a 40 bp GC clamp (Sigler and Turcob, 2002). PCR amplification was performed using Ready-To-Go Taq polymerase (Pharmacia). The cycles were initial denaturation at 94°C for 5 minutes, 35 cycles at 94°C for 30 seconds, 50°C for 1 minute, 72°C for 2 minutes and a final elongation step at 72°C for 10 minutes using a GenAmp PCR System 2400 (Perkin-Elmer). The PCR products were fingerprinted as described above and PCR amplicons were determined for different positions by DGGE (Denaturing Gradient Gel Electrophoresis) using the DcodeTM Universal Mutation Detection System (Biorad, Richmond, CA). For DGGE, 20 µl of product was loaded onto 8% (w/v) polyacrylamide gel with a linear gradient of urea and formamide denaturants increasing from 35%-60% (100% denaturant contains 7 M urea and 40% (v/v) formamidine). Gel electrophoresis was carried out at 20V for 10 minutes then 75V for 16 hours. Gels were run in 0.5X TAE at a constant temperature of 60°C. The gels were stained using an ethidium bromide solution and photographed under UV transillumination using GelDoc 2000 system (Biorad).

2.6 Statistical analyses

The mean soil fungal densities in the four different zones were compared and the statistical difference was measured by the ANOVA-Tukey test using the R program version 2.7.0. The 28S rDNA DGGE banding patterns were analyzed based on the band appearance
and disappearance using image analysis software Quantity One (Bio-rad). Banding data from the DGGE gel was used to evaluate the genetic similarity of fungi from the four different sampling zones. The genetic similarity of fungal 28S rDNA was measured using Dice coefficients and a similarity dendrogram was created using UPGMA (Unweighted Pair Group Method of Arithmetic means) and Quantity One, version 4.2.1.

3. Results

3.1 Soil fungal enumeration and selected strains

The fungal community was evaluated using a plate dilution method. The number of culturable fungal strains in zone H were higher than in zone B but no significant difference in fungal density was observed. However, the number of fungal strains in zone BH was slightly higher than in zone HH but, again, there was no significant difference in fungal density (Table 1).

Five fungal strains were selected based on their abundance in the soil under trees affected by TPN (B and BH). The selected fungi were identified by comparing the DNA sequence with the Genbank database using the Blast program. This revealed that Paecilomyces lilacinus and Thielavia hyrcaniae were dominant in zone BH while Trichoderma asperellum and Corynascus kuwaitiensis were dominant in zone B (Table 2).
Table 1. Number and density of fungi in soil in near the collars of trees affected by TPN (B), midway between diseased trees and healthy trees (BH), near the collars of healthy trees (H) and midway between healthy trees (HH).

<table>
<thead>
<tr>
<th>Sampling zones</th>
<th>Number of fungal strains</th>
<th>Density (Log_{10} CFU/g of dry weight soil)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>25</td>
<td>3.46 ±0.22 a</td>
</tr>
<tr>
<td>BH</td>
<td>22</td>
<td>3.33 ±0.27 a</td>
</tr>
<tr>
<td>H</td>
<td>30</td>
<td>3.55 ±0.22 a</td>
</tr>
<tr>
<td>HH</td>
<td>19</td>
<td>3.40 ±0.20 a</td>
</tr>
</tbody>
</table>

Means within the same row followed by different letters are significantly different using Tukey HSD test (P<0.05) (+: standard deviation, n=9)

Table 2. Proportion of each selected fungal strains out of the total number of fungi and their taxonomic position by matching ITS regional sequence in GenBank

<table>
<thead>
<tr>
<th>Strain</th>
<th>Proportion of total number of fungi (%)*</th>
<th>Referenced in GenBank</th>
<th>Similarity (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>B</td>
<td>BH</td>
<td>H</td>
<td>HH</td>
</tr>
<tr>
<td>P</td>
<td>27</td>
<td>32</td>
<td>0</td>
</tr>
<tr>
<td>C</td>
<td>4</td>
<td>0</td>
<td>1</td>
</tr>
<tr>
<td>Th</td>
<td>3</td>
<td>12</td>
<td>0</td>
</tr>
<tr>
<td>Tr1</td>
<td>13</td>
<td>0</td>
<td>0</td>
</tr>
<tr>
<td>Tr2</td>
<td>12</td>
<td>4</td>
<td>0</td>
</tr>
</tbody>
</table>

*The proportion of selected fungi near the collars of diseased trees (B), midway between diseased trees and healthy trees (BH), near the collars of healthy trees (H) and midway between healthy trees.

3.2 Genetic diversity of soil fungi

The total DNA was extracted directly from the soil under rubber trees and was pure enough for PCR amplification of 28S rDNA fragments using primers 403f and 662r. The PCR products were separated by DGGE so that the genetic diversity in soil near trees
affected by TPN (B and BH) and healthy trees (H and HH) could be compared. The DGGE banding patterns were obtained by migrating PCR products in gel. The DGGE patterns in each soil layer showed that the fungal diversity differed slightly between the zones (Fig. 1). The DGGE profiles of soil at a depth of 0-5 cm showed that the genetic diversity of fungi in zone B (37 bands) was higher than in zone H (33 bands). The fungal soil in zone BH (31 bands) had greater genotypic diversity than in zone HH (27 bands) (Fig. 1a). The DGGE profiles of soil at a depth of 5-10 cm revealed that the genetic diversity in trees affected by TPN (34 bands in zone B and 30 in zone BH) were higher than in healthy trees (28 bands in zone H and 29 in zone HH) (Fig. 1b). The DGGE profiles of soil at depths of both 0-5 cm and 5-10 cm showed that some bands of PCR products had migrated a similar distance in all zones, indicating the presence of soil fungi common to all zones.
Fig. 1. Denaturing gradient gel electrophoresis (DGGE) banding profiles of fungal 28S rDNA from soil at a depth of 0-5 cm (a) and 5-10 cm (b) near the collars of trees affected by TPN (lanes 1-3), midway between diseased trees and healthy trees (lanes 4-6), near the collars of healthy trees (lanes 7-9) and midway between healthy rubber trees (lanes 10-12).
Fig. 3. The similarity dendrograms of the 28S rDNA-DGGE banding profiles of four different zones (B: near the collars of trees affected by TPN and BH: midway between diseased trees and healthy trees, H: near the collars of healthy trees, HH: midway between healthy trees) at a depth of 0-5 cm (a) and 5-10 cm (b). The dendrograms were constructed using Dice’s coefficient and clustered by the unweighted pair group method using arithmetic averages (UPGMA).
3.3 Structure of soil fungal community

The cluster dendrogram was created by UPGMA using 28S rDNA-DGGE banding patterns. The dendrogram was used to describe the similarities between the soil fungal communities in four zones (B, BH, H and HH). The fungal communities near trees affected by TPN-trees were closer to each other than they were to those near healthy trees. The similarity dendrograms clearly showed two distinct clusters in soil at a depth of both 0-5 cm and 5-10 cm. The UPGMA dendrogram showed that the fungal community in soil near to trees affected by TPN (B and BH) was distinct from that near to healthy trees (H and HH) with 0.67 similarities for soil at a depth of 0-5 cm and 0.72 similarities for soil at a depth of 5-10 cm (Fig. 2). Clustering indicated that the composition of soil fungal communities near healthy trees was dissimilar from that near to trees affected by TPN.

4. Discussion

This is the first study on the composition of fungal community in soil in rubber plantations with a high incidence of TPN. The soil fungal community was used as a biological indicator of ecological change because it responds to environmental disturbance (Dick, 1992; Dilly and Blume, 1998; Houston et al., 1998; Kasela et al., 2008; Lodge, 1997; Persiani et al., 1998). The fungal communities in soil at a depth of 0-10 cm under healthy trees (H and HH) and diseased trees (B and BH) were compared. The fungal populations in each zone were analyzed using the culture-dependent plate-counting method. Although fungal diversity was highest in zone H, the density was not significantly different from that in other zones. This may be due to the propagation capacity of fungal species. Some fungi such as *Paecilomyces* sp. and *Trichoderma* sp. can produce propagules rapidly and easily. The method of fungal enumeration used in this study may include the distribution of mycelia or spores and thus overestimate these species by comparison with others. However, in this case, the fungal diversity in areas with a low abundance of organic resources (B and H) was higher than in areas with a high abundance of resources (BH and HH). Because soil with abundant substrates (litter or dead plant materials) influences microbial growth by providing nutrient sources, microbial populations with high growth rate and activity can colonize quickly more than microbial populations with low growth rate and activity. In the absence of litter or dead plant material, some soil microorganisms have a high potential for seeking food. For instance, some fungi of the *Paecilomyces* genus are nematophagous (Bhat et al., 2009; Carneiro and Cayrol, 1991) (Gaspard et al., 1990; Hewlett et al., 1988). Many studies have reported that some of the *Trichoderma* genus are parasitic on nematodes and fungi (Elad and Kapat, 1999;
Nagayama et al., 2007; Sharon et al., 2007). Fungi with a low colonization capacity and low activities may, therefore, be difficult to detect by the method used in this study. As the soil with lower resources was not adequate for microbial growth, the influence of fast growing fungi was less and so it was easy to detect fungal diversity. The same fungal diversity pattern was also found using the biological molecular approach.

The fungal diversity was analyzed using PCR-DGGE on DNA directly extracted from soil, amplifying the 28S rDNA region (approximately 260 bp). The 28S rDNA-DGGE fingerprints revealed that the diversity obtained by detecting ribotypes was higher than that obtained by detecting morphotypes. Some authors have reported the detection of unculturable fungi using DDGE (Bates and Garcia-Pichel, 2009; Bougoure and Cairney, 2005; Gao et al., 2008; Li et al., 2008; Meroth et al., 2003). However, the results obtained using PCR-DGGE to study community complexity depends on the primer used (Green et al., 2004; Marshall et al., 2003; Zuccarob et al., 2003). For evaluating soil fungi, using 28S rDNA as the target gives a higher diversity than 18S rDNA (Diouf et al., 2005) and may provide higher resolution for DGGE (Bonanomi et al., 2008; Möhlenhoff et al., 2001; Zuccaro et al., 2008).

The structure of the fungal community in soil (0-5 cm and 5-10 cm depth) obtained from similarity dendrogram of 28S rDNA-DGGE fingerprints showed that the fungal groups near healthy trees (H and HH) were clearly separated from those near diseased trees (B and BH). This indicated that some fungal strains in trees affected by TPN were different from those near healthy trees. The soil microbial community within the ecosystem may be affected by many stresses including changes in environment under abiotic conditions and changes in quality resources (Brussaard et al., 2005). In this study, determining factors for the soil fungal community structure may arise from root exudates or litter component of trees. Root exudates are chemical compounds produced by plant roots which secrete into the surrounding rhizosphere. Root exudation can regulate the soil microbial community (Bais et al., 2006; Walker et al., 2003). The type and quantity of root exudates depend on many factors such as the plant species (Ström et al., 1994; Zhang et al., 2007), the plant varieties (Huang et al., 2008; Rae and Castro, 1967) and the plant development stage (Aulakh et al., 2001; Mougel et al., 2006). The chemicals released by plant roots affect the soil fungal community by both increasing and decreasing the abundance of fungal species (Broeckling et al., 2008; Kumar et al., 2007). The root exudates from trees affected by TPN may induce some new fungal species in the rhizosphere. However, the differences in chemical compounds of root exudation from healthy trees and diseased trees have not yet been demonstrated.
The analysis of plant chemical components can identify plant stresses. Many woody plants respond to stresses by producing secondary metabolites such as tannins (Hernes and Hedges, 2004; Yu and Dahlgren, 2000) which are water soluble polyphenols (Spencer et al., 1988). Tannins can be induced by several factors including abiotic factors (Carter et al., 1999) and human factors (artificial xylem injury) (Eyles et al., 2004). A recent study showed the effect of TPN on soil enzymatic activities: polysaccharidase activities including cellulase, xylanase and amylase were lower in soil under trees affected by TPN and suggested that tannins affect extracellular microbial enzymes by disrupting their activities (paper in press).

Because TPN is a plant physiological disorder which may be caused by multiple stresses (Nandris et al., 2004), a tree affected by TPN may produce tannin in at least one of its parts, such as bark, leaves or roots. Tannins do not reduce soil fungal populations (Baptist et al., 2008) but the high tannin concentration produced by some plants can inhibit fungal growth (Nichols-Orians, 1991). However, this biochemical plant defense may induce new strains or abundance of fungi in soil under trees affected by TPN that can degrade tannin, indicating a difference from soil under healthy trees. The soil microbial community composition was found to be correlated to the concentration of tannin in the soil and fungi that have the ability to degrade tannin can adjust easily to a high concentration of phenol (Bhat et al., 1997; Gamble et al., 1996; Makkar et al., 1993; Mutabaruka et al., 2007; Pinto et al., 2001). It has also been reported that some fungi in soil, such as fungal species in the Paecilomyces genus (Battestin and Macedo, 2007; Guedegbe et al., 2008; 2009; Mahendran et al., 2006) and Trichoderma (Bajpai and Patil, 1996; Bajpai and Patil, 1997) can produce enzymes to degrade tannins. These genera were dominant in soil near trees affected by TPN possibly because of their ability to tolerate the secondary metabolites of diseased trees. Furthermore, the area round diseased trees had a greater accumulation of wood from dead bark which also had a predominance of Paecilomyces, Trichoderma, Thielavia and Corynascus, these fungi being able to degrade plant residues (Eslyn et al., 1975; Gerber et al., 1997; Kluczek-Turpeinen et al., 2003; Lopez et al., 2006; Maheshwari et al., 2000; Martínez et al., 2005; Ryckeboer et al., 2003) and also to resist or degrade polyphenol.

This study suggested that TPN affected the soil fungal community. The similarity dendrogram of 28S rDNA-DGGE explained the structure of the fungal community composition indicating that some fungi were positively correlated in the soil round trees affected by TPN. The secondary metabolites produced by diseased trees may affect fungal populations. Little is known about the interaction between biochemical plant defenses and
diseased trees and soil fungal populations. It would be useful to study the secondary metabolites generated by rubber trees affected by TPN.

5. Conclusions

The fungal UPGMA dendrogram of 28S rDNA-DGGE gel showed that the cluster of fungal populations in soil under trees affected by TPN was distinct from that under healthy trees. Although the fungal density in diseased trees was not significantly different from healthy trees, the number of fungal strains in diseased trees (BH) was slightly higher than in healthy trees (HH) using both culture-dependent and molecular approaches. This may indicate that some soil fungi in diseased trees were different from those in healthy trees. Some fungi including *Paecilomyces* sp. *Thielavia* sp. and *Trichoderma* sp. were dominant in soil under trees affected by TPN. These fungal species may be used as biological indicators of the impact of TPN on soil ecology. This study suggested that TPN may have a long-term effect on the fungal community.
III.1.7 DISCUSSION

III.1.7.1 TPN and soil properties

In this study, organic carbon, total N, C:N ratio and organic matter content were investigated for indicating status of soil fertility. The soil under midway between healthy trees (HH) had fertility higher than under midway between TPN affected tree and healthy trees (BH). However, no significant difference was observed. This may occur from the area under midway between TPN affected tree and healthy trees (BH) have only one TPN affected tree among healthy trees and thus, ratio of TPN affected tree and healthy tree is 1:4. In his ratio, the expression of impact of TPN disease may be not distinctly. Thus, the impact of TPN on soil biological activities and soil chemical properties was unclear. However, the impact of TPN on soil ecology may express in the rainy season more than in the dry season because the trees are continuously stressed by tapping during the rainy season. The TPN, caused of damaging phloem system, may directly affect translocation soluble organic materials to all parts of tree. When the phloem system is dysfunctional, the organic compounds in parts of tree (especially, litter) may be reduced. The litter quality depends on chemical component in leave (Bardgett and Shine, 1999; Warren and Zou, 2002; Abraham and Chudek, 2008). Thus, the litter quality of TPN affected tree may lower than healthy tree and consequently, soil fertility rather low in area under TPN affected tree.

III.1.7.2 Impact of trunk phloem necrosis on macrofauna community

The macrofauna diversity has used as biological indicator of soil disturbance because it reflects the impact of environmental change (Nahmani et al., 2006; Velasquez et al., 2007; Yankelevich et al., 2007). In the rainy season, most invertebrates were associated with areas between trees (BH and HH) more than areas near collars of trees (B and H). The areas between trees in this plantation were accumulated with organic materials, especially, litter. The higher food source influences macrofauna population and diversity. In the rainy season, the difference in Coleoptera abundance between BH and HH was not obviously different but the diversity was the highest in area between healthy trees (HH). Likewise, the earthworm population was mostly found in soil under between healthy trees (HH). For the tropical forest, the quality of litter has been showed to affect macrofaunal population (Zou, 1993; Mboukou-Kimbatsa et al., 1998; Gonzalez and Zou, 1999). The quality of plant materials depends on
chemical component which is the important factor to regulate the macrofaunal abundance and diversity. As mentioned above, the TPN affected tree may contain tannins more than healthy tree. The exceeding tannins can adversely affect some invertebrates. Tannins are binding protein agents, which are able to bind digestive enzymes to limit assimilation in insects. This has effect on survival and growth of macrofauna. Consequently, macrofaunal abundance and diversity was low (Loranger-Merciris et al., 2007). However, (Ayres et al., 1997) suggested that the same tannin had different effect on different herbivore insects. In this plantation, Coleoptera genus *Cerambycidae, Elateridae, Scarabaeidae* and *Dytiscidae* were not found in both soil and litter under TPN affected trees (B and BH). These Coleoptera genera may be affected by tannins from trees affected by TPN. On the other hand, some insect species in Carabidae and Tenebrionidae were detected in all sampling zones. These macrofaunas may resist polyphenols.

The termite density was higher under TPN affected trees (B and BH). Severe TPN causes the bark to slough off at the collar. As only healthy bark containing latex and cyanide is able to block boring insects, the unprotected wood can be attacked by termites to become part of their habitat. Consequently, the zones with the highest termite population were strongly associated with areas under the affected trees.

Soil macrofauna diversity in the dry season was lower than in the rainy season. The Coleoptera was detected in only soil under between TPN affected tree and health trees (BH) but its diversity was very low, only Tenebrionidae was found. This may be due to climate condition. The climate is considered to be the most important factor in the regulation of macrofauna community. In drought circumstance, soil was arid and hard because of decreasing soil water levels. Soil moisture, which is greatly associated with seasonal variation, significantly affects soil macrofauna (Sroka and Finch, 2006). In this season, no earthworm was found in all sampling zones because the earthworms maybe migrate to deeper soil layers. In during winter and summer, the earthworms can reach a depth of 40-45 cm and was in quiescent stage (Reddy and Pasha, 1993) whereas macrofauna was collected in a maximum depth of 30 cm in this study.

### III.1.7.3 Relationship between trunk phloem necrosis and soil enzyme activity

Principal component analysis (PCA) revealed that most of enzyme activities were low in soil under near collars of tree (H and B). These sampling zones did not accumulate leaf litter, which is a major source of soil organic matter. Plant litter is an important source of
energy for soil microorganisms (Bardgett and Shine, 1999; Ruan et al., 2005; Abraham and Chudek, 2008) which mainly produce soil enzymes (de Caire et al., 2000; Stark et al., 2008). In contrast, most of enzyme activities were high in soil under midway between trees (HH and BH). These zones were accumulated with abundant litter and had high humidity and thus the environmental conditions stimulated producing soil enzymes. There has been reported that the soil with litter addition can increase soil microbial activities (Dornbush, 2007).

In the rainy season, cellulase, xylanase and amylase activities were higher in soil under zones of healthy trees (H and HH) than under zones of TPN affected trees (B and BH). Thus, soils with healthy trees were strongly associated with polysaccharidases, which can break down carbohydrates and release glucose as a source of available energy for plants and microorganisms. On the other hand, the polysaccharidase activities were low in soils with TPN affected trees. This result may be due to the difference in chemical composition of plant materials between healthy and TPN affected trees. This difference may have an impact on the quality of litter and could, therefore, explain the differences in soil enzyme activities. Numerous studies revealed that soil enzyme activities depend on organic input. The litter quality influences various enzyme activities in litter decomposition and also results in increasing or decreasing the litter decomposition in soil (Fioretto et al., 2000; Kourtev et al., 2002; Güsewell and Freeman, 2005). In *H. brasiliensis* plantation in this study, dead plant materials or plant components (such as bark, branches, litter and latex) were the main source of organic matter decomposition. However, TPN affected tree may produce defensive chemicals and the accumulation of these chemicals in plant components may affect soil enzyme activities. Plant defensive chemicals are secondary metabolites, which are produced when plants suffer from stresses, e.g., pathogenic attack, adverse environmental factors, and mechanical and chemical injuries.

Some papers have reported that TPN may be caused by multiple exogenous and endogenous stresses (Nandris et al., 2004; 2005; Nandris et al., 2006). Tapping bark for latex collection (too frequently, badly carried out or using chemicals for latex stimulation) can conduce the cumulative stress of tree. Rubber tree can respond to such wounds or stresses by producing defensive chemicals including polyphenols and chitinase. However, some plant defensive chemicals, such as tannins, can disrupt enzyme activities which occur during the decomposition of organic input. Tannins are secondary plant metabolites, found in higher plant including woody plants, in any parts of plant such as leaves (Yu and Dahlgren, 2000), bark (Matthews et al., 1997; Hernes and Hedges, 2004) and root (Peterson et al., 1999; Kosola et al., 2004). Many forest plants produce large quantities of tannin in the litter (Kraus
et al., 2003). Tannins in plants can be induced by many factors and they have been shown to be defense mechanisms against pathogens (Hewlett et al., 1997) herbivores (Coley and Barone, 1996; Ayres et al., 1997; Nomura and Itioka, 2002), abiotic factors (Carter et al., 1999) and human factors; tannin was detected in a wound in eucalyptus caused by an artificial xylem injury (Eyles et al., 2004).

Tannins are water soluble polyphenols that can precipitate proteins (Spencer et al., 1988). Tannins can inhibit the activity of extracellular microbial enzymes (Scalbert, 1991) including polysaccharidases such as cellulase and xylanase (Gamble et al., 2000; Barahona et al., 2006). Both the quality and quantity of litter tannins can cause enzyme inhibition or reduce enzyme expression in soil (Joanisse et al., 2007) and high concentration of tannins in leaves appears to inhibit decomposition (Driebe and Whitham, 2000; Lorenz et al., 2000). It was reported that tannins are readily transferred to the soil by leaching from litter (Schofield et al., 1998). Furthermore, high levels of tannins may influence the decomposition of soil organic matter by extracellular enzymes, soil mineralization, humus formation and decreasing soil fertility (Bradley et al., 2000; Driebe and Whitham, 2000; Kraus et al., 2004; Nierop et al., 2006; Wurzburger and Hendrick, 2007).

PCA showed that N-acetyl-glucosaminidase activity was associated with area between TPN affected tree and healthy trees (BH). N-acetyl-glucosaminidase is one of chitinases that degrade chitin (Ekenler and Tabatabai, 2002). Some chitinase activity in soil may be produce by plants as a defensive mechanism. Plant chitinase has been found in plant tissues such as seed, bark, leaves and root (Albrecht et al., 1994; Hietala et al., 2004; Rakwal et al., 2004; Santos et al., 2004). This enzyme plays a role of a chemical defense mechanism (Mauch et al., 1988) and is rarely detectable in healthy plants but can be produced in reaction to damage or stress caused by various biotic or abiotic factors. Some studies have showed that H. brasiliensis can produce chitinase (Bokma et al., 2002). Hevein, a chitinase from H. brasiliensis (Gidrol et al., 1994), has been showed to be an antifungal agent inhibiting microorganisms (Giordani et al., 2002; Kanokwiroon et al., 2008) and several pathogenic fungi (Parijs et al., 1991). The chitinase level is high in laticifers producing latex but very low in healthy tissues such as leaves, stem and roots (Martin, 1991). However, chitinases in plants can increase rapidly when stimulated by phytopathogenic microorganisms, pest attacks and abiotic stress such as wounding (Clarke et al., 1994; Clarke et al., 1998; Hietala et al., 2004; Rakwal et al., 2004; Hartz et al., 2005).

In contrast, in the dry season, PCA showed that xylanase and cellulase activities in soil with TPN affected trees (BH and B) were higher than with healthy trees (HH and H).
This indicated that soil enzyme activities may be not greatly affected by tannins. At the time of taken samples, the *H. brasiliensis* shed leaves and is not collected latex, the trees are not stressed by tapping and thus, the tannins may be not induced. In addition, the concentration of soil tannins in the dry season may be not enough to inhibit enzyme activities because tannins, the water soluble polyphenols, are readily leached and dispersed into soil by rainfall (Hättenschwiler and Vitousek, 2000; Teklay, 2004). Furthermore, the areas with TPN affected trees (B and BH) were high in species richness of Coleoptera, termites and ants. These macrofaunas are epigeic species, which comminute the plant materials (such as litter and bark). The comminuted materials are readily degraded by xylanase and cellulase. Meanwhile, PCA showed that amylase activity was rather high in soil under healthy trees. This may depend on plant carbohydrate metabolism, which can be influenced by stresses. Several biotic and abiotic stresses can change the concentration of carbohydrate in plants (Jeun and Hwang, 1991; Abdalla and El-Khoshiban, 2007; Morsy et al., 2007). In this case, the TPN may involve with decreasing amylase synthesis of *H. brasiliensis*. Moreover, many fungi in soil can express the amylase activity (Domingues and Peralta, 1993). In this plantation, the numeration of fungi showed that the fungal diversity in soil under midway between healthy trees (HH) was the highest. This may result in the highest amylase activity in this sampling zone and the highest β-glucosidase and urease activities were also included.

This study shows that TPN of *H. brasiliensis* has strong long-term effects on the soil ecosystem and soil fertility. PCA showed that the soil under TPN affected trees has lower polysaccharidase activities in rainy season. This may be the result of secondary metabolites produced by affected trees. As little is known about the relationship between TPN and defensive chemicals and the impact of these on soil enzymes in *H. brasiliensis* plantation, research is required into the secondary metabolites of TPN affected tree.

### III.1.7.4 The influence of trunk phloem necrosis on soil microbial community

In this study, the assessment of soil microbial community in rain season was concentrated because soil moisture greatly influences microbial activities and propagation. In addition, the result of soil enzymes assay indicated that soil enzyme activities were greatly affected by TPN in rainy season.

Soil fungal community has been used as biological indicator of ecological change because it can respond to environmental disturbance (Dick, 1992; Lodge, 1997; Dilly and Blume, 1998; Houston *et al.*, 1998; Persiani *et al.*, 1998; Kasela *et al.*, 2008). In this
plantation, the fungal community in soil at deep layer of 0-10 cm with healthy trees (H and HH) and TPN affected trees (B and BH) were compared. Each zone, the fungal community composition were analyzed by cultivation-dependent using plate-count method. Although the fungal density between soils under healthy trees (H and HH) and TPN affected trees (B and BH) were not significantly different, the fungal diversity in soil under near collar of healthy tree (H) was the highest. This result may be involved with soil sodicity because the sodium concentration levels adversely affect soil microbial community (Rietz and Haynes, 2003; Wong et al., 2008). The soil under near collar of healthy tree (H) in rainy season was the lowest in sodium concentration whereas the fungal diversity was the highest. In contrast, the sodium concentration in this zone in dry season was the highest whereas fungal diversity was the lowest. Furthermore, no significant difference in fungal density may be due to capacity of fungal species in their propagation. Some fungi in soil such as Paecilomyces sp. and Trichoderma sp. can rapidly produce propagules. Thus using method of fungal enumeration in this study may consider the distribution of mycelia or spores and the overrating of these species were observed when compared with others. However, in this case, the fungal diversity in areas with low abundance of organic resources (B and H) was rather high than in areas with high abundance of resources (BH and HH). The soil with abundant substrates (litter or plant dead materials) greatly influences microbial growth in sense of theirs nutrient sources. In addition, soil moisture, which greatly affects microbial reproduction, was higher in soils under between trees (BH and HH) than in soil under near collars of trees (B and H). The microbial with high properties of growth rate and activity can colonize quickly more than others microbial with low properties of those. Except litter or plant dead materials, some soil microbial has high potential in seeking food. For instance, some fungal species in genus Paecilomyces are nematophagous (Carneiro and Cayrol, 1991; Bhat et al., 2009) (Hewlett et al., 1988; Gaspard et al., 1990) which can feed nematodes as theirs nutrient. Several studies revealed that some strains in genus Trichoderma were characterized as parasitism on nematodes and fungi (Elad and Kapat, 1999; Nagayama et al., 2007; Sharon et al., 2007). Thus fungi with low capacity of colonization may be difficult to detect by using method in this study. In addition, slow growing fungi in soil with lower nutrient resources or moisture may be less affected by fast growing fungi because the soil condition is not suitable for fungal colonization and consequently the most culturable fungi was easily detected.

In this study, investigation of soil fungal diversity by biological molecular approach using PCR-DGGE technique was also included. The DNA directly extracted from soil and the region of 28S rDNA (approximately 260 bp) was amplified. The 28S rDNA-DGGE
fingerprints revealed the diversity obtained by detecting ribotypes was higher than diversity obtained by detecting morphotypes. Some authors reported that unculturable fungi were detected by DDGE technique (Meroth et al., 2003; Bougoure and Cairney, 2005; Gao et al., 2008; Li et al., 2008; Bates and Garcia-Pichel, 2009). However, using PCR-DGGE technique for studying community complexity, the obtained results were various depend on primer used (Marshall et al., 2003; Zuccaro et al., 2003; Green et al., 2004). To evaluate the soil fungal community, using 28S rDNA as the target had diversity higher than 18S rDNA (Diouf et al., 2005) and may present the higher resolution for DGGE procedure (Mohlenhoff et al., 2001; Bonanomi et al., 2008; Zuccaro et al., 2008).

The relationship between fungal community and sampling zones was investigated using similar dendrogram of 28S rDNA-DGGE fingerprints. The dendrogram revealed the fungal group in soil (both 0-5 cm and 5-10 cm) under healthy trees (H and HH) was clearly separated from under TPN affected trees (B and BH). This indicated that some fungal strains in soil under TPN affected trees may be different from under healthy trees. The soil microbial community within ecosystem can change by many stresses including changing environment under abiotic conditions and changing quality resource (Brussaard et al., 2005). In this study, determining factors of soil fungal community structure may arise from root exudates or litter component. Root exudates are chemical compounds that produce by plant root and secrete into surrounding rhizosphere. Root exudation can regulate the soil microbial community (Walker et al., 2003; Bais et al., 2006). Type and quantity of root exudates depend on many factors such as plant species (Ström et al., 1994; Zhang et al., 2007), plant varieties (Rae and Castro, 1967; Huang et al., 2008) and stage of plant development (Aulakh et al., 2001; Mougel et al., 2006). The releasing plant root chemistry can positively and negatively influences soil fungal species (Kumar et al., 2007; Broeckling et al., 2008). The root exudates by TPN affected tree may induce some fungal species in growth and reproduction. For instance, Paecilomyces lilacinus, Thielavia hycaniae, Corynascus kuwaitiensis and Trichoderma asperellum were abundant in soil under TPN affected trees. Nevertheless, the differences in chemical compounds of root exudation from healthy tree and TPN-tree have not yet been demonstrated.

Determination of plant chemistry component has related plant stresses. Numerous woody plants can respond to stresses by producing some of secondary metabolites such as tannins (Yu and Dahlgren, 2000; Hernes and Hedges, 2004) which are water soluble polyphenols(Spencer et al., 1988). Tannins can be induced by several factors including abiotic factors (Carter et al., 1999) and human factor: making artificial xylem injury (Eyles et
al., 2004). However, impact of tannins on soil fungal community may be difficult to conclude. Tannins did not negatively affect on soil fungal populations (Baptist et al., 2008) but high tannin concentration, which produced by some plants, can inhibit fungal growth (Nichols-orianns, 1991). In this study, tannins may influence soil fungi and thus discrimination of fungal community between healthy trees and TPN affected trees was observed. Some fungi in soil can degrade tannins and can adjust itself to high concentration of phenol (Makkar et al., 1993; Gamble et al., 1996; Bhat et al., 1997; Pinto et al., 2001; Mutabaruka et al., 2007). In this study, some fungi in soil have ability in enzymatic production for degrading tannins such as fungal species in genus of *Paecilomyces* (Mahendran et al., 2006; Battestin and Macedo, 2007; Guedegbe et al., 2008) and *Trichoderma* (Bajpai and Patil, 1996; 1997). These fungal genera existed dominantly in soil with TPN affected tree. Furthermore, *Paecilomyces*, *Trichoderma*, *Thielavia* and *Corynascus* also have ability in degrading plant residues (Eslyn et al., 1975; Gerber et al., 1997; Maheshwari et al., 2000; Kluczek-Turpeinen et al., 2003; Ryckeboer et al., 2003; Martínez et al., 2005; Lopez et al., 2006).

This study suggested that the TPN has influenced soil fungal community. Similar dendrogram of 28S rDNA-DGGE revealed that fungal cluster under TPN affected trees (BH) was discriminated with under healthy trees (HH). The secondary metabolites and root exudation of TPN affected trees may influence fungal diversity.

### III.1.8 CONCLUSION

The soil biological activities in *H. brasiliensis* plantation seem to be affected by TPN in rainy season because of climate condition. Soil biological activity analysis carried out in this study showed for the first time that some biological activity indicators can help to distinguish between healthy trees (H and HH) and trees affected by TPN (B and BH). This distinguish seems to be useful for observing the impact of TPN on the ecology of *Hevea brasiliensis* plantations. The principal component analysis showed that High cellulase, xylanase and amylase activities were associated with healthy trees and were lower near diseased trees. PCA also indicated that N-acetyl-glucosaminidase activity was associated with the zone between healthy and affected trees (BH). In addition, macrofaunal investigation revealed that the termite density was strongly associated with TPN. Soil microbial analysis showed that *Paecilomyces lilacinus* and *Trichoderma asperellum* were abundant in soil with TPN. Furthermore, soil microbial community structure was investigated using 28S DNA-DGGE. The similar dendrogram revealed that fungal cluster of TPN affected trees (B and
BH) was clearly separated from fungal cluster of healthy trees (H and HH). It would seem that polysaccharides activities, N-acetyl-glucosaminidase activity, termite density, and *Paecilomyces lilacinus* and *Trichoderma asperellum* abundance can be used as an indicator of the impact of TPN on soil ecology. However, the production of secondary metabolites as a result of TPN must be investigated.
CHAPTER III.2
IMPACT OF SOIL ORGANIC AMENDMENTS
ON SOIL BIOLOGICAL ACTIVITIES
Abstract

Soil quality is necessary to sustain *Hevea brasiliensis* productivity. Soil amendment has been known to improve soil fertility. In this study, biological activities in soil with different amendments including no organic input, external organic input and *Pueraria* cover crop + external organic input were investigated. The biological and soil chemical parameters measured can separate soils according to their management. Most enzymes measured were high in amended soils which had high organic carbon and organic matter. Principal components analysis revealed that high xylanase and cellulase activities were associated with soil under *Pueraria* cover crop + external organic input. The highest FDA hydrolysis was the highest in soil with external organic input probably because of commercial compost added to soil. Investigating macrofauna community showed that total macrofauna abundance was the highest in plantations with organic amendment. Ants, termites, Arachnida and Diplopoda densities were higher in plantations with organic amendments than in plantation without organic amendment. Soil organic carbon and organic matter were higher in amended soils than in unamended soil and had the highest in soil with *Pueraria* cover crop + external organic input. The results show that plantation with *Pueraria* cover crop + external organic input had high enzyme activities, macrofauna density and soil chemistry. This plantation, *Hevea* tree had also the highest growth and latex yield. Thus, this study suggests that *Pueraria* cover crop + external organic input seem to be proper soil management for *H. brasiliensis*. 
Ill.2.1 INTRODUCTION

*Hevea brasiliensis* or para rubber tree is an economically important tree in Thailand because its latex can increase public and rural revenue. The natural rubber from *H. brasiliensis* is used as component of many products, especially, transportation such as tires and tire products. The *H. brasiliensis* was the first established in south of Thailand and nowadays is extended to northeast. However, many areas in northeast Thailand have the problem in low soil fertility because of soil characteristic. The soils in this region have the most proportion of sand, which is the low capacity to hold mineral nutrients and moisture existing in soil. This can affect plant productivity in both quantitative and qualitative. The *H. brasiliensis* is a humid tropical tree and good soil fertility significantly affects latex yields (Akpan *et al.*, 2007). Therefore, agricultural practices that enhance soil quality are needed particularly for this region. Improvement of soil quality has the aim in enhancing physical, chemical and biological properties of soil (Lavelle, 1988; Bandick and Dick, 1999; Lado *et al.*, 2004). Enhancing these soil properties can lead to sustainable agriculture by increasing crop productivity and maintaining long-term soil fertility.

Organic amendment can improve soil quality and sustain crop productivity by increasing organic matter. Organic manures are acknowledged that it can use to solve the adverse effects on ecosystem because it can modify the properties of soil to be favorable on living organisms in ecosystem. Soil organic amendment using external organic materials and green manure/cover crop are generally practiced in agricultural system to enhance soil fertility. Using exogenous organic materials is increasing diversity of organic resources and consequently, the biological diversities are increased. Studies have been suggested that organic manure can increase soil organic matter and provide long-term soil productivity (Chander *et al.*, 1997a; Chander *et al.*, 1998). It has been known that the organic matter in soil is important source of energy for living organisms in soil. For instance, the diversity and quantity of organic matter in soil can increase diversity and abundance of soil macrofauna. Under the cover cropping system, the density of cover plant roots increase soil pore sizes and enhance soil environment to suite for soil macrofauna (Carof *et al.*, 2007). The macrofauna is useful to soil fertility and soil structure because its activities play role in soil nutrient cycling, which result in increasing soil fertility (Lobry de Bruyn and Conacher, 1990; Lavelle *et al.*, 1993; Mora *et al.*, 2005). Additionally, the macrofauna activity can enhance soil structure including soil porosity and soil aggregate stability (Lobry de Bruyn and Conacher, 1990; Mando *et al.*, 1996; Mando, 1997; Lavelle, 1997a; Pitkänen and Nuutinen, 1998;
The good soil aeration and water infiltration contribute to plant root development and crop productivity is consequence. Furthermore, microbial activities were stimulated by soil macrofauna (Scheu, 1987; 1990; Binet et al., 1998; Holt, 1998; Daube and Wolters, 2000; Scullion and Malik, 2000; Chaou et al., 2003; Desjardins et al., 2003; Ndiaye et al., 2004; Stadler et al., 2006). Some invertebrates in soil associate with soil microorganisms. For instance, the higher nitrifying bacteria were detected in earthworm burrows (Parkin and Berry, 1999). Additionally, some microbial decomposers, which can degrade cellulose and lignin, inhabit symbiotically in guts of earthworm and termites (Rouland et al., 1988; Rouland et al., 1991; Zhang et al., 1993; Rouland-Lefevre et al., 2002; Hyodo et al., 2003). Organic amendments not only enhance soil quality but also decrease plant pest levels such as nematodes and weeds (Isaac et al., 2007).

Organic matter in soil can increase diversity and activity of soil microorganisms. However, quality of organic materials is an important to enhance soil microbial biomass. The microbial activity contributes to soil fertility (Wander et al., 1994; Nelson and Mele, 2006; Fliessbach et al., 2007; Hamer et al., 2008) because it can decompose organic matter to available forms for plants. The soil microbial activity was often measured using values of enzymes in soil. Soil enzyme activity is an important to sustain fertility of soil because it plays role in degradation of organic materials. The organic decomposition greatly enhances available nutrients and cycling of nutrients, which benefit to plant productivity (Dick et al., 1988; Bandick and Dick, 1999; Hu and Cao, 2007). However, values of enzymes in soil depend on quality and type of organic amendment because soil practices directly affect soil biomass and activities (Wander et al., 1994; Stark et al., 2007; Stark et al., 2008; Liu et al., 2009). Enzyme activities have important roles in biochemical functioning of soil including decomposition of soil organic matter and nutrient cycling (Gianfreda et al., 2005; Acosta-Martínez et al., 2007; Chen et al., 2008; Sardans et al., 2008). Additionally, the potential of soil degradation can be detected by information of soil enzyme activities (Trásar-Cepeda et al., 2000). The measurement of integrative activity of several soil enzymes is greatly efficient to predict the quality of soil because soil enzyme expression is closely related to soil physical and chemical properties, and soil microbial community (Decker et al., 1999; Andersson et al., 2004; Roldán et al., 2005; Acosta-Martínez et al., 2007; Iovieno et al., 2009). Further, change in soil quality can be predicted by soil enzymes before it is detected by other soil analyses (Ndiaye et al., 2000). Soil enzyme activity is nowadays used to monitor the quality of soil under different management because it is sensitive to change in soil environment (Acosta-Martínez and Tabatabai, 2000; Acosta-Martínez et al., 2007). In addition, soil enzyme
activity can be used as sensitive index for soil biological stress (Garcia and Hernandez, 1997). Therefore, it is useful as early indicators of biological changes (Bandick and Dick, 1999). Examination of enzyme activities affected by soil organic amendments can provide understanding the progress of ecological interaction between soil enzymes and physiochemical properties. This knowledge can be applied for soil fertility development for \textit{Hevea} plantation.

The objectives of this study were: i) to evaluate the impact of different organic amendments on soil macrofauna community and soil enzyme activities in \textit{Hevea} plantation, and ii) to examine interaction between soil biological activities and soil physiochemical properties in different soil organic amendments. Soil biological activities greatly influence sustainable agriculture because it significantly associates with soil fertility. Information of soil organic amendments of \textit{Hevea} plantation is important to long-term maintenance of soil quality and to sustain latex production.
III.2.2 EXPERIMENTAL DESIGN

Three *Hevea* plantations were selected by depending different soil managements as described in table 7. Samples were taken during the rainy season (July to October 2006) and dry season (December 2008). In plantation without organic amendment (S1), 10 sampling zones under midway between trees were selected randomly. In plantation with organic amendment (S2 and S3), each plantation, the 40 sampling zones under midway between trees were selected. The experimental design is an ordered block consisting of 20 x 20 meters.

**Table 7** Description of three experimental sites with different soil managements

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<th>S1</th>
<th>S2</th>
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<td>Date of planting</td>
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<td><em>Hevea</em> varieties</td>
<td>RRIM 600</td>
<td>RRIT 251</td>
<td>RRIT 251</td>
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<td>Before plantation</td>
<td>Sugarcane</td>
<td>Sugarcane</td>
<td>Sugarcane</td>
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<td>Soil managements</td>
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<tr>
<td>- Chemical fertilizers</td>
<td>22-78-0</td>
<td>15-7-18</td>
<td>15-7-18</td>
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<td></td>
<td>(624 kg/ha)</td>
<td>(357 kg/ha)</td>
<td>(357 kg/ha)</td>
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<td>- Organic amendment</td>
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<td>2000-2002</td>
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<td>Cover cropping with <em>Pueraria phaseoloides</em> (6.25 kg of dry seed/ha)</td>
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<td>2003 eucalyptus bark residue (62.5 t/ha)</td>
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<td>2003 eucalyptus bark residue (62.5 t/ha)</td>
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<td>2003 sugarcane filter cake (waste from sugar production) (12.5 t/ha)</td>
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<td>2003 sugarcane filter cake (waste from sugar production) (12.5 t/ha)</td>
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<td>2003 Commercial compost (made from eucalyptus residue) (223.19 kg/ha)</td>
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<td>Latex yield (kg/ha/day)</td>
<td>13.75</td>
<td>26.9</td>
<td>33.1</td>
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III.2.3 IMPACT OF SOIL AMENDMENTS ON PLANT GROWTH

The averages of perimeter of *Hevea* tree were 46.8, 45.1 and 47.2 cm under plantation without organic input (S1), with external organic input (S2) and with *Pueraria* cover crop + organic input (S3), respectively (Figure 7). The *Hevea* growth in plantation with *Pueraria* cover crop + organic input (S3) was significantly higher ($P>0.05$) than in plantation with external organic input (S2).

![Figure 7](image_url)  

*Figure 7*  *Hevea* growth under different soil amendments; without organic amendment (S1), with external organic input (S2) and with *Pueraria* cover crop + organic input (S3). Means and standard deviation ($n=40$ for S1 and $n=140$ for S2 and S3); bars with the same letter are not significantly different at $P<0.05$
III.2.4 IMPACT OF SOIL ORGANIC AMENDMENTS ON SOIL PROPERTIES

Soil properties in rainy and dry seasons were showed in table 8.

Soil pH
The assay of soil pH showed that the soils were more acidic in rainy season than in dry season. In each season, no significant difference was observed for soil with no organic amendment (S1), organic input (S2) and *Pueraria* cover crop + organic input (S3).

Soil organic carbon
In each plantation, organic carbon in soil was higher in rainy season than in dry season. In both seasons, organic carbon in soil with *Pueraria* cover crop + external organic input (S3) was the highest and soil without external organic input (S1) was the lowest.

Total nitrogen
For total nitrogen, there was no significant difference between the seasons or the amendments.

C:N ratio
In each plantation, soil C:N ratio was higher in rainy season than in dry season. In both seasons, soil C:N ratio was the highest in plantation with *Pueraria* cover crop + organic input (S3) and was the lowest in plantation without external organic input (S1).

Soil organic matter
In each plantation, soil organic matter was higher in rainy season than in dry season. In both seasons, soil organic matter was the highest in plantation with *Pueraria* cover crop + organic input (S3) and was the lowest in plantation with no organic amendment (S1).

Cation exchange capacity (CEC)
In both seasons, the cation exchange capacity was the highest in soil with *Pueraria* cover crop + organic input (S3). The cation exchange capacity of soil was not significantly influenced by tested soil amendments or by the season.
Soil moisture
In rainy season, the soil with no organic amendment (S1) had the highest in moisture and was significantly higher (P<0.01) than soil with *Pueraria* cover crop + organic input (S3) and with organic input (S2).

Clay content
In rainy season, the clay content was the highest in soil with no organic amendment (S1) and was significantly higher (P<0.01) than soil with *Pueraria* cover crop + organic input (S3). There was no significant difference in clay content for samples taken from soil with organic amendments (S2 and S3).

In dry season, the soil clay contents in various plantations were not significantly different.
Table 8  Soil characteristics in three different soil amendments; no organic amendment (S1), amended with external organic input (S2), amended with *Pueraria* cover crop + external organic input (S3) during rainy and dry season

<table>
<thead>
<tr>
<th></th>
<th>S1</th>
<th>S2</th>
<th>S3</th>
</tr>
</thead>
<tbody>
<tr>
<td><strong>Rainy season</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>4.50 (0.15) a</td>
<td>4.45 (0.26) a</td>
<td>4.55 (0.13) a</td>
</tr>
<tr>
<td>Organic carbon (%)</td>
<td>0.78 (0.08) a</td>
<td>1.04 (0.27) a</td>
<td>1.50 (0.56) a</td>
</tr>
<tr>
<td>Total nitrogen (%)</td>
<td>0.08 (0.01) ab</td>
<td>0.08 (0.01) a</td>
<td>0.09 (0.00) bc</td>
</tr>
<tr>
<td>C/N ratio</td>
<td>9.94 (0.54) a</td>
<td>14.03 (4.53) a</td>
<td>17.22 (6.52) a</td>
</tr>
<tr>
<td>Organic matter (%)</td>
<td>1.34 (0.14) a</td>
<td>1.79 (0.47) a</td>
<td>2.58 (0.97) a</td>
</tr>
<tr>
<td>Cation exchange capacity (%)</td>
<td>4.43 (0.20) a</td>
<td>4.36 (0.54) a</td>
<td>5.09 (0.96) a</td>
</tr>
<tr>
<td>Moisture (%)</td>
<td>16.22 (0.98) a</td>
<td>9.60 (0.79) c</td>
<td>11.63 (1.05) b</td>
</tr>
<tr>
<td>Clay (%)</td>
<td>9.54 (0.68) bc</td>
<td>8.65 (0.35) ab</td>
<td>7.66 (0.62) a</td>
</tr>
<tr>
<td><strong>Dry season</strong></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>pH</td>
<td>5.02 (0.14) a</td>
<td>5.41 (0.28) a</td>
<td>5.29 (0.31) a</td>
</tr>
<tr>
<td>Organic carbon (%)</td>
<td>0.71 (0.13) a</td>
<td>0.82 (0.08) a</td>
<td>0.90 (0.27) a</td>
</tr>
<tr>
<td>Total nitrogen (%)</td>
<td>0.11 (0.01) a</td>
<td>0.10 (0.02) a</td>
<td>0.09 (0.01) a</td>
</tr>
<tr>
<td>C/N ratio</td>
<td>7.68 (0.99) a</td>
<td>8.87 (3.15) a</td>
<td>10.63 (2.68) a</td>
</tr>
<tr>
<td>Organic matter (%)</td>
<td>1.23 (0.22) a</td>
<td>1.41 (0.13) a</td>
<td>1.55 (0.47) a</td>
</tr>
<tr>
<td>Cation exchange capacity (%)</td>
<td>5.20 (0.08) a</td>
<td>6.82 (0.85) a</td>
<td>7.08 (1.39) a</td>
</tr>
<tr>
<td>Moisture (%)</td>
<td>3.99 (0.58) a</td>
<td>4.04 (1.38) a</td>
<td>2.30 (0.59) a</td>
</tr>
<tr>
<td>Clay (%)</td>
<td>9.97 (1.11) a</td>
<td>8.12 (1.09) a</td>
<td>8.37 (0.55) a</td>
</tr>
<tr>
<td><strong>n</strong></td>
<td>3</td>
<td>5</td>
<td>4</td>
</tr>
</tbody>
</table>

* Mean within a column followed by same lower case letter are not significantly different at P<0.05 using Tukey HSD. Numbers in parentheses indicate standard deviation.
Principal Component Analysis

Principal component analysis on soil variables in rainy and dry seasons was showed in figure 8. In rainy season, PCA showed axis 1 and axis 2 for 53.8% and 17.3% of the variance, respectively. In dry season, PCA showed axis 1 and axis 2 for 50.5% and 24.5% of the variance, respectively. Testing the significance of groupings (sites) was carried out on 10,000 permutations and was significantly discriminated (P<0.002) in both seasons.

Principal component analysis revealed that, both seasons, soil with no organic amendment (S1) was clearly separated from the amended soils (S2 and S3) by the highest soil clay content and moisture. In the dry season, the highest total N was associated with soil under no organic amendment (S1). Furthermore, the highest values of C:N ratio, organic matter and organic carbon were associated with soil under Pueraria cover crop + external organic input (S3), particularly in rainy season. High values of cation exchange capacity (CEC) in soil were associated with plantations with organic amendments (S3 and S2 for the rainy and dry seasons, respectively).
Figure 8 Principal component analysis showed the relationship between soil variables and organic managements during rainy (A) and dry (B) seasons
III.2.5 IMPACT OF SOIL ORGANIC AMENDMENTS ON MACROFAUNA COMMUNITY

III.2.5.1 Macrofauna density

For each *Hevea* plantation, macrofauna abundance in soil was higher than in litter, except macrofauna abundance in plantation without organic amendment (S1) in dry season where macrofauna cannot be detected (Figure 9). The highest density in soils was due to the large numbers of termites and ants.

The determination of macrofauna in litter showed that in each plantation, ant group was the largest taxonomic unit, except for S1 in dry season where “others” (particularly hemiptera) is the most abundant group. Except this plot, the litter macrofauna was not greatly different: ants, Arachnida and termites densities were the highest in plantation independently of the season.

The investigation of soil macrofauna showed that macrofauna in plantations with organic amendment (S2 and S3) was higher abundant than in plantation without organic amendment (S1). Moreover, no macrofauna population in plantation without organic amendment (S1) was observed in dry season. Each plantation, proportions of termite, ant and Coleoptera were greater than others. Earthworms were collected only in rainy season and the most abundant was in plantation without organic amendment (S1).

The analysis of macrofauna by combining results between litter and soil macrofauna revealed that the macrofauna density in plantations with organic amendment (S2 and S3) was higher than in plantation without organic amendment (S1), the highest density was in plantation with *Pueraria* cover crop + external organic input (S3) in rainy season. For S2 and S3 plantation, termites, ants, Coleoptera and Arachnida were the largest taxonomic groups. Coleoptera, earthworm and “others” populations were the most present in plantation without organic amendment (S1).
Figure 9  Macrofauna densities (individual m$^{-2}$) in litter and soil under plantation with no organic input (S1), with external organic input (S2) and with *Pueraria* cover crop + external organic input (S3)
### III.2.5.2 Macrofauna diversity

Macrofauna diversity in each plantation was investigated by combining results of litter and soil macrofauna obtained during the two seasons (Table 9). Plantations with organic amendment (S2 and S3) were the highest macrofauna diversity.

Coleoptera, termites and ants were the most abundant groups and found in both litter and soil, these were studied separately. The level of genus, possible, to species was determined. The Coleoptera diversity showed that six genera were identified in plantation without organic amendment (S1) and with organic amendment (S2), seven in S3. The genus *Cerambycidae* was not found in S1 and S2 than *Dytiscidae* was not observed in plantation with external organic input (S2). For termite, two genera (*Ancistrotermes* and *Macrotermes*) were observed in all plantations.

Concerning ant diversity, 11 species were identified in plantation without organic amendment (S1), 18 in plantation with external organic input (S2), 16 in plantation with *Pueraria* cover crop + external organic input (S3). The *Odontoponera transversa, Paratrechina longicornis, Anoploepis gracilipes, Oecophylla longinoda, Camponotus* sp 1, *Camponotus* sp 3, *Camponotus* sp 5 and *Pachycondyla* sp. were found in all plantations. *Hypoponera* sp 1, *Hypoponera* sp 2, *Odontomachus* sp 1 and *Camponotus* sp 2 were detected in plantations with organic amendment (S2 and S3). The *Morphospecies* sp 2 was only found in plantation without organic amendment (S1). *Ectatomma, Odontomachus* sp 2, *Odontomachus* sp 3, *Camponotus* sp 4 and *Camponotus* sp 6 were only detected in plantation with external organic input (S2). *Pheidole* sp 1, *Morphospecies* sp 1 and *Morphospecies* sp 3 were only found in plantation with *Pueraria* cover crop + external organic input (S3).
Table 9  Diversity of macrofauna under different soil organic managements; no organic input (S1), external organic input (S2) and *Pueraria* cover crop + external organic input (S3)

<table>
<thead>
<tr>
<th></th>
<th>Litter</th>
<th>Soil</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Rainy Dry</td>
<td>Rainy Dry</td>
</tr>
<tr>
<td>S1</td>
<td>S2 S3</td>
<td>S1 S2 S3</td>
</tr>
<tr>
<td><strong>Coleoptera</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Carabidae sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Cerambycidae sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Coccinellidae sp.</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Elateridae sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Scarabaeidae sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Staphylinidae sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Tenebrionidae sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Dytiscidae sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Larva</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td><strong>Termites</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Ancistrotermes sp.</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Macrotermes sp.</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td><strong>Ants</strong></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Odontoponera transversa</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Hypoponera sp. 1</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Hypoponera sp. 2</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Ectatomma sp.</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Odontomachus sp. 1</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Odontomachus sp. 2</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Odontomachus sp. 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Pachycondyla sp.</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Platythyrea sp.</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Pheidole sp. 1</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Morphospecies sp. 1</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Morphospecies sp. 2</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Morphospecies sp. 3</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Calyptomyrmex sp.</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Paratrechina longicornis</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Anoploepis gracilipes</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Oecophylla longinoda</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 1</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 2</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 3</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 4</td>
<td></td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 5</td>
<td>*</td>
<td>*</td>
</tr>
<tr>
<td>Camponotus sp. 6</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Larva</td>
<td></td>
<td>*</td>
</tr>
</tbody>
</table>
Diversity Index

The specific data obtained on the major group were used to calculate several diversity indexes: specific richness, Shannon index and diversity Simpson index (Table 10).

The specific richness, which considers only the number of species or morphospecies, was higher in plantation with soil organic managements (S2 and S3) than in plantation without organic management (S1).

Shannon and Simpson indexes indicated that the repartition is more homogenous in S1 and S3 than in S2, which the macrofauna community was dominated by few species.

Moreover, this investigation notice that the species composition of the macrofauna is closer between S2 and S3 (55% common species) than between S1 and the two other plots (39 % for S2 and 43% for S3 respectively).

Table 10  Specific richness, Shannon index and Simpson index of macrofauna in Hevea plantation without external organic input (S1), with external organic input (S2) and with Pueraria cover crop + external organic input (S3)

<table>
<thead>
<tr>
<th></th>
<th>Specific richness</th>
<th>Shannon</th>
<th>Simpson</th>
</tr>
</thead>
<tbody>
<tr>
<td>S1</td>
<td>19</td>
<td>1,10a</td>
<td>0,98a</td>
</tr>
<tr>
<td>S2</td>
<td>26</td>
<td>0,82b</td>
<td>0,67b</td>
</tr>
<tr>
<td>S3</td>
<td>25</td>
<td>0,98ab</td>
<td>0,95a</td>
</tr>
</tbody>
</table>
III.2.6 IMPACT OF SOIL ORGANIC AMENDMENTS ON SOIL ENZYME ACTIVITIES

The expression of soil enzyme activities from samples in rainy and dry seasons was showed in table 11.

Polysaccharidases

Cellulase activity is not significantly different between the different plots neither in relation to the season. Xylanase activity is higher during rainy season and is in soils with organic amendment (S2 and S3). The amylase activity is also higher in rainy season but activity was not significantly different in soils without organic amendment (S1).

Heterosidases

The β-glucosidase activity in soil without organic amendment (S1) was significantly higher \( (P<0.05) \) than in soils with organic amendments (S2 and S3) in rainy and dry season.

N-acetyl-glucosaminidase activity is higher during the dry season but it was not significantly between the plots.

Urease

Urease activity was higher in dry season than in rainy period for all the plots. Plantation with external organic input (S2) had the highest urease activity and it was significantly higher \( (P<0.05) \) than plantations with \textit{Pueraria} cover crop + organic input (S3) and without organic amendment (S1). In rainy season, urease activity was highest in soil without amendment (S1) and was significantly higher \( (P<0.05) \) than in soil with \textit{Pueraria} cover crop + external organic input (S3).

Fluorescein diacetate (FDA) hydrolysis

In each plantation, FDA hydrolysis activity was higher in rainy season than in dry season. However, the significantly higher difference \( (P<0.05) \) was observed only in plantation without external organic input (S1). In each plantation, FDA hydrolysis activity was not significantly different among plantations.
Table 11  Soil enzyme activities (µg of product g\(^{-1}\) soil h\(^{-1}\)) in three different organic amendments; no organic amendment (S1), amended with external organic input (S2), amended with *Pueraria* cover crop + external organic input (S3) during dry and rainy periods.

<table>
<thead>
<tr>
<th></th>
<th>S1 Rainy</th>
<th>S1 Dry</th>
<th>S2 Rainy</th>
<th>S2 Dry</th>
<th>S3 Rainy</th>
<th>S3 Dry</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellulase</td>
<td>1.52 (0.33) (^a)</td>
<td>1.76 (0.53) (^a)</td>
<td>3.38 (1.42) (^a)</td>
<td>3.27 (1.06) (^a)</td>
<td>3.30 (1.46) (^a)</td>
<td>2.91 (0.85) (^a)</td>
</tr>
<tr>
<td>Xylanase</td>
<td>3.79 (0.55) (^b)</td>
<td>0.75 (0.79) (^c)</td>
<td>5.08 (1.73) (^a)</td>
<td>3.51 (1.03) (^b)</td>
<td>7.38 (3.88) (^a)</td>
<td>4.28 (1.92) (^ab)</td>
</tr>
<tr>
<td>Amylase</td>
<td>12.90 (1.13) (^a)</td>
<td>5.57 (0.56) (^c)</td>
<td>10.14 (2.61) (^ab)</td>
<td>7.21 (1.02) (^bc)</td>
<td>13.83 (2.40) (^a)</td>
<td>3.92 (2.04) (^c)</td>
</tr>
<tr>
<td>β-glucosidase</td>
<td>32.15 (10.88) (^a)</td>
<td>33.97 (8.05) (^a)</td>
<td>12.33 (4.52) (^b)</td>
<td>28.83 (3.19) (^ab)</td>
<td>8.21 (2.00) (^b)</td>
<td>15.69 (2.43) (^b)</td>
</tr>
<tr>
<td>N-acetyl-glucosaminidase</td>
<td>1.25 (0.17) (^b)</td>
<td>7.18 (0.54) (^ab)</td>
<td>0.84 (0.56) (^b)</td>
<td>9.09 (2.18) (^a)</td>
<td>1.46 (0.43) (^b)</td>
<td>5.73 (1.55) (^ab)</td>
</tr>
<tr>
<td>Urease</td>
<td>1.70 (0.29) (^b)</td>
<td>3.17 (0.13) (^a)</td>
<td>1.31 (0.20) (^bc)</td>
<td>5.51 (1.23) (^a)</td>
<td>1.00 (0.26) (^c)</td>
<td>2.42 (0.52) (^ab)</td>
</tr>
<tr>
<td>FDA hydrolysis</td>
<td>5.30 (1.22) (^a)</td>
<td>1.17 (0.14) (^b)</td>
<td>6.63 (1.69) (^a)</td>
<td>3.19 (0.78) (^ab)</td>
<td>4.79 (1.07) (^ab)</td>
<td>2.46 (0.54) (^b)</td>
</tr>
</tbody>
</table>

\(^a\) Mean within a row followed by same lower case letter are not significantly different at \(P<0.05\) using Tukey HSD. Numbers in parentheses indicate standard deviation.
III.2.7 DISCUSSION

### III.2.7.1 Soil quality and plant productivity

Soil quality is generally used to evaluate sustainable land management in agroecosystems (Carter, 2002). Soil organic matter is an important indicator for agricultural soil (Lal and Kimble, 1997). Soil organic matter is efficient storage for available nutrients (such as C, N, P and S) and it can improve soil properties, soil fertility and crop production. Thus, improving soil organic matter can improve soil quality (Reeves, 1997; Freixo et al., 2002). However, soil organic matter is influenced by agricultural management (Liu et al., 2006). In this study, the parameters for assessing soil quality including organic carbon, C:N ratio and organic matter were higher in soils with organic amendments (S2 and S3) than in soil without organic amendment (S1). This indicated that unamended soil (S1) had the lowest soil quality. The lower carbon and organic matter contents in unamended soil (S1) may due to lower organic materials in soil (Golchin et al., 1995). In contrast, addition of organic materials to soil can increase the level of soil organic matter (Carter et al., 1998). Furthermore, low residue inputs have caused rapid losses of soil organic matter (Lal and Kimble, 1997) while plant residue additions can increase soil carbon content (Bossuyt et al., 2002). In this study, the quality of organic materials in both amended soil (S2 and S3) may be higher than in unamended soil (S1). It has documented that quality of organic residue can influence the quantity of soil organic matter (Juma, 1993).

Principal components analysis revealed that soil amended with Pueraria cover crop + organic input (S3) was associated with highest organic carbon and organic matter. Studies suggested that cover crops can increase organic carbon and nitrogen because amounts of organic residue in soil were increased (McVay et al., 1989; Kuo et al., 1997; Seo and Lee, 2008). It has been reported that the legume has high aboveground biomass and foliar N content (Agamuthu and Broughton, 1985). Cover crop also increases soil organic matter due to increased plant residue addition to soil (Sainju et al., 2000).

However, using legume cover crop need long-time for improving soil quality. In this study, Pueraria cover crop was terminated when Hevea canopy was close (approximately 3 years). Studies have reported that the long-term P. phaseoloides (more than 10 years) cover crop significantly affected soil organic C and N. (Dinesh et al., 1999; Dinesh et al., 2004; Dinesh et al., 2009). Wander et al. (1994) reported that the legume cover cropped soil with 10
year-period significantly increased in soil carbon and organic matter compared to conventional soil. It has demonstrated that continuous legume cover crop can improve soil organic matter and sustain crop yield (Sainju et al., 2002; Seo and Lee, 2008). Hu et al. (1997) demonstrated that short-term cover crop affected transient soil microbial dynamics and nutrient availability. Several studies have documented that microbial biomass and nitrogen mineralization were rapidly increased in first few weeks after cover crop incorporation and then returns to pre-incorporation (Wyland et al., 1996; Lundquist et al., 1999; Jackson, 2000; Sung et al., 2008).

Increasing soil organic matter and soil organic carbon have affect on soil properties and increasing available nutrients and consequently, crop yield is improved (Berzsenyi et al., 2000; Onemli, 2004). In this study, the growth of Hevea tree was evaluated using perimeter estimation. The Hevea growth from plantation with Pueraria cover crop + organic input (S3) was significantly higher (P<0.05) than from plantation with organic input (S2). In addition, latex yield under plantation with Pueraria cover crop + organic input (S3) was the highest. It has been documented that the P. phaseoloides has ability in N-fixation (Vesterager et al., 1995). In addition, legumes foliar generally have high abundance of other nutrients such as P, K and Mg (Franchini et al., 2004; Njunie et al., 2004; Armecin et al., 2005). Therefore, the legume cover cropped soil contains the available nutrients for plant development. Studies have been reported that legume cover crop can increase crop yields (Abdul-Baki and Teasdale, 1993; Armecin et al., 2005; Venkateswarlu et al., 2007). Under plantation with Pueraria cover crop (S3) in this study, the P. phaseoloides was planted as cover crop in the interspaces of plantation in the early establishment of plantation. The trees in the early stage may be positively affected from ability of legume in N-fixation process and releasing nitrogen forms after its decomposition (Lundquist et al., 1999; Jackson, 2000). It has demonstrated that cover crop can provide nitrogen source affecting plant growth at early stage (Sung et al., 2008). Additionally, when the legume was terminated, it can release the nutrients by microbial decomposition. This can greatly contribute the growth of tree. The perfect development of trees in the early stage may have significant effect on long-term plant productivity. In this study, plantation with organic input (S2) was the older than plantation with Pueraria cover crop + organic input (S3) but Hevea growth rate and latex yield were lower. In addition, principal components analysis showed that most enzymes tested were high in soil under plantation with Pueraria cover crop (S3). This may supply nutrients available for Hevea productivity.
This study observed that plantation without organic amendment (S1) had the lowest latex yield. This may due to high incidence of trunk phloem necrosis (TPN) in this plantation. Trunk phloem necrosis is the main constraint of latex production (Nicole et al., 1991; Nandris et al., 2005). High incidence of TPN may due to coarse quality of soil. (Nandris et al., 2004; 2005) concluded that TPN disease may cause by low soil fertility which increasing the risk of the syndrome.

This investigation suggests that organic amendment can increase soil quality and using *P. phaseoloides* cover crop in the early stage of *Hevea* plantation positively affect soil fertility and potentially increase tree growth and latex yield.

**III.2.7.2 Impact of soil amendment on macrofauna community**

The principal components analysis revealed that systems with organic amendments (S2 and S3) were clearly separated from system without organic amendment (S1) in both rainy and dry seasons (Figure 10). Most macroinvertebrates were abundance in plantations with organic amendments, this due probably to the quality of organic materials adding to system. This study observed that the total macrofauna density was the highest in rainy season. It had been reported that the increasing organic matter in rainy period positively influenced macrofauna community by given new opportunities in colonization, refuge and foraging (Callisto et al., 2002). Furthermore, plantation with *Pueraria* cover crop + organic input (S3) had the highest macrofauna density. It has been documented that *Hevea* plantation with continuous *P. phaseoloides* cover crop soon restores suitable conditions for soil macrofauna (Gilot et al., 1995). In this study, the highest of macrofauna density under plantation with *Pueraria* cover crop-organic input (S3) in rainy season including, especially, ant, termite, Diplopoda and Coleoptera may be an integral part to sustain the growth of *H. brasiliensis* by improving soil structure and removing organic materials into deeper soil.
Among macrofauna taxonomic groups in this study, ants and termites were mainly abundant organisms. Density of ants was higher in plantations with organic amendments (S2 and S3) than in plantation without organic amendment (S1) in both rainy and dry season. In addition, ant diversity was higher in plantations with organic amendments (S2 and S3) than in plantation without organic amendment (S1). These may result from the higher quantity and quality of organic matter present in these plantations. The abundance of ant is beneficial to soil structure because it can enhance soil porosity, aeration, infiltration and drainage (Lavelle and Spain, 2001b). This is important for plant root development and consequently, better
plant growth. Furthermore, ant density was the highest in plantation with *Pueraria* cover crop + organic input (S3) in rainy season. Laossi et al. (2008) suggested that some legume species significantly affected ant density but it did not influence density of other fauna group. This investigation found that diversity of ants under all plantations was decreased in dry period. However, some ants species including *Odontomachus* sp1. and *Camponotus* sp1. were only found in plantation with organic amendment (S2 and S3) in both rainy and dry season. It has been reported that, the number of species of ants depended on their ability in reproductive maintenance and adaptation on unfavorable environment (Bruyn, 1999). In addition, *Anoploepis gracilipes* and *Oecophylla longinoda* were found in all plantations in both rainy and dry season but they were not found in plantation without organic amendment (S1) in dry season. These species (*Odontomachus* sp1., *Camponotus* sp1. *Anoploepis gracilipes* and *Oecophylla longinoda*) may strongly associate with quality of organic amendments.

Termites were most abundance in soils with organic amendments (S2 and S3). Few species of termites had been observed. *Macrotermes* sp. was constantly associated with amended soils (S2 and S3). In addition, *Macrotermes* sp. was not found in soil without organic amendment (S1) in dry season. This indicated that quality of organic matter in soils with organic amendments (S2 and S3) may sustain this termite species. In addition, *Ancistrotermes* sp. was found in all soils in rainy season but it was not found in dry season. The decrease of soil moisture in dry season may negatively influence this species. However, the reason for termite distribution is not clear.

The rarity of earthworms and Coleoptera were found in this study. It has been reported that organic matter was necessary to keep an earthworm population (Gilot *et al.*, 1995). Earthworm community in soil is positively correlated with the quality of mulch cover (Edwards and Lofty, 1982). During rainy period, density of earthworm and Coleoptera were the lowest in plantations with organic amendments (S2 and S3). In addition, diversity of Coleoptera was the highest in plantation without organic amendment (S1). The eucalyptus bark, which is one of organic matter sources in amended plantations (S2 and S3) in this study, may be weakly palatable for macrofauna. High lignin and phenolic compounds contents of eucalyptus may be important factor in control of macrofauna population (Ganihar, 2003). Phenolic compounds, particularly, tannins are known to be defense mechanisms against herbivores (Coley and Barone, 1996; Ayres *et al.*, 1997; Nomura and Itioka, 2002). It has been shown that among tested tree plantations, eucalyptus plantation had the lowest in abundances and biomass of soil macrofauna (Warren and Zou, 2002). In addition, low palatability of *Hevea* litter may negatively affect occurrence of epigeic earthworm species
Several studies have reported that the quality of organic material significantly affects earthworm populations (Edwards and Lofty, 1982; Zou, 1993; Tian et al., 1995; Gonzalez and Zou, 1999). Additionally, low earthworm population in plantations with organic amendments (S2 and S3) may have resulted from competition with termites (Gilot et al., 1995). Furthermore, each experimental site in this study, soil was classified as sandy. (Mboukou-Kimbatsa et al., 1998) found that the sandy soil had very few earthworms. In contrast, during dry period, the earthworm density was the lowest in plantation without organic amendment (S1) in dry season. (Lobry de Bruyn, 1997) suggested that the fallacious results on earthworm density due to sampling at inappropriate times. However, Lobry de Bruyn (1997) has been reviewed that earthworms were not convinced as indicators of soil sustainability because identical management practice can gave the resulting in different response to earthworms.

In this study, density and diversity of Coleoptera were varied change between seasons and between organic amendment systems. The distributions of Coleoptera are heterogeneous. Studies have been reported that Coleoptera abundance can alter between years (Holland et al., 1998), within fields (Thomas et al., 1998; Holland et al., 1999) and between fields (Hance et al., 1990).

The Diplopoda was presented in plantation with Pueraria cover crop + external organic input (S3) in both rainy and dry seasons while it was not found in plantation with external organic input (S2) in dry season and was not found in plantation without organic amendment (S1) in both rainy and dry seasons. It has been reported that the Diplopoda is probably associated with greater organic matter received from cover crop (Merlim et al., 2005). Vohland and Schroth (1999) reported that P. phaseoloides litter had a favorable effect on Diplopoda abundance. Furthermore, it has documented that Diplopoda generally prefer soil with higher polyphenol (Lavelle and Spain, 2001a). The polyphenol containing in eucalyptus bark residues in this study may had effect on Diplopoda. For plantation with Pueraria cover crop + external organic input (S3), the large numbers of millipedes may be due to higher substrate palatability (Warren and Zou, 2002). Studies revealed that Diplopoda is an important factor to increase nutrient mineralization (Anderson et al., 1983; Anderson et al., 1985). This study observed that Diplopoda density under plantation Pueraria cover crop + external organic input (S3) was greatly lower in rainy season than in dry season. This probably is affected by ant and termite predators, which are higher abundance in rainy season.
Principal components analysis showed that Arachnida was the most abundance in plantations with organic amendments (S2 and S3). This group may be positively associated with quality of organic amendments. It has been reported that spider density was higher in organic farming system than in conventional farming system (Feber et al., 1998). Spider community structure is depended on habitat differences (Bultman and Letz, 1982; Marc and Canard, 1997; Bonte et al., 2002). Studies have been revealed that soil organic matter had a significant influence on some species of Arachnida population (Bardgett and Cook, 1998). Arachnida can contribute to decomposition and nutrient cycling by transporting fungal propagules (Heneghan et al., 1999), comminuting organic matter (Seastedt, 1984) and mixing plant debris (Bardgett and Chan, 1999). In addition, spiders can be a good indicator group of ecological quality because they sensitive to ecological system such as vegetation structure and disturbance (Jocque et al., 2005).

When both organic amendments (S2 and S3) are compared, individual densities of main macrofauna taxonomic groups were not greatly different between plantations with organic amendments (S2 and S3). Since environment is one of important factors controlling macrofauna community (Lavelle, 1997b; Nahmani et al., 2006; Velasquez et al., 2007; Yankelevich et al., 2007), environmental condition in these plantations may be not outstandingly different. In addition, studies have been documented that macrofauna density is strongly affected by resource quantity (Vohland and Schroth, 1999; Laossi et al., 2008). At the time of sampling, the amount of organic materials adding to both plantations (S2 and S3) may be similar.

Furthermore, macrofauna communities in this study were very change between rainy and dry seasons. For example, most macrofauna groups had the highest density under plantation with Pueraria cover crop-organic input (S3) in rainy period but they had the highest in plantation with organic input (S2) in dry season. Sroka and Finch (2006) found that droughty condition greatly affected macrofauna. It has been suggested that fallacious results can be obtained when macrofauna was sampled in improper period. (Lobry de Bruyn, 1997). However, the period for using Pueraria cover crop in this study may be insufficient to significantly increase macrofauna communities. It has been revealed that long-term legume fallows had positive impact on the diversity and functions of soil invertebrate (Silesli and Mafongoya, 2006a).

In this study, macrofauna distributions were patchy. Soil property may be a cause of patchy distribution. It has been documented that low level of nutrients in sandy soils is a limiting factor for macrofauna (Mboukou-Kimbatsa et al., 1998). The macrofauna cannot
readily respond to environmental change due to low fecundity, the interpretation of data were
difficult because their distributions were patchy (Lobry de Bruyn, 1997). Additionally, the
knowledge concerning effect of environmental factors on their distributions was not enough.
It has been suggested that the measurement of macrofauna should be operated in short-term
events (weeks to months) because macrofauna are active for short periods of time (Anderson,
1995).

Furthermore, macrofauna populations in this study were very low. It has been
reported that most macrofauna populations in *Hevea* plantation were decreased after five
years because the quality of litter due to ageing of the trees (Gilot *et al.*, 1995). However,
chemical quality of organic materials (eucalyptus bark and sugarcane filter cake) needs to
obtain for understanding the role of organic materials as substrate palatability.

**III.2.7.3 Impact of soil organic amendment on soil enzyme activities**

Numerous studies have been reported that enzyme activities are associated with
increase in soil organic matter, organic C, total N content (Dick *et al.*, 1988; Chander *et al*.,
1997a; Pascual *et al.*, 1999; Acosta-Martínez *et al.*, 2007). Analysis of different enzyme
activities together can provide information of the status of soil processes (Acosta-Martínez *et al*.,
2003). This investigation, activities of enzyme tested were generally higher in soils with
organic amendments (S2 or S3) than in soil without organic amendment (S1). The higher
activities in the soils with organic amendments (S2 or S3) may be due to addition of organic
materials. The quantity and quality of organic substrates influence soil microbial processes,
which are the contributing factor in increase of soil enzyme activities (Chander *et al*., 1997b;
Chander *et al*., 1998).

Enzyme activities were strongly associated with microbial organisms, which are the
main source of enzymes in soil and act important role in organic decomposition (Barbhuiya *et al*.,
2004; Ingram *et al*., 2005; Devi and Yadava, 2006; Faterrigo *et al*., 2006; Fosu *et al*.,
2007; Lucas *et al*., 2007). Since expression of enzymes is microbial processes, the quantity of
enzyme can generally indicate the levels of soil microbial population. The FDA (3′,6′-
diacetylfluorescein) is hydrolyzed by different enzymes such as proteases, lipases and
esterases. The ability to hydrolyze FDA can provide correlation information on amounts of
microbial decomposers and activities (Brunius, 1980; Ingham and Klein, 1982; Schnürer and
Rosswall, 1982; Gaspar *et al*., 2001). Numerous studies revealed that higher soil microbial
activity caused by addition of exogenous source of organic matter (Saffigna *et al*., 1989;
García-Gil et al., 2000; Nilsson et al., 2005; Stark et al., 2007). Principal component analysis showed that the highest FDA hydrolysis activity was associated with amended soil with external organic input (S2). This indicated that population of microorganisms was the most in soil with external organic input (S2). The highest population may due to addition of commercial organic compost to soil. This compost may have large numbers of microorganisms. The FDA hydrolysis activity was decreased in dry season, probably, because this season is unfavorable for stimulation of microbial processes. However, this enzyme activity was significantly higher (P<0.05) in soils with organic amendments (S2 and S3) than in unamended soil (S1). This indicated that the higher quality of organic matter may associate with amended soil (S2 and S3) because it can protect loss of microbial processes.

Cellulose is an important carbon source for microorganisms in agricultural soil and it is degraded into glucose, cellobiose and high molecular weight oligosaccharides by cellulase. In addition, hemicellulose is a component of plant cell wall and it is degraded into glucose by xylanase, as the important carbon source. Principal component analysis revealed that high cellulase and xylanase activities were associated with soils under organic amendments (S2 and S3). High activities may due to accumulate the cellulolytic and xylanolytic materials in soil matrix, which derived from adding exogenous organic residues. Furthermore, xylanase activity was consistently the highest under soil with *Pueraria* cover crop+ organic input (S3) in both rainy and dry seasons. This result may due to *Pueraria* cover crop as the organic material. It has demonstrated that legume residue can increase soil xylanase activity (Rodíguez-Kábana, 1982).

The β-glucosidase is predominant in soil and it play important role in providing carbon source for microbes in soil. This enzyme hydrolyses cellobiose and various β-glucosides present in plant materials (Martinez and Tabatabai, 1997). This investigation showed that soil without organic amendment (S1) had the highest β-glucosidase activity. This result may due to decomposition rate of organic matter, which may be higher in soils with organic amendments (S2 and S3). The fast decomposition in organic residues may have lead to a decrease in available substrate for the β-glucosidase activity (Kourtev et al., 2002). Likewise, this study found that β-glucosidase activity was increased in dry season. It has been reported that β-glucosidase is very sensitive to change in pH (Acosta-Martínez and Tabatabai, 2000). Increasing pH in dry season may result in increasing β-glucosidase activity in this study. Since β-glucosidase is sensitive to pH change, it can be used as biochemical indicator for determining changes in ecology causing from soil acidification.
The α-amylase is produced by plants, animals and microorganisms. This enzyme hydrolyzes starch to glucose which is the carbon source for plant and living soil organisms. The amylase expression is influenced by different factors such as agricultural practices, type of plant, and environmental (Fioretto et al., 2000). This study, amylase activity lacked sufficient resolution to discriminate the differences in soil managements because values of amylase activity in soil without organic amendment were in between soils with organic amendments (S2 and S3). Furthermore, soil with *Pueraria* cover crop (S3) had the highest amylase activity in rainy season while it had the lowest in dry season.

N-acetyl-glucosaminidase is one of chitinolytic enzyme. This enzyme involve in C and N cycling in soil because it degrade chitin to amino sugars which easily mineralize into C and N in soil (Ekenler and Tabatabai, 2002). In this study, N-acetyl-glucosaminidase activity cannot discriminate between soils with organic amendments (S2 and S3) and no organic amendment (S1) because values of this enzyme of soil without organic amendment (S1) were in between soils with organic amendments (S2 and S3). The highest N-acetyl-glucosaminidase activity in soil amended with *Pueraria* cover crop + organic input (S3) in rainy season may associate with highest soil organic carbon and total N content. It has been reported that higher N-acetyl-glucosaminidase activity was associated with higher soil organic carbon and total N content (Acosta-Martínez et al., 2007). In contrast, in dry season, N-acetyl-glucosaminidase activity was the lowest in amended soil with *Pueraria* cover crop + organic input (S3). This season, the total N in soil with *Pueraria* cover crop + organic input (S3) was the lowest and this may be one of reason for the lowest values of N-acetyl-glucosaminidase in this soil.

Urease is an enzyme that catalyzes the hydrolysis of urea to \( \text{NH}_3 \) and \( \text{CO}_2 \). This enzyme is produced by large numbers of microorganisms in soil (Mobley and Hausinger, 1989). This investigation, urease activity was the least in soil with *Pueraria* cover crop + organic input (S3) in both rainy and dry seasons. High urease activity in soil with external organic input (S2) may due to addition of commercial compost, which may possibly contain high urease. It has been reported that urease enzyme was higher in the soil amended with composts than amended with fresh materials (Pascual et al., 2002 ). Furthermore, this study found that urease activity was increased in dry season. This may occur form soil pH because urease activity is influenced by pH (Byrnes and Amberger, 1989). This study, the increase in soil pH in dry season may positively affect urease activity.
Principal component analysis on soil enzyme activities with the results obtained in rainy and dry seasons were constructed (Figure 11). In rainy season, PCA showed axis 1 and axis 2 for 65.1% and 15.4% of the variance, respectively. In dry season, PCA showed axis 1 and axis 2 for 45.7% and 28.3% of the variance, respectively. Testing the significance of groupings (sites) was carried out on 10,000 permutations and was significantly discriminated (P<0.00) in both seasons.

Principal component analysis showed that, in each season, the soil without amendment (S1) was characterized by low enzymatic activities. Enzyme activities were generally high in soil amended with *Pueraria* cover crop + external organic input (S3) in rainy season and were generally high in soil amended with external organic input (S2) in dry season. This indicated that enzyme tested could vary according to the season.

For example, soil with *Pueraria* cover crop + external organic input (S3) had significantly higher (P<0.05) in amylase and N-acetyl-glucosaminidase activities than soil with external organic input (S2) in rainy season while, in dry season, it had significantly lower (P<0.05). Bandick and Dick (1999) suggested that obstacle of using biological measurements as part of an index probably occurs from seasonal variations. They found that in the legume cover crop site, β-glucosidase activity changed significantly between seasons whereas FDA hydrolysis did not change. Collins et al. (1992) observed that microbial biomass C increased in Fall and decreased in Spring. Moscatelli et al. (2005) also suggested that time is the factor mostly affecting soil biological properties and nutrients availability.
Figure 11 Principal component analysis showed relationship between soil enzyme activities and different organic managements during rainy (A) and dry (B) seasons.

The *Pueraria phaseoloides* is a leguminous and it can use as cover crop to enhance soil microbial biomass (Dinesh *et al.*, 2009). Studies have revealed that the soil with legume cover crops can increase soil enzyme activities such as β-glucosidase, cellulase and urease (Bandick and Dick, 1999; Dinesh *et al.*, 1999; Dinesh *et al.*, 2004). In this study, soil with *Pueraria* cover crop + organic input (S3) were lower in these enzymes activities than soil with external organic input (S2) in both in rainy and dry season. The period of soil amendment may be one factor as possible to explain this result. In this study, period of using *Pueraria* cover crop (approximately 3 years) may be inadequate to increase microbial
activities greater than other soils (S1 and S2). It has been reported that the soil with legume cover crop over 12 years was significantly increase in enzyme activities and microbial biomass (Dinesh et al., 2004; Dinesh et al., 2006). Bandick et al. (1999) found that enzyme activities were higher in soil with 6-year legume cover crop than soil without organic amendments.

The investigation of different enzyme activities together can provide information of status of soil processes (Acosta-Martínez et al., 2003). The assay for cellulase and xylanase activities can be used as indicator to discriminate between soil with organic amendment and soil without organic amendment. Xylanase activity may be used to monitor effect of Pueraria cover crop because it was consistently high under soil with Pueraria cover crop (S3). In addition, FDA hydrolysis activity can applied to be indicator for impact of external organic materials used in this study.

However, the monitoring enzyme activity over time need for investigating activity change and eliminate problem in seasonal change. This can obtain potential of enzyme activities to be early indicators of soil management.
III.2.8 CONCLUSION

The biochemical and macrofaunal parameters measured can separate soils according to their management. Principal components analysis revealed that most enzymes tested were higher in soil with organic amendment (S2 or S3) than in soil without organic amendment (S1). The quality of external organic materials may effect on higher enzyme activities. In this study, xylanase and cellulase activities were high in soil with *Pueraria* cover crop + external organic input (S3). This indicates that these enzymes probably associate with quality of *Pueraria* residue and they may be used as indicators for consequence of *Pueraria* cover crop. In addition, especially dry season, most enzymes tested were lower in soil without organic amendment (S1) than in soils with organic amendments (S2 and S3). The organic materials used in this study probably maintain microbial processes.

The macrofauna in this study was scanty and patchy. This may due to soil property as sandy or quality of organic materials used as unpalatable for macrofauna. However, ants, termites, Arachnida and Diplopoda were higher in plantations with organic amendments (S2 and S3) than plantation without organic amendment (S1). The infertility of unamended soil may have effect on macrofauna abundance or diversity. These macroinvertebrates may sensitive to quality of soil and thus, they probably used for indicators of organic materials used in this study.

Since most enzyme activities and macrofauna densities were high in plantations with organic amendments (S2 or S3), organic matter and organic carbon contents in soil were high. However, soil with *Pueraria* cover crop + organic input (S3) had the highest organic matter and organic carbon. These properties of soil may lead to the highest *Hevea* productivity when compared with other soils (S1 and S2).

This study suggest that using *P. phaseoloides* as cover crop combined with organic materials can increase *Hevea* productivity and sustain long-term soil fertility because of higher diversity of organic materials. The *Pueraria* cover crop may stimulate *Hevea* growth at the early stage because of ability in N-fixation. Further studies are recommended to investigate organic materials chemistry and soil microbial communities as interacting controls on decomposition rate. These researches can obtain a better understanding of role of macrofauna and microorganisms in organic materials decomposition. Furthermore, to characterize the synchronization between nutrient availability of organic materials and *Hevea* nutrient demand may provide.
GENERAL CONCLUSION
AND
FUTURE PERSPECTIVES
General conclusion

This dissertation demonstrates that trunk phloem necrosis (TPN) seems to have effect on ecosystem of *Hevea* soil (Chapter III.1). The biological activities in soil were probably affected by TPN under rainy season, which is important period for organisms and plant growths. Enzyme activities were distinguished between soil with healthy tree and with TPN affected tree when it was analyzed by principle component analysis (PCA). High cellulase, xylanase and amylase activities were associated with healthy trees and were lower near diseased trees. The lower values of polysaccharides in soil under TPN affected tree may be due to increase in plant defensive chemistry, especially polyphenol, which accumulated in plant component of TPN affected tree. Under dry period, polysaccharides activities may be not inhibited by plant defensive mechanism because *Hevea* trees were not induced the stress by tapping, and therefore higher cellulase and xylanase were expressed in soil under TPN affected tree (B and BH). PCA also revealed that N-acetyl-glucosaminidase activity in rainy period was associated with soil under midway between TPN affected tree and healthy trees (BH). This may be also affected by plant defensive process. N-acetyl-glucosaminidase is one of chitinases and many plants can produce when they are in stressed situation.

Investigation of macrofauna revealed that termite group was the most abundant in soils under TPN affected trees (B and BH). Higher population of termite under zones of TPN probably associated with higher accumulated barks, which were shredded by disease attack.

Determining numeration of fungi showed that *Paecilomyces lilacinus* and *Trichoderma asperellum* were abundant in soil with TPN. The mechanism in tannins degradation may affect on the distribution of these fungi. The structure of soil microbial community obtained from similar dendrogram of 28S rDNA-DGGE showed that fungal cluster under TPN affected trees (B and BH) was clearly separated from fungal cluster under healthy trees (H and HH). The root chemical exudates, which produced from diseased trees and healthy tree, may influence microbial community.

In investigating the soil organic amendment (Chapter III.2), several trends were observed. Cellulase, β-glucosidase, urease and FDA hydrolysis activities in soil under *Hevea* plantation with organic input (S2) were higher than with *Pueraria* cover crop+ organic input (S3). Higher expression of these enzyme activities may be due to longer-period accumulation of exogenous organic materials, which added into plantation. The xylanase activity was consistently high in soil under plantation with *Pueraria* cover crop + organic input (S3). The macrofauna community was varied by season variation. Macrofauna abundance was the
highest under plantation with *Pueraria* cover crop + organic input (S3) in rainy period but it was the lowest in dry period. Coleoptera and Diplopoda abundances seem to associate with cover cropped plantation (S3). The investigation of soil properties revealed that organic carbon, organic matter and CEC were higher in soil under plantation with cover crop-organic input (S3) than with organic input alone (S2). The productivity of *Hevea* tree including growth rate and latex yield appears to increase under plantation with cover crop-organic input (S3). The capacity in N\textsubscript{2}-fixation of *Pueraria* cover crop probably strongly influenced the trees growth at the early stage, which positively affected long-term latex production.

The organic carbon and organic matter content in soil were the lowest under plantation without organic management (S1) compared to plantation with organic management (S2 and S3) and were significantly lower than plantation with cover crop-organic input (S3). This fertility parameters of soil may indicates that the high apparition of TPN was associated with lower soil fertility. Therefore, this study can sustain the hypothesis that lower soil fertility is a partial determinant of TPN incident. Hence, this study is beneficial for *Hevea* grower and institute both of government and private. The obtained information from this study can be used as practical knowledge for development of *Hevea* production and protection of TPN apparition.

This study succeeded in the objectives and can conclude as follows:

1. The TPN disease of *H. brasiliensis* has effects on soil macrofauna, soil enzyme activities and soil microbial community in the rainy season. The low polysaccharidase (cellulase, xylanase and amylase) activities and high N-acetyl-glucosaminidase activity seem to be used as biological indicators of TPN.

2. Soil organic amendment using *Pueraria* cover crop + external organic input (which used in this study including eucalyptus barks and sugarcane filter cake) positively associate with high organic matter and organic carbon content in soil and has great influence on productivity of *H. brasiliensis* including tree growth, and latex yield.

3. The lower soil fertility, which indicated by lower soil chemical properties and biological activities, associate with high incident of TPN disease.
Future perspectives

The further investigation which would be appropriate to pursue as follows:

1. Determination of role of *Hevea* secondary metabolites and root exudates.

The data presented in this study showed that reduction of polysaccharidase activities in rainy season were strongly associated with soils under TPN affected trees. The secondary metabolites produced by TPN affected tree may be a considerable factor to inhibit polysaccharidase activities. The tapping in wet season may induce tree to produce defensive chemistry. In addition, obtained data from fungal genetic analysis revealed that some fungi probably related root chemical exudation of TPN affected tree. Investigation of secondary metabolites and root exudates compared between TPN affected tree and healthy tree can verify the findings reported in this work.

2. Investigation of chemical component of organic material.

Agricultural residues are the important source of energy for soil organisms. Chemical compound of organic material have a great potential in regulation of biological activities in soil. Knowledge of organic residue characteristic added into plantation will give a better understanding about macrofauna and microbial communities in the ecosystem.

3. Investigation of dynamic soil enzymes.

Soil enzymes are sensitive to change in biochemical component of organic matter and organic carbon but their activities were controlled by several factors. Investigating soil enzyme activities before and after organic material addition, in addition, evaluating soil enzyme activities over time (week or month) will clarify the effect of soil organic management. Long-term study may warrant the determination of ecological consequences of changes in enzyme activities and microbial community

4. Determination of microbiological role in soil.

Microorganisms have greatly efficiency in organic degradation mechanism. Determining the role of microorganisms in the biogeochemical cycling of organic material would provide information to support the soil chemical property in this work. To determine the specific microbial populations affected by both soil organic management and TPN
disease, and how these microbes associate with the rate of C and N cycling, it will be helpful to study microbial community composition and function.
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APPENDICES
Appendix 1  Density of the macrofauna (mean and standard deviation, individual m\(^{-2}\)) in four different sampling zones; near collar of affected tree (B), midway between affected trees and healthy trees (BH), near collar of healthy trees (H) and midway between healthy trees (HH) during rainy period of 2006 and dry period of 2008

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**Appendix 2** Density of the macrofauna (mean and standard deviation, individual m\(^{-2}\)) in *Hevea* plantations with different soil organic managements; no organic input (S1), external organic input (S2) and *Pueraria* cover crop + external organic input (S3) during rainy period of 2006 and dry period of 2008.

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