Microbial Biomass and Enzymatic Activity of Major Cropping Systems in Soils of Inceptisols and Vertisols at Northern Telangana

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Authors’ contributions

This work was carried out in collaboration among all authors. All authors read and approved the final manuscript.

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ABSTRACT

Under cropping systems, microbial biomass plays a major role in nutrient and energy flow of soil. Similarly, urease and dehydrogenase activities are essential for nitrogen cycle and determining biological index of soils, respectively. However, their information is minimal at major cropping systems of this region. Therefore, surface soil (00-15 cm) samples were collected after 8 years from rice-rice, rice-maize, cotton and turmeric-sesame cropping systems at soils of Inceptisols and Vertisols of Northern Telangana zone during kharif 2019. A five replicated soil samples were collected, assesseed and statistically analyzed with factorial randomized block design. The results revealed that the forms of microbial biomass carbon (14%) and nitrogen (22%), urease (29%) and dehydrogenase (20%) activity were found to be higher in cropping systems under Vertisols compared to Inceptisols. Among the cropping systems, rice-rice showed significantly higher biological properties than others. The interactions are significant for urease activity. Urease and dehydrogenase activity is positively correlated with soil available nitrogen and organic carbon content of soils, respectively of cropping systems.
Keywords: MBC; MBN; urease, dehydrogenase; cropping systems; rice-rice; rice-maize; turmeric-sesamum; cotton fallow.

1. INTRODUCTION

The predominant cropping systems grown in Northern Telangana Zone are rice-rice, rice-maize, cotton-fallow and turmeric-sesame. Rice-based cropping systems are the major contributing food production systems and largely cultivated, contributing 84% of total production of the World. Soils of rice–rice system are heavy textured, slow infiltration rate, high water-holding capacity, rich in soil organic matter (SOM) and easy to puddle. In agro-ecosystem, different crops are grown in sequence and they contribute different amounts of crop residues and root exudates [1]. They may ultimately help to build up different amount of organic carbon in soils. Nature of such organic C also varies because of variation in the nature of root deposition, quality of crop residues and simultaneously there is a build-up microorganisms. Mostly the microbial biomass nitrogen, microbial biomass carbon, urease and dehydrogenase population are found to be higher at root residues when compared to other microbes. 

Microbial population is always a sign of healthy soil and they helps in soil quality maintenance. This study helps to know which cropping systems helps in soil health and soil quality maintenance.

Microorganisms play a crucial part in soil nutrient cycling, maintenance of soil structure, degradation of agrochemicals and pollutants, and plant pest control [2] hence it has often been indicated as an important component of soil fertility [3]. Enzymatic activities in the soil highly affect nutrient cycling and organic matter decomposition [4]. Moreover, ureases are in charge of releasing inorganic N in the N-cycle [5]. A case study indicated that excessive cultivation decreased both microbial biomass and its activities [6].

Soil enzyme activities related to SMBC and soil organic carbon (SOC) are often used for comparison of different land use with varying SOM content [7, 8]. Measurement of soil enzyme activities in key nutrient cycling (C, N, and P) and oxidation–reduction processes have been used widely as a potential indicator for determining the effect of land use conversions and management practices on soil health [9, 10, 11]. Therefore, an investigation of soil microbial characters and enzyme activities is critical in studying the land conversions and focusing the soil management.

2. MATERIALS AND METHODS

An experiment was conducted during kharif, 2019 at Agricultural College, Polasa, Jagtial, Professor Jayashankar Telangana State Agricultural University. The experiment was laid out in randomized block design with factorial concept (FRBD) with five replications. Soil samples were collected from four cropping systems viz., rice-rice (CS1), rice-maize (CS2), cotton-fallow (CS3) and turmeric-sesame (CS4) under two soil types viz., inceptisols (S1) and vertisols (S2) from surface soil (0-15cm). Selection of sites was based on continuous cultivation of the same cropping system (at least for 8 year), in Northern Telangana Zone of Telangana State.

2.1 Microbial Biomass Carbon

Field-moist soil samples (25.0 g) were exposed to CHCl3 vapour for 24 h and extracted with 0.5 M K2SO4. Under same conditions the second set of non-fumigated samples were extracted. The difference between C obtained from the fumigated and from the non-fumigated ones was taken to represent the microbial C-flush and converted to MBC using the relationship: MBC = 1/0.41 × C-flush (Voroney and Paul, 1984). All results are expressed on an oven-dry soil basis (105°C, 24 h) and are the mean of three replicate analyses.

2.2 Microbial Biomass Nitrogen

Microbial biomass nitrogen (MBN) was also estimated using the same principle of microbial biomass carbon [12]. The K2SO4 extractant of both fumigated and unfumigated soil was digested for 3 hr. by adding H2SO4 and digestion mixture. After cooling, distillation was carried out to find the nitrogen content. The difference between fumigated and unfumigated extracted nitrogen of soil divided by a calibration factor ($K_{EC}$) 0.38 gives the measure of microbial biomass nitrogen in soil and expressed as micro gram of microbial biomass-N per gram of dry soil.

2.3 Urease

This method is based on determination of NH3 released other incubation of soil with urea solution for 2 hours at 30°C [13]. Five gram of soil was taken in duplicate in 50 ml volumetric
flask. 0.2 ml toluene and 9 ml of THAM buffer (pH- 9; 0.05M) were added to it. The flasks were swirled for few second to mix the content. Then 1 ml of 0.2 M urea solution was added and swirled again for a few second. The flasks were then stoppered and placed in an incubator at 30°C for 2 hours. After 2 hours the stoppers were removed and approximately 35 ml of KCl – Ag₂SO₄ solution was added and the flasks were swirled for a few seconds and allowed to stand until the contents have cooled to room temperature. The volume of the flasks was made upto mark (50 ml) by addition of KCl – Ag₂SO₄ solution. In order to mix the contents the flasks were inverted several times with the help of a stopper in the flake. To perform control, the above procedure was followed, but 1 ml 0.2M urea solution was added after the addition of 35 ml KCl-AgSO₄ solution.

A 40 ml aliquot of the suspension was pipetted out into 100 ml distillation flask and 0.2g MgO was added to it for the determination of NH₄-N in the resulting soil suspension. The content of the flask was then distilled for 15 minutes and the distillate was collected in a 50 ml conical flask containing 5 ml of 2 per cent boric acid indicator solution. The distillate was then titrated with 0.005(N) H₂SO₄. The urease activity is expressed as micro gram NH₄ – N per gram dry soil per hour at 30°C (µg NH₄⁺ released g soil⁻¹ h⁻¹).

2.4 Dehydrogenase

Five grams soil was weigh into glass tubes and mixed with 5 ml TTC solution. The tubes were sealed with rubber stopper and inoculated for 24 hours at 30°C. The control contain only 5 ml tris buffer (without TTC). After incubation 40 ml acetone was added to each tube and tubes were shaken thoroughly and further incubated at room temperature for 2 hour in dark (shaking the tubes at intervals). The suspension was then filtered and optical density of clear supernatant was measured against the blank at 546 nm (red colour). The activity of dehydrogenase is expressed in µg TPF formed per gram of dry soil per day (µg TPF produced g⁻¹ soil h⁻¹).

2.5 Statistical Analysis

The data were analysed using analysis of variance (ANOVA) – two way classification. Two factor factorial ANOVA was used to determine the existence of interaction effect between soil orders and cropping systems. Simple correlation coefficient was also developed to evaluate relationships between the response variables using the same statistical package. The 5% probability level was regarded as statistically significant [14].

3. RESULTS AND DISCUSSION

The higher content of soil organic carbon, the more active the soil microorganisms. Microorganisms accelerate the degradation of organic matter, which is reflected in soil respiration and release of carbon dioxide from the rhizosphere (Zhang et al., 2010). The results on the effect of cropping systems and soil type on microbial biomass and enzyme activity are in the Table.1.

3.1 Microbial Biomass Carbon

Microbial biomass carbon (MBC) content in surface soils was significantly higher in vertisols (132 µg C g⁻¹ soil) than inceptisols (116 µg C g⁻¹ soil) the results were in agreement with Prasad et al. 2013. The high clay and organic matter contents in vertisols might have contributed for higher MBC values compared to inceptisols. Different cropping systems were found to have significant effect on MBC. MBC was in the range of 89 to 152 µg C g⁻¹ soil MBC values was found highest in CS₁ (141 µg g⁻¹) followed by CS₂ (134 µg g⁻¹), CS₃ (123 µg g⁻¹) and CS₄ (99 µg g⁻¹). Higher MBC values are commonly found in cropping systems that include high residue-producing crops [15] crops with intensive root growth and root density ([16,17]) Thus CS₁ has found to maintain significantly higher MBC values than other cropping systems this may be due to the amount of crop residues, the proportion of easily decomposable organic compounds returned to the soil, root density and microclimate in rice-rice cropping systems [18,19]. Interactional effect of soil types and cropping system were found to be non significant.

3.2 Microbial Biomass Nitrogen

Microbial biomass nitrogen (MBN) content in surface soils was significantly higher in vertisols (11.21 µg NH₄⁺ g⁻¹ soil) than inceptisols (9.16 µg NH₄⁺ g⁻¹ soil). The result is also in accordance with Prasad et al. [20]. The high clay and organic matter contents in vertisols might have contributed for higher MBN values compared to inceptisols. Different cropping systems were found to have significant effect on MBN. The values of MBN ranged from 8.85 to 11.64 µg
NH$_4^+$ g$^{-1}$ soil. MBN values were found highest in CS$_1$ (11.64 µg NH$_4^+$ g$^{-1}$ soil) followed by CS$_2$ (10.63 µg g$^{-1}$), CS$_4$ (9.6 µg g$^{-1}$) and CS$_3$ (8.86 µg g$^{-1}$). Thus MBN count was higher in CS$_1$ as this system has more substrate production when compared to other systems above. The microbial biomass C and N pools will increase in one cropping system relative to another only if microbes have access to sufficient substrates and non-growth requirements have been satisfied. Therefore, greater microbial biomass in diversified cropping systems may be a consequence of increased substrate availability, where greater retention and recycling of C and N enhance available substrate to support microbial growth and biosynthesis [21]. Interactional effect of soil types and cropping system were found to be non significant.

3.3 Urease

Soil types had significant impact on the activity of urease. Greater urease activity was recorded in vertisols (3.32 µg NH$_4$-N g$^{-1}$ hr$^{-1}$) than inceptisols (2.58) as biological activities are more in vertisols than inceptisols Prasad et al. [20]. In cropping systems, rice-rice (CS$_1$) has maintained higher amount of urease activity followed by CS$_2$ > CS$_4$ > CS$_3$ with the activity of 3.84, 3.62, 2.41 and 1.92 µg NH$_4$-N g$^{-1}$ hr$^{-1}$ respectively. Biogenic elements were more in rice-rice cropping system compared to other systems [22] which helps in increase of microbial properties, such as urease [22,23,24] Urease participate in ammonification, during which ammonia is released from urea, amino acids, and purine bases. Soil fertility and productivity depend on soil organic matter, which is a reserve of nutrients and is very important in nutrient cycling [25] as well as improves soil physical, chemical, and biological properties [26]. Processes associated with organic matter transformations in soil occur with the participation of soil microorganisms and their enzymes [27].

Interactional effect of cropping system and soil types were found to have a profound influence on urease activity. CS$_1$ cropping system maintained higher urease activity followed by CS$_2$, CS$_4$ and CS$_3$ in inceptisol. Similar trend also recorded with vertisols. Soil available nitrogen content in soil had positive influence on urease activity. With higher the content of nitrogen in soil more will be the urease activity, such results were also found with [22,23,24]. Relationship between urease with available nitrogen content was positively correlated ($Y=26.80X+128.2; R^2=0.499$), is shown in the Fig. 1. Nitrogen stimulates soil microorganisms which produce more soil enzymes when biogenic elements become more available [22]. As urease is the enzyme that catalyzes hydrolysis of urea to CO$_2$ and NH$_3$, which is a vital process in the regulation of N supply to plants after urea fertilization [28].

![Fig. 1. Correlation studies a) Soil available nitrogen Vs Urease b) Organic carbon Vs Dehydrogenase](image-url)
1. Soil organic carbon has a positive relationship with dehydrogenase activity. Higher biological activities in soil have been observed under rice ecology. Under cropping systems, urease activity was highest in CS1, followed by CS2, CS3, and CS4, respectively. The results were in line with the findings of Prasad et al. [30]. The greater amount of organic carbon, nutrients, and stimulated microbial activity in vertisols compared to inceptisols. Different cropping systems were found to have significant effects on dehydrogenase activity. Irrespective of soil order, dehydrogenase activity was found to have significant effects on dehydrogenase activity. This activity was highest in CS1, as biological activity was found highest in rice ecology. Under CS1, the very labile pool of SOC was higher, which might have been used as feed for microorganisms [31,32] and enhances soil enzymatic activity. The relationship between organic carbon and dehydrogenase activity was positively correlated.

### Table 1. Influence of soil types and cropping systems on soil microbial biomass carbon (µg C g⁻¹ soil), microbial biomass nitrogen (µg NH₄-N g⁻¹ soil), urease (µg NH₄-N g⁻¹ hr⁻¹) and dehydrogenase (µg TPF g⁻¹ hr⁻¹)

| Soil order | MBC  | MBN  | Urease | Dehydrogenase |
|------------|------|------|--------|---------------|
| S₁         | 116.15 | 9.16 | 2.58   | 2.25          |
| S₂         | 132.41 | 11.21| 3.32   | 2.71          |
| S.Em       | 4.73   | 0.35 | 0.09   | 0.09          |
| CD@5%      | 13.71 | 1.00 | 0.27   | 0.27          |

**Cropping System**

| CS₁       | 141.14 | 11.64 | 3.84   | 3.52          |
| CS₂       | 123.24 | 10.63 | 3.62   | 2.65          |
| CS₃       | 98.73  | 8.86  | 1.92   | 1.48          |
| CS₄       | 134.02 | 9.60  | 2.41   | 2.27          |
| S.Em      | 9.00   | 0.49  | 0.13   | 0.13          |
| CD@5%     | 26.07  | 1.42  | 0.39   | 0.38          |

**Interactions**

| S₁S₂      | 130.74 | 10.78 | 3.65   | 3.32          |
| S₁S₃      | 119.15 | 9.82  | 3.44   | 2.25          |
| S₁S₄      | 88.58  | 7.91  | 1.52   | 1.32          |
| S₂S₃      | 126.12 | 8.11  | 1.71   | 2.12          |
| S₂S₄      | 151.55 | 12.49 | 4.04   | 3.72          |
| S₃S₄      | 127.31 | 11.45 | 3.80   | 3.05          |
| S₂S₄      | 108.85 | 9.80  | 2.32   | 1.64          |
| S₃S₄      | 141.94 | 11.08 | 3.12   | 2.43          |
| S.Em      | 9.47   | 0.69  | 0.19   | 0.18          |
| CD@5%     | NS     | NS    | 0.55   | NS            |

**Soil order abbreviations:** S₁: Inceptisols, S₂: Vertisols, S₃: Rice-Rice, S₄: Rice-Maize, S₅: Cotton-Fallow, S₆: Turmeric-Sesame, SE: Standard error of mean, CD: Critical difference, CV: Critical Variance

### 3.4 Dehydrogenase Activity

Activity of dehydrogenase reflects oxidative activity of soil microflora and is a good indicator of microbial [29-30]. Activity of dehydrogenase enzyme was significantly higher in vertisols (2.71 µg TPF g⁻¹ hr⁻¹) over inceptisols (2.25 µg TPF g⁻¹ hr⁻¹). The results were in the same line with the findings of Prasad et al. [30]. The greater amount of organic carbon, nutrients, and stimulated microbial activity in vertisols might have contributed to the higher biological activities in soil. Different cropping systems were found to have significant effects on dehydrogenase activity. Irrespective of soil order, dehydrogenase activity was found to have significant effects on dehydrogenase activity. This activity was highest in CS1, as biological activity was found highest in rice ecology. Under CS1, the very labile pool of SOC was higher which might have been used as feed for microorganisms [31,32] and enhances soil enzymatic activity. The relationship between organic carbon and dehydrogenase activity was positively correlated, as shown in the Fig. 1. Similar correlation was also observed by Bergstrom et al. [33] and Roldan et al., 2005. Soil organic carbon has been considered as an indicator of soil quality, because of its character of nutrient sink and source that can enhance soil physical and chemical properties, and also promote biological activity [34] and highest carbon and biological activity were found in rice ecology when compared to other cropping systems. Interational effect of soil type and cropping system were found to be non significant.

### 4. CONCLUSION

Biological properties viz., microbial biomass carbon, microbial biomass nitrogen, dehydrogenase and urease enzyme activities of soils showed higher values under vertisols over inceptisols. Under cropping systems compared rice-rice cropping systems showed significantly higher biological activities in soil. Lowest activities were recorded in cotton-fallow cropping system. Urease was positively correlated with available nitrogen and dehydrogenase was positively correlated with SOC content. Microbial
activities in soil were found higher under vertisols and in rice-rice cropping system.

COMPETING INTERESTS

Authors have declared that no competing interests exist.

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