Biodegradation of Phenolic Compounds in Creosote Treated Wood Waste by a Composting Microbial Culture Augmented with the Fungus *Thermoascus aurantiacus*

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**Abstract:** Problem statement: Creosote is used as a wood preservative and water proof agent in railway sleepers, utility poles, buildings foundations and fences and garden furniture. It is a mixture of over 300 hydrocarbons which include 75% polycyclic aromatic hydrocarbons, 2-17% phenolic compounds and 10-18% heterocyclic organic compounds. Exposure to creosote may result in several health problems including damage to kidney, liver, eyes and skin. Potential contamination of soil and water exist from creosote treated wood from construction and demolition sites. Approach: The possibility of using an invessel composting process augmented with the ascomycetous fungus *Thermoascus aurantiacus* as a mesophilic/thermophilic bioremediation option for the degradation of phenolic compounds in creosote treated wood waste was evaluated. Results: The temperatures of bioremediation process reached thermophilic phase and the mesophilic and thermophilic lag phases were clearly identified. The moisture content decreased significantly indicating that the water produced by microbial respiration did not compensate for the water lost as vapor with the exhaust gases. Initial increases in pH due to the breakdown of organic nitrogen to ammonium and final drop in pH due to the formation of organic acids and the loss of ammonium with the exhaust gases in the latter stage were observed. Different degradation rates were observed in the mesophilic and thermophilic stages of composting. The control experiment achieved higher reductions of volatile solids, total carbon and TKN and higher degradation of phenolic compounds, cellulose and lignin, indicating a higher level of activity of microorganisms during the composting process compared with the inoculated experimental trial. The stability and maturity of the product of the control experiment were also better than those of the product from the inoculated experimental trial. Conclusion: The inoculation of the cellulolytic-thermophilic fungus *Thermoascus aurantiacus* did not accelerate the bioremediation process in degrading phenolic compounds and the fungus may have inhibited the growth and metabolic activities of composting organisms.

**Key words:** Composting process, bioremediation bioremediation, phenolic compounds, wood waste, *Thermoascus aurantiacus*, Moisture Content (MC), Polycyclic Aromatic Hydrocarbons (PAHs), Total Kjeldahl Nitrogen (TKN)

**INTRODUCTION**

Creosote is distilled from crude coke oven tar consisting of around 75% of Polycyclic Aromatic Hydrocarbons (PAHs), 2-17% phenolic compounds, 10-18% heterocyclic organic compounds and minor aromatic amines. However, because of different sources and preparation procedures in manufacturing processes, the components of creosote may vary in concentration as well as type (ATSDR, 2002). Creosote is widely used as a preservative of wood products and water-proofing agent (Bedient *et al*., 1984; CEPA, 1994; ATSDR, 2002). Creosote-treated wood has been widely used in railway sleepers, utility poles, buildings foundations, building fences, bridges stakes for agricultural and fruit products, garden furniture and outdoor recreational facilities in parks (CICAD, 2004; Ikarashi *et al*., 2005).

Creosote is toxic to human being and has carcinogenic and genotoxic potentials due to the presence of PAHs as the main component of creosote (ATSDR, 2002; CICAD, 2004). Phenolic compounds contained in creosote are also carcinogens and are toxic to aquatic living creatures (CICAD, 2004). Because of its toxicity, creosote-treated wood does not degrade easily in the environment and requires special disposal...
Methods. A possible disposal option is incineration which is problematic because of the release of components such as PAHs and furans (CICAD, 2004). The pollutants contained in the creosote-treated wood waste are also barriers to its use as landfill cover due to potential migration of contaminants into groundwater. Therefore, a proper disposal method should include an effective degradation of the pollutants in creosote-treated wood waste.

Composting (as a bioremediation technique) has the advantage of degrading wood waste. Through mineralization and humification, the wood waste can be converted into a substance rich in humus and plant nutrients while at the same time result in reducing the volume of the final product. If the contaminants in the waste are degraded during composting, the final product could be used as a soil amendment (Loser et al., 1999; Borazjani et al., 2000; McMahon et al., 2008). However, a stable product requires a long period of temperature higher than 55°C. Although the temperature during thermophilic phase of composting could reach 70°C which indicates a fast degradation of organic matter inside the composting pile, the phase typically lasts no longer than 2-3 days while a maturing phase of at least 3 weeks is required to result in a stable and mature compost product (Haug, 1993; CCME, 2005; Gajalakshmi and Abbasi, 2008). In order to accelerate the maturing process, a controlled prolonged thermophilic phase is needed.

The inoculation of cellulytic microorganisms may help to achieve a fast bioremediation process. Given that mesophilic cellulytic microorganisms would be severely deactivated under temperature higher than 37°C, thermophilic fungi or bacteria would serve as better decomposers under elevated temperature environment (Cooney and Emerson, 1964). The thermophilic ascomycetous fungus *Thermoascus aurantiacus* can produce all cellulytic enzymes required for complete degradation of cellulose to glucose (Brienzo et al., 2008). Cellulytic enzymes produced by *Thermoascus aurantiacus* (especially endo-glucanase) have been applied in industrial production because of their superior thermostability, high rates of substrate hydrolysis and stability over a wide range of pH values (Mamma et al., 2009; Yu et al., 1987).

The main aim of this study was to evaluate the possibility of using a composting process augmented with the ascomycetous fungus *Thermoascus aurantiacus* as a bioremediation option to facilitate the degradation of phenolic compounds in creosote treated wood waste. Phenolic compounds were chosen as the target contaminants because extensive studies have been conducted on PAHs, but few reports were focusing on phenolic compounds.

MATERIALS AND METHODS

Experimental apparatus: The experiments were carried out in a specially designed multiple bioreactor composting/bioremediation system (Fig. 1). Three bioreactors were horizontally fastened into a main frame. Each bioreactor was made of a polyvinyl chloride cylinder (PVC/711, IPS Corporation, Gardena, CA). The cylinder had an inside diameter of 203 mm, a length of 520 mm and a wall thickness of 5 mm which provided a space for 3.5 kg (wet-basis) of the compost mixture plus 25% of the volume as a head space. A fixed circular PVC plate of 203 mm diameter and 6 mm thickness was glued into the back end of the cylinder. A removable circular plexiglas plate of 203 mm diameter and 6 mm thickness was installed on the front end of the cylinder. A circular window of 64 mm diameter was left on the removable circular plate for sampling purposes. The window was closed with a rubber stopper (No. 13) when it was not in use. The cylinders were insulated with a 38.1 mm thick Fibreglass while the removable circular plates were insulated with a 38.1 mm thick Styrofoam layer.

Each bioreactor had three holes at the bottom which were used for supplying air to the bioreactor. A top hole was used for the exhaust gas. The pressure regulated air passed through a desiccator and then through a flow meter (No 32461-14, Cole-Parmer Instrument Company, Vernon Hills, Illinois, USA). The flow meter had a 10 cm (4-inch) scale and a range of 0.0566-0.566 m³ h⁻¹ (2-20 standard cubic feet per h.). When the gas left the bioreactor, it passed through a scrubber that contained water to get rid of aerosol and water soluble organic compounds, then through another scrubber to eliminate possible airborne PAH compounds in the exhaust gas.

![Fig. 1: The experimental set up of the bioremediation system](image-url)
Inside each bioreactor, a removable 10.5 mm diameter solid stainless steel shaft was mounted on two bearings. There were 5 stainless steel collars on the shaft (Fig. 2). A bolt of 69 mm in length and 6 mm in diameter was mounted on each collar. A thermally protected electric motor (Model No. 127P1486/B, D. C., Sigma Instruments Inc., Braintree, Mass, USA) provided power to rotate the mixing shaft at speed of 6 rpm.

The data acquisition unit consisted of a master unit (Multiscan 1200, Omega, Stamford, CT), thermocouple/volt scanning card (MTC/24, Omega, Stamford, CT), Tempview software (Omega, Stamford, CT), temperature sensors (type T thermocouples, Cole Parmer, Chicago, IL) and a personal computer (Pentium IV, 256 MB of RAM). The thermocouples were inserted through specially constructed fitting. The thermocouples on the bottom of all bioreactors were located far enough from the inlet air (65 mm away) to minimize the negative influence of inlet air temperature.

Collection and preparation of wood waste: The C and D wood waste was obtained from C and D site in Yarmouth, Nova Scotia. It was screened to remove visible non-biodegradable materials and was sieved using USA Standard Testing Sieve with 12.5 mm opening (USA Standard Testing Sieve, ATM, Milwaukee, Wisconsin). Then, the wood chips were well mixed with fresh compost (Miller Compost Corporation, Dartmouth, NS) at a ratio of 1:1. The C:N ratio was adjusted using urea [CO (NH₂)₂] and the moisture content was adjusted to 60% using distilled water.

Preparation of inoculums: Thermoascus aurantiacus (ATCC 204492) was obtained from the Manassas, Virginia and used as inoculum in one of the experiments. Potato Dextrose Agar (PDA) medium was used for slants and Petri dishes and potato dextrose broth was used as liquid medium. Both solid and broth media were made in the laboratory using distilled water and the reagents listed in Table 1. The steps for preparing the Thermoascus aurantiacus spore suspension are showing in Fig. 3. The freeze dried culture was hydrated in dextrose broth and then plated on dextrose agar. After 24 h, colonies with bright orange color start forming on the dextrose agar (Fig. 4a). Spore suspension of Thermoascus aurantiacus was prepared by cutting 1 cm² Petri dish culture into 25 mLsterile potato dextrose broth medium and incubating in an incubator (Isotemp® oven, Model 106G, Fisher Scientific, Hampton, New Hampshire) at 45°C in order to activate the fungal culture. After 48 h of incubation, plate counts were conducted to detect Colony Forming Units (CFU). An amount of 5 mLMedia containing the CFU of 2.0×10⁴ mL⁻¹ was transferred into Fernbach flasks containing 250 mLof liquid cultural media and agitated on rotary shaker (Series G-25 Incubator Shaker, New Brunswick Company, New Jersey, USA) at 120 rpm and 45°C for 48 h.

Figure 4b shows a higher magnification of asci containing ascospores. The final cultures were used as inoculums in the experiment in the amount of 10% (by weight) of composting mixture (wood waste+municipal solid waste compost).

Experimental protocol: Approximately 3 kg of the final mixture were placed in each bioreactor. Two experiments were carried out: (a) A control experiment with the no fungus Thermoascus aurantiacus added and (b) An experiment with the fungus Thermoascus aurantiacus added to the mixture. Three replicates were conducted for each experiment using the three bioreactors simultaneously. Each group of experiments lasted for 15 days. The pressure-regulated air was supplied continuously to the bottom of the bioreactor. The flow rate was measured by a flow meter (No 32461-14, Cole-Parmer Instrument Company, Vernon Hills, Illinois, USA) and adjusted to 0.05 m³ h⁻¹ (3 V V h⁻¹). Temperature was monitored throughout the process and the data was stored in a Microsoft Excel® file in the computer every 30 min. The temperature of each experimental group was then generated using average temperature from 12 thermocouples in three
bioreactors. An amount of 36 mL used cooking oil were added into the bioreactor every 12 h for the duration of each experiment as recommended by Alkoaik (2005). This was done in order to achieve a thermophilic condition of 60°C. The pH and moisture content were also monitored. After 15 days of bioremediation, the quality, degradation of cellulose and lignin, the degradation of phenolic compounds and the quality, stability and maturity of the end product were evaluated. The quality of the end product was determined by quantifying C: N ratio, pH, CO₂ evolution, phytotoxicity.

Moisture content and pH: The Moisture Content (MC) was measured following the ASTM (D4442-07) oven-drying method (ASTM, 2007a). Slurry contained about 10 g of material and 50 mL distilled water was used to measure the pH value using a pH meter (Fisher Accumet®, Model 805 MP, Fisher Scientific, Hampton, New Hampshire).

Total carbon: Approximately 1.0 g of the material was sampled for total carbon analysis. Carbon dioxide was determined with a Leco carbon analyzer (Model 516-000, Leco Corporation, St. Joseph, Michigan) along with an induction furnace (Leco HF2O Furnace, Leco Corporation, St. Joseph, Michigan) at the Mineral Engineering Centre of Dalhousie University, Halifax, Nova Scotia.

Solids: The solids analyses were performed according to the procedures described in the USEPA Method 1684 (USEPA, 2001). Ash contents were determined by burning samples in muffle furnace (Isotemp® Muffle Furnace, Model 186A, Fisher Scientific, Hampton, New Hampshire) at a temperature of 550°C for 20 min.

Total Kjeldahl Nitrogen (TKN): The Total Kjeldahl Nitrogen (TKN) was determined at Maxxam Analytical Testing Laboratory in Mississauga, Ontario, following the procedure of USEPA Method 351.2 (USEPA, 1993).

Phenolic compounds: The phenolic compounds were extracted from 3 g material with 50 mL deionized water and centrifuged for 20 min at 2400 rpm. The supernatant was vacuum filtered through a 0.4 µm polycarbonate filter paper (47 mm diameter polycarbonate filter paper, Fisher Scientific, Montreal, Quebec) as described by Carter and Gregorich (2008). The supernatant was transferred into a flask and analyzed for the presence of phenolic compounds using the 4-aminoantipyrine colorimetric test following the ASTM procedure (ASTM, 2007b) at 510 nm using spectrophotometer (Spectronic 601, Milton Roy, Ivyland, PA). A standard curve was prepared as shown in Fig. 5. The linear relationship of phenolic compounds concentration to the absorbance reading at 510 nm was described by Eq. 1 (R² = 0.9972):

\[ \text{AU510} = 0.0908 \text{PC} \]  
(1)

Where:
\[ \text{AU510} = \text{Absorbance reading at 510 nm} \]
\[ \text{PC} = \text{Concentration of phenolic compounds (mg/L)} \]
**Fig. 4:** *Thermoascus aurantiacus* (a) Colonies with bright orange colour elliptical on potato dextrose agar (b) Asci containing ascospores

**Fig. 5:** Standard curve for the phenolic compounds

### The Germination Index (GI)

The Germination Index (GI) was measured following the procedure described by Iannotti *et al.* (1994) and Jiang *et al.* (2006). About 10 g of compost sample were mixed with 100 mL distilled water. Ten cress seeds (*Lepidium sativum L.*) were evenly placed on the filter paper (Whatman® 40, Whatman Inc., Clifton, New Jersey) in a sterilized Petri dish. Then, 5.0 mL of the extract was transferred into the filter paper. Three replicates were carried out for each sample. The Petri dishes were incubated at 25°C in the dark for 48 h. The results were evaluated by counting the number of germinated seeds and measuring the length of roots. The Germination Index (GI) was determined as Eq. 2:

\[
\text{GI (\%)} = \left( \frac{\text{Seed germination} \times \text{Root length of treatment (cm)}}{\text{Root length of control (cm)}} \right) \times 100
\]

### CO₂ Evolution

CO₂ evolution was determined as described by Benito *et al.* (2003). Approximately 25 g of the material were sampled and pre-incubated at room temperature for 3 days. The moisture content was adjusted to 60% and each sample was separately sealed in containers containing a beaker with 10 mL of 1 M NaOH solution. The samples were incubated at 25°C and the CO₂ generated was determined by titrating NaOH solution with 1 M HCl solution every day for 5 consecutive days. The rate of CO₂ evolution was calculated as mg C-CO₂ g⁻¹ compost per day.

### Cellulose and lignin contents

The cellulose and lignin contents were measured following the Standard Methods published by AOAC (1980) for Acid Detergent Fiber (ADF) and Acid Detergent Lignin (ADL). Cellulose was estimated as the difference between ADF and ADL. Lignin was estimated as the difference between ADL and ash content as described by Yu *et al.* (2007).

### RESULTS

#### Temperature

The temperature profile was generated by plotting time series of the average temperature data of the control experiment and the inoculated experiment with inoculation of *Thermoascus aurantiacus* as shown in Fig. 6. The fluctuation in the temperature during the thermophilic phase (between 40 and 50°C) was due to the process of adding bio-available carbon source (used cooking oil) which required opening the system every 12 h. The peak temperatures for control and the inoculated experiments were 50 and 48.8°C, respectively. The temperature was maintained above 45°C for 59 and 99 h and above 40°C for 146 and 186 h for the inoculated and control experiments, respectively. The room temperature was stable around 22-24°C.

#### Moisture content

The moisture content decreased gradually during the bioremediation process as shown in Fig. 7. The final moisture contents for the control and inoculated experiments were 42.9 and 48.0%, respectively.

#### pH

The changes in pH are shown in Fig. 8. The initial material was acidic (a pH of 6.0). In the first week of bioremediation, the pH value increased to the basic range of 7.4-8.5 and then decreased to a weak acidic (5.5-6.8).
**Solids:** The changes in volatile solids and ash contents are presented in Fig. 9. The initial volatile solids content of the materials was 805±10 g kg\(^{-1}\) material (dry basis) which decreased slowly resulting in reductions of 4.6 and 0.1% for the control and the inoculated experiments, respectively. The ash content basically stayed constant at 195±12 g kg\(^{-1}\) material (dry basis) till the end of the bioremediation process as the variations in the final ash contents were within the experimental errors.

**Total carbon:** The changes in total carbon content are shown in Fig. 10. The initial total carbon was 392 g kg\(^{-1}\) material (dry basis). The final total carbons for the control and the inoculated experiments were 358 and 386 g kg\(^{-1}\) material (dry basis) resulting in total carbon reductions of 8.7 and 1.5%, respectively.

**Total Kjeldahl Nitrogen (TKN):** The changes in total Kjeldahl nitrogen (TKN) are shown in Fig. 11. The initial TKN was 24.94 g kg\(^{-1}\) material (dry basis) which decreased to 12.5 and 17.1 g kg\(^{-1}\) material (dry basis) for the control and the inoculated experiments resulting in reductions in TKN of 50.2 and 31.9%, respectively.

**C:N Ratio:** The changes in C:N ratio (Fig. 12) were calculated from total carbon and total Kjeldahl nitrogen data. The initial C:N ratio was 15.6:1. Due to the consumption of continuously added used cooking oil and the rapid consumption of nitrogen by microorganisms (to metabolize bio-available carbon for cell growth and energy production), the C: N ratio increased reaching 28.6: 1 and 22.6: 1 for the control and the inoculated experiments, respectively.

**Lignocellulose:** The results of the degradation of cellulose and lignin are shown in Table 2. The initial content of cellulose was 24.8±1.5% which decreased after 15 days to 19.2 and 20.2% for the control and inoculated experiments resulting in reductions of 19.8 and 20.2%, respectively. The initial content of lignin was 19.8±0.9% which decreased after 15 days to 16.3 and 17.2% for the control and the inoculated experiments resulting in reductions of 17.7 and 13.1%, respectively.

**Phenolic compounds:** The initial concentration of PC was 0.222±0.010 mg g\(^{-1}\) material (dry basis). During the bioremediation process, the concentration of PC decreased gradually in both experimental trials, reaching a final concentration of 0.058±0.006 mg g\(^{-1}\) material (dry basis) for the control and 0.071±0.005 mg g\(^{-1}\) material (dry basis) for the inoculated experiment as shown in Fig. 13. The inoculated experimental trial had a lower degradation (68.0%) of phenolic compounds than the control experiment (73.9%).

**Maturity and stability tests:** The maturity and stability of the final bioremediation product were evaluated by analyzing the CO\(_2\) evolution (as mg of CO\(_2\)-C/g VS-day and mg of CO\(_2\)-C/g C-day), the Germination Rate (GR) and the Germination Index (GI) of the final product. The results are shown in Table 3.
Table 2: Degradation of cellulose and lignin

| Trial              | Cellulose Content (% DM) | Lignin Content (% DM) |
|--------------------|--------------------------|-----------------------|
|                    | Initial | Final | Degradation* | Initial | Final | Degradation* |
| Control            | 24.8±1.5 | 19.2±0.7 | 22.6      | 19.8±0.9 | 16.3±0.3 | 17.7      |
| Inoculated experiment | 24.8±1.5 | 19.8±1.2 | 20.2      | 19.8±0.9 | 17.2±0.7 | 13.1      |

* Standard deviation: * Percent reduction

Table 3: CO₂ evolution, germination rate and germination index

| Trial              | CO₂ Evolution | GR (%) | GI (%) |
|--------------------|---------------|--------|--------|
|                    | (mg CO₂-C/g VS-d volatile solid-day) | (mg CO₂-C/g C-d carbon-day) | Initial | Final | Initial | Final |
| Control            | 3.18±0.19     | 0.89±0.41 | 0      | 90    | 0      | 20    |
| Inoculated experiment | 4.22±0.24     | 8.79±0.50 | 3      | 62    | 0      | 5     |

* Standard deviation

Fig. 9: Changes in volatile solids and ash contents during bioremediation process

Fig. 10: Changes in total carbon content during the bioremediation process

The CO₂ evolution rates from the final product were 3.18 and 4.22 mg CO₂-C/g VS-day for the control and inoculate experiments, respectively. The initial GI was 0% which improved significantly after bioremediation reaching 20 and 5% for the control and inoculated experiments, respectively.

Fig. 11: Changes in TKN content during the bioremediation process

Fig. 12: Changes in C: N during the bioremediation process

DISCUSSION

**Temperature:** The peak temperature achieved was above 50°C in the control bioreactors and 48.8°C in the inoculated bioreactors. Loser et al. (1999) used a pilot scale percolator to compost PAH-contaminated pine wood waste with liquid hog manure as a nitrogen and mineral source and obtained a maximum temperature of 42°C.
McMahon et al. (2009) composted a mixture of board waste (C and D wood waste) with poultry manure, green waste, top soil and compost and observed a peak temperature above 70°C which is significantly higher than the peak temperature achieved in this study. This could be due to the fact that more nutrients were added in the system (poultry manure), thereby increased the bio-available carbon and nitrogen. Although the addition of used cooking oil maintained the temperature within the thermophilic range for the first few days, the temperature decreased at the end of each trial indicating the decline of bio-available carbon in the composting mixture.

Chen and Chen (1988) reported a very fast growth for *Thermoascus aurantiacus* at a temperature of 40-50°C. Cooney and Emerson (1964) reported minimum, optimum and maximum temperatures for *Thermoascus aurantiacus* growth of 20, 45 and 55°C, respectively. Deploey (1995) reported minimum, optimum and maximum temperatures for the growth of ascospores of 32, 47.5 and 60°C, respectively. The temperature recorded in this study was within the optimum range of temperature for the growth of *Thermoascus aurantiacus*.

**Moisture content:** Water is the media for nutrient transportation and metabolic reactions and, therefore, the availability of nutrients and the contaminant to microorganisms is affected by the water content in their micro-environment (Golueke, 1977; Gajalakshmi and Abbasi, 2008). The optimum moisture content for metabolic activity is in the range of 50-70% (Tiquia et al., 1996; Epstein, 1997; Gajalakshmi and Abbasi, 2008). In this study, the initial moisture content of the mixture was 59.69±0.77% which was within the optimal range. However, the moisture content decreased significantly for both trials during the 15 days of bioremediation. The decreased moisture content was due to the fact that the loss of moisture with the exhaust gas was higher than the moisture produced by metabolic activity as a result of declining bioavailable carbon. According to Haug (1993) and Walker et al. (1999), an intense decrease of moisture content will reduce the metabolic rate and affects the effectiveness of the bioremediation process. If the moisture content is lower than 30%, the microbial activity will be significantly limited (Haug, 1993). The moisture contents in both trials were lower than the minimum optima for metabolic activity but still high enough (42.9-48.0%) for the bioremediation process to proceed.

**pH:** The pH of the mixture was slightly acidic at the beginning of the experiments and increased to 7.9 on the 4th day for the control and then gradually decreased back to a weakly acidic pH (6.5) by the end of the experiment. For the inoculated experimental trial, the increase in pH was not significant until the 7th day (8.5) but followed similar pattern to that of the control experiment. The changes in the pH were due to the decomposition process and the production of various byproducts. The initial nitrogen content in the material was relatively high (C: N ratio of 15.7:1) and the breakdown of organic nitrogen to ammonium resulted in the initial increase in the pH. The final drop in the pH was due to the formation of organic acids from decomposition of fats and grease (Epstein, 1997). A similar pH trend was observed by Khan and Anjaneyulu (2006) who explained the rise in pH as due to the breakdown of protein into ammonia.

The optimum pH range for the growth of *Thermoascus aurantiacus* is 3.5-4.5 (Tong et al., 1980; Upadhyay et al., 1984; Grajek, 1988; Maheshwari et al., 2000). Upadhyay et al. (1984) studied the effect of pH on the growth of *Thermoascus aurantiacus* at the optimum growth temperature of 50°C and found the fungus to grow the fastest within a pH range of 3.5-4.0 with the shortest lag occurring at pH 4.0 but the growth rate decreased when the pH was increased from 3.0 to 11.0. Maheshwari et al. (2000) reported an optimum growth of *Thermoascus aurantiacus* at pH of 4.5 which then declined with increased pH.

**Solids:** The degradation of organic matter by microorganisms during bioremediation process is the major energy source for microbial growth (De Bertoldi, 1987; Lemus and Lau, 2002). Thus, the decrease of volatile solids is common in the composting process. Saludes et al. (2007) achieved 43.89% reduction of volatile solids of dairy cattle manure after 35 days in a controlled thermophilic-mesophilic composting system.
Lu et al. (2008) reported reductions of volatile solids of 16.71-22.97% after 7 days of composting barley dregs and sewage sludge. In the current study, 36 mL of the used cooking oil was added into the system every 12 hrs as bio-available carbon which was utilized by microorganisms. The bioavailable carbon in the feedstock (cellulose material) was limited and as a result the reduction in volatile solid was small (0.1 and 4.6%). The ash contents stayed relatively constant, the changes observed in the ash were within the experimental errors.

**Total carbon:** In this study, the reduction in total solids were in the range of 0.1-4.6%. Wang et al. (2003) achieved a total carbon reduction of 14% while composting sewage sludge with solid food waste in a bioreactor for 5 days. Gomez-Brandon et al. (2008) reported a total carbon reduction of 3.5% while composting cattle manure for 15 days. Tiquia and Tam (2000) reported total carbon reductions of 50-63% in turned windrows and 30-54% in unturned windrows windrow while composting manure for 42 days. Michel et al. (1995) reported a carbon reduction of 24% while composting yard trimming waste in a bioreactor for 45 days. The higher reductions of total carbon in these studies may be due to the longer composting time and higher biodegradability of the materials used in their experiments.

The pH recorded in the study was in the range of 5.5-8.5 which is higher than the optimum pH range of Thermoascus aurantiacus. According to Upadhyay et al. (1984) higher pH will negatively impact the growth of fungus causing it to sporulate. There were significant differences in the total carbon between the control and the inoculated experimental trial. The results indicated a higher rate of decomposition in the control experiment. The reported pH is however, suitable for the mixed culture of composting microorganisms.

**Total Kjeldahl Nitrogen (TKN):** The reductions in TKN were more rapid and much higher than the reductions in the total carbon. There was significant difference between the control and the inoculated experiments. A lower reduction of TKN was observed with the inoculated experiment which is in agreement with the volatile solids and total carbon results. Zorpaş et al. (1999) reported a TKN reduction of 0.99 mg g⁻¹ (from 2.12-1.13 mg g⁻¹ dry sludge or 46%) in the sewage sludge composting process. Alkoaik and Ghaly (2006) observed a TKN reduction of 11.0% during composting of greenhouse tomato plant residues. Tiquia and Tam (2000) reported a TKN reduction of 50% during the 168 days of composting of chicken litter in forced-aeration piles. Kumar et al. (2008) reported a TKN reduction of only 0.53-0.64% during the composting of paddy straw which is low in nitrogen content.

**C: N Ratio:** The C:N ratio decreases in a biological decomposition system because: (a) the organic carbon is oxidized to CO2 faster than ammonium is oxidized to NO3- and (b) nitrogen can remain relatively stable if the balance between mineralization of organic nitrogen to NH4 and the immobilization of NH4 to organic nitrogen (microbial growth) is maintained during the process (Wang et al., 2003; Alkoaik, 2005). However, if the initial concentration of nitrogen is high, the decrease in nitrogen will surpass the decrease in total carbon resulting in a higher C: N ratio. The higher temperature and/or longer thermophilic phase will result in higher rate of organic nitrogen decomposition and increased nitrogen loss (Bishop and Godfrey, 1983; Tiquia and Tam, 2000; Wang et al., 2003).

Bioremediation/composting that starts with low C:N ratio results in significant losses of nitrogen (Tiquia and Tam, 2000; Beck-Friis et al., 2001). Tiquia and Tam (2000) reported a nitrogen reduction of 59% while composting chicken litter with an initial C: N ratio of 14.5:1. Beck-Friis et al. (2001) reported nitrogen reduction of 24-33% while composting household wastes with an initial C: N ratios of 21-23:1 under controlled conditions for 22-31 days. Due to the continuous addition of used cooking oil and the rapid consumption of nitrogen by microorganisms for cell growth, the C: N ratio increased by 83 and 45% for the control and the inoculated experimental trial, respectively.

**Lignocellulose:** The results indicated that the inoculation of cellulytic-thermophilic microorganisms had little effect on the degradation of cellulose and lignin during the bioremediation process. Yu et al. (2007) studied the degradation of lignocellulose in a laboratory composting system during which the temperature reached a peak of 65°C and reported degradation of cellulose and lignin of 11 and 18% on day 15 and 32 and 23% on day 45, respectively. Vikman et al. (2002) used controlled composting system to test the biodegradation of lignin-containing pulp and paper products and observed a lignin reduction of approximately 17% at 35°C, 25% at 50°C and 15% at 58°C on the 15th day and were 41.8% at 35°C, 39.6% at 50°C and 24.8% at 58°C on day 45. The current study resulted in similar degradation of cellulose and lignin but in a shorter period of time. This may be due to the addition of bio-available carbon used cooking oil.
Thermoascus aurantiacus has been shown to grow efficiently on lignocellulosic biomass as it is capable of producing all cellulolytic and hemicellulolytic enzymes (cellulases, hemicellulases, amylases, pectinases) required for the complete degradation of cellulose and hemicellulose to glucose (Da Silva et al., 2005; Kalogeris et al., 2003; Milagres et al., 2004; Santos et al., 2003; Yu et al., 1987). These enzymes operate at a temperature of 50-55°C and acidic pH (Yu et al., 1987; Adams, 1992). The lack of degradation of cellulose in the experiment augmented with Thermoascus aurantiacus could be the result of higher pH which may have retarded the production of cellulolytic and hemicellulolytic enzymes required for the complete degradation of cellulosic material to glucose.

**Phenolic compounds:** About 73.9 and 68.0% of Phenolic Compounds (PC) were degraded after 15 days of bioremediation in the control and the inoculated experiments, respectively. The higher final phenolic compounds observed in the inoculated experimental trial was the result of the lower temperature and shorter thermophilic phase observed with this trial. This is in agreement with reduction in the total carbon, nitrogen and volatile solids.

The degradation of phenolic compounds in wood substrate was reported by Galli et al. (2008) who used white-rot fungus Pleurotus ostreatus to degrade phenolic compounds in creosote-treated wood and observed a decrease of phenol compounds greater than 75% after 30 days. In the present study, the 68-73% degradation of PC was achieved in a much shorter period of time. Khan and Anjaneyulu (2006) achieved complete phenolic compounds and benzene degradation after 40 days of composting contaminated soil and sediment.

Stoilova et al. (2008) studied the biodegradation of mixtures of phenolic compounds by monocultures and mixed culture of Thermoascus aurantiacus and Aspergillus awamori and reported a negative interaction between the studied species. Higher reductions of phenolic compounds were obtained with the mono cultures compared to the mixed culture. There have been also some reports on the toxicity of the enzyme xylanase produced by the fungus Thermoascus aurantiacus on animals (Bedford, 2000; Kongbuntad et al., 2006). There is a possibility that this enzyme could also have a negative impact on the composting microorganisms in the inoculated experiment, resulting in a lower reduction in a phenolic compounds in the augmented experiment compared to the control experiment.

![Graphical determination of rate constant (k) for PC degradation](image)

**Fig. 14:** Graphical determination of rate constant (k) for PC degradation

**Table 5:** Rate constants for PC degradation

| Trial                      | Rate constant \( (h^{-1}) \) |
|----------------------------|--------------------------------|
| Rising phase \((T<45^\circ C)\) | 0.0027                           |
| Thermophilic phase \((45^\circ C<T<55^\circ C)\) | 0.0047                           |
| Inoculated Experiment       | 0.0033                           |
| Control                    | 0.0029                           |

The degradation of organic substrate can be described with the following first order model (Haug, 1993):

\[
C_t = C_0 e^{-Kt}
\]

Where:

- \( C_t \) = Concentration of the organic substrate at time \( t \) (mg/kg)
- \( C_0 \) = Initial concentration of the organic substrate (mg/kg)
- \( K \) = Rate constant

The value \( \ln(C_t/C_0) \) has a linear relationship with the time \( t \) within given temperature range. The linear relationship between \( \ln(C_t/C_0) \) and time for phenolic compounds was determined graphically for the mesophilic and thermophilic temperature zones as shown in Fig. 14. The rate constant \( (k) \) was determined from the slope of the lines and presented in Table 5. The mesophilic rate constant was 0.0027 and 0.0033 h^{-1} for the control and the inoculated experimental trials, respectively. During the thermophilic phase, the rate constant of the inoculated experimental trial (0.0029 h^{-1}) was lower compared with the control (0.0047 h^{-1}). The rate constant of PC degradation was correlated well with reductions in volatile solids, total carbon and TKN.

**Maturity and stability tests:** The maturity and stability of the bioremediation product were evaluated by analyzing the \( CO_2 \) evolution rate of the final product.
and comparing the Germination Index (GI) of the final product with that of the feedstock. CO₂ evolution is a good indicator to determine the level of microbial activity and the stability of compost while GI provides information about the decomposition of phytotoxic organic substances which indicates the maturity of compost. The lower the CO₂ evolution, the more stable the product is and the higher the GI, the more mature the product is (Iannotti et al., 1994; Wu et al., 2000; Boulter-Bitzer et al., 2006). There was significant difference between the control and the inoculated experimental trial in the GI and CO₂ evolution rate. The inoculated experimental trial had a higher CO₂ evolution rate and a lower GI. This indicates that the product of this experiment was less stable than the control. This result correlates well with the reductions in the volatile solids, total carbon and total kjeldahl nitrogen.

Cooperband et al. (2003) conducted windrow composting of sawdust for 1 year and observed a CO₂ evolution of approximately 2 mg CO₂-C/g final stable compost carbon-day. The CO₂ evolution in the current study was higher but the period of bioremediation only lasted 15 days which is much shorter than the processes reported by other researchers. The final product for the control was within the threshold of 8 mg CO₂-C/g carbon-day and is considered stable according to Korner et al. (2003) and Gomez et al. (2006). Rekha et al. (2006) used composting technology for bioremediation of contaminated lake sediments for 14 weeks and reported GIs of 49-95%. The GI of 20-30% obtained in this study after only a short period of time (15 days) is reasonable. Longer bioremediation may be required to achieve more mature products as the phytotoxicity still existed in the final product in both trials. The product of the control was stable but not mature whereas the product of the inoculated experimental trial was neither stable nor mature. The pH of both products was 5.5-6.8 which was within the range of 5-7 for stable compost.

CONCLUSION

The temperatures of bioremediation process reached thermophilic phase (>45°C) and the mesophilic and thermophilic lag phases were clearly identified. The moisture content decreased significantly indicating that the water produced by microbial respiration did not compensate for the water lost as vapor with the exhaust gases. The final moisture content remained in the range of 43-48% which is within the optimum range for composting of 40-60%. Initial increases in the pH due to the breakdown of organic nitrogen to ammonium and final drops in the pH due to the formation of organic acids from decomposition of fats and grease and the loss of ammonium with the exhaust gases in the latter stage were observed. Different degradation rates were observed in the mesophilic and thermophilic stages of composting. The control experiment achieved higher reductions of volatile solids, total carbon and TKN indicating a higher level of activity of microorganisms during the composting process compared with the inoculated experimental trial. As a result, higher degradation of phenolic compounds, cellulose and lignin were observed in the control experiment. The stability and maturity of the product of the control experiment were also better than those of the product from the inoculated experimental trial. The inoculation of cellulolytic and thermophilic fungus Thermosascus aurantiacus did not accelerate the bioremediation process in degrading phenolic compounds and may have inhibited the metabolic activity of composting organisms.

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