Interactive effects of non-fodder litter and fungal species on soil enzymes: A microcosm temporal assessment from Indian arid zone

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ABSTRACT
The interactive effects of three non-fodder Indian arid plant species, Tephrosia purpurea, Aerva persica, and Calotropis procera, and four Aspergillus fungal species on soil enzymes (acid and alkaline phosphatase, -glucosidase, dehydrogenase, urease, and amidase activities) were temporally assessed (15 and 30 days withdrawals). The results were statistically analysed using ANOVA, Principal Component Analysis (PCA), and Canonical Correlation Analysis (CCoA). Aside from these, a biochemical soil quality index was created by assigning a weighted score to each enzyme and analysing it using PCA. This study found that various litter-fungal species complexes acted differently and that their effects changed over time, specifically for acid phosphatase, alkaline phosphatase, beta-glucosidase, and amidase. Dehydrogenase and urease activities increased with predictors over time. With temporal backwash, all four fungal species with C. procera inhibit acid phosphatase, alkaline phosphatase, and beta-glucosidase activities (i.e., more at 15 days and lesser after 30 days). Our current findings suggest that (a) urease activities were modulated by A. persica in cooperation with fungi like A. terreus, A. niger, and A. flavus at specific enzyme levels; (b) In assistance with fungi such as A. fumigatus, A. niger, and A. persica, amidase concentration was successfully managed through litter of the legume plant species T. purpurea. (c) When C. procera and A. fumigatus, A. niger, and A. flavus worked together, they were most effective at supporting beta-glucosidase and dehydrogenase (d) Alkaline phosphatase and acid phosphatase was more responsive to T. purpurea-A. terreus complexes than were T. purpurea-A. flavus and C. procera-A. terreus complexes.

Introduction
Soil health is the net result of ongoing conservation and degradation processes; which is heavily dependent on the biological elements of the soil ecosystem and affects plant health, environmental health, food safety, and quality (Laishram et al., 2012). Numerous indicators can be used to gauge this, but soil enzymes are among the most significant because they are essential for preserving soil fertility, soil health, and soil ecology (Mathur and Sundaramoorthy, 2019). They react quickly to changes in environmental circumstances and soil management practices. Because of this, they are used as sensors, and they have been researched all over the world as a measure of soil fertility (Utobo & Tewari, 2015), a measure of microbial biomass (Ren et al., 2018), as indicators of vegetation effects and capability to conduct bio-geochemical cycling, total microbial activity (Luo et al., 2018), as a predictor of bio-remediation (Basak et al., 2016), and to understand the consequence of

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degradation (Pajares et al., 2011; Samuel, 2017). Their responses have also been assessed in terms of litter type (Xiang et al., 2018; Jagadish et al., 2001), soil fungi (Oseni, 2011), soil management practises (Maharjan et al., 2017), and temporal variations (Ren et al., 2018; Veeraragavan et al., 2018). Some of the above attributes have been addressed in Indian arid and semi-arid zones by Tarafdar et al. (2002) and Gaur et al. (2012). Few studies have been done on the interactions between different types of plant litter and different fungi species and extracellular soil enzymes in hot dry region of India. In light of this, our study closes a knowledge gap regarding the impact of fungus species and litter type on extracellular soil enzymes. We hypothesized in this work that the effects of particular plant litter and fungus species on soil enzymes would vary from their combined effects and that these variances would change over time. The objectives of this study were (a) to evaluate the temporal effects of litter from three wild arid plant species on acid phosphatase, alkaline phosphatase, dehydrogenase, urease, and amidase, i.e. after 15 and 30 days withdrawal periods. (b) to visualise the studied enzymes in a cumulative approach that takes into account the interactive effects of the litter-fungi-time complex, and (c) to develop a biochemical soil quality index.

Material and Methods
Species and experimental setup
As litter source experimental, three non-fodder arid wild plant species were used: Tephrosia purpurea (L.) Pers., Aerva persica (Burm.f.) Shult, and Calotropis procera (Aiton) W.T.Aiton. These species can be found in abundance in wastelands, fallow lands, crop-fences, and community grazing lands (Mathur and Pandey, 2016). As litter, total aboveground biomass of T. purpurea and A. persica, as well as a mixture of stem (1 cm width) and leaves of C. procera, were used. These species were collected in and around natural agroforestry field located at six different provinces of arid eco-regions, namely Shergarh (Site 1; 26°19' 32.66" N and 72°17' 13.16" E), Balesar (Site 2; 26°24.0' 69" N and 72°28' 59.83" E), Osiyan (Site 3; 26°43' 53.46" N and 72°54' 30.79" E), Chawa (Site 4; 26°22.06' 62" N and 72°09.05' 58" E), Baitu (Site 5; 25°54' 13.72" N and 71°46' 15.69" E) and Kalyanpur (Site 6; 26°0.1’ 19.67" N and 72°34' 34.21" E). After being collected, each plant material was air dried in comparable climatic conditions. After achieving a homogeneous moisture level, they were completely blended to create homogeneous material. For a microcosm experiment, four different fungi—Aspergillus flavus, Aspergillus fumigates, Aspergillus niger, and Aspergillus terreus—were chosen and inoculated with plant litter from non-fodder plants. The plant pathology lab of the Department of Botany at Jai Narain Vyas University in Jodhpur, India, provided these fungal cultures. The same department's experimental field of ecology laboratory provided the soil, which was then pulverized and sieved through a 2 mm mesh size in preparation for the microcosm experiment. Then, over the course of three days, this natural soil was tyndalized by autoclaving at 121°C for an hour and overnight oven drying at 80°C. (Eivazi and Tabatabai, 1988).

100g tyndalized soil was placed in 250ml cotton plugged conical flasks, followed by litter at a fertility level of 2000 kg N h⁻¹ and the fungal culture (on the basis of colony forming unit 3 X 10⁷ per treatment). After 15 and 30 days of laboratory incubation, this microcosm experiment setup was recovered, and six different extracellular enzymes were quantified. Acid and alkaline phosphatase (Eivazi and Tabatabai, 1977), glucosidase (Eivazi and Tabatabai 1988), dehydrogenase (Tabatabai, 1982), urease (Douglas and Bremner, 1977a and b), and amidase (Frankenberger and Tabatabai, 1980) were all quantified using standard methodologies. After incubation and enzyme reaction termination, each enzyme activity was kept under control by adding substrate to blank samples (Mathur, 2005). All the experiments were conducted in triplicates.

Statistical analysis
To analyze the effects of three major sources of variation (litter types, fungal species, and withdrawal time) and their interactions on the concentration of the examined enzymatic activities, we utilized Statsoft's (2011 Version 10) three-way ANOVA (Strip-Split design) tool. The intersection plot is divided into subplots in the strip-split-plot design to accommodate a third element, extending the capabilities of the strip-plot design. This design stands out due to the use of four levels of precision for measuring the effects of various factors, with the highest level corresponding to the sub-plot.
factor and its interactions with other factors, as well as more than three plot sizes (such as horizontal, vertical strip, intersection plot, and subplot). The percentage temporal deviation in various soil enzymes was calculated in relation to litter type and fungal species by using the following formula.

\[
\% \text{ Temporal Deviation} = \left( \frac{W_2 - W_1}{W_1} \right) \times 100 \quad (1)
\]

Where \(W_1\) and \(W_2\) are first and second withdrawal periods. In this study, principal component analysis (PCA) was used for two distinct purposes: (a) to visualize the litter-fungal complex with reference to their proximity or distance with withdrawal periods for each studied enzyme; and (b) to develop a soil quality index with weighted scores assigned to each enzyme. The distance of each litter-fungal complex from the centroid of the PCA bi-blot developed in earlier steps was calculated. A cumulative data set that includes the litter-fungi complex for each enzyme under consideration was produced using these centroid distances. This cumulative data set was used to create the PCA bi-plot, which graphically depicts the overall scenario for various enzymes that is naturally adjusted by litter-fungal withdrawal as well as their interactions. Such PCA strategies have been supported by many researchers (Laliberte and Legendre, 2010, Mathur and Sundaramoorthy, 2018). The connections between soil enzymes detected during the first and second withdrawal periods, as well as between these two periods, were established using Canonical Correlation Analysis (CCoA). Correlation matrices were converted into a network-like structure for ease of graphical interface. PCA was used to quantify correlations among enzymes with weighted scores using the PAST (Hammar et al. 2001) and XLSTAT (2017) software.

Weights in the soil quality index were determined by dividing the percent of variation in the data set explained by the principal component analysis that contributed the indicated variable by the total percentage of variance explained by all the PCs with more than one eigenvector (Mathur and Sundaramoorthy, 2018).

\[
SQI = \sum_{i=1}^{n} W_i x S_i \quad (2)
\]

\(W_i\) = weighting factor of soil enzyme \(i\) and \(S_i\) = value of soil enzyme \(i\).

Results and Discussion

**Acid phosphatase (µg p-Nitrophenol released h\(^{-1}\) g\(^{-1}\) of soil)**

This enzyme had a higher concentration (0.97) with *C. procera + A. terreus* after 15 days, but it had a lower concentration (0.65) after 30 days with a different plant-fungal complex (i.e., *A. persica + A. niger* Table 1). Except for *A. persica + A. niger* and *T. purpurea + A. fumigatus*, we found higher concentrations of this enzyme during 15 days withdrawal compared to 30 days with most treatments. We noticed +7.40 and +5.41 percent temporal deviation with these two complexes. (Table 2).

**Alkaline phosphatase (µg p-Nitrophenol released h\(^{-1}\) g\(^{-1}\) of soil)**

Higher concentrations were found in *C. procera + A. fumigatus* (1.87) and *C. procera + A. terreus* (1.38), respectively, at 15 and 30 day intervals (Table 1). While its minimum concentration was recorded at 15 and 30 day intervals with *A. persica + A. niger* (0.58) and *A. persica + A. fumigatus* (0.87), respectively. With the exception of *C. procera* and *T. purpurea + A. terreus*, all four fungal species showed negative temporal deviation (Table 2), indicating that they were recorded more during the first sampling period.

**Beta-glucosidase (µg p-Nitrophenol released h\(^{-1}\) g\(^{-1}\) of soil)**

During the two sampling periods, higher concentrations of this enzyme (1.19 and 1.14) were detected with the *C. procera + A. fumigatus* complex (Table 1). Its minimum concentrations with *T. purpurea + A. flavus* (0.35 at 15 days) and *T. purpurea + A. fumigatus* (0.35 at 15 days) were determined (0.42 at 30 days). Similar to Alkaline Phosphatase, all four fungal species with *C. procera* and *T. purpurea + A. fumigatus* exhibited negative temporal deviations (Table 2).

**Dehydrogenase (µg TPF released h\(^{-1}\) g\(^{-1}\) of soil)**

*Calotropis procera* was found to be more conducive for this enzyme with four studied fungi, with higher concentrations recorded with *C. procera-A. flavus* (12.68) and *C. procera-A. fumigatus* (12.67) during second sampling periods (i.e., 30 day). During the first sampling period (15
Table 1: Effects of soil fungi and litter types on various soil enzymes during two sampling times.

<table>
<thead>
<tr>
<th>Variables</th>
<th>AcP</th>
<th>AlP</th>
<th>BG</th>
<th>De</th>
<th>Ur</th>
<th>Am</th>
</tr>
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<td>1</td>
<td>2</td>
<td>1</td>
<td>2</td>
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<td>Control</td>
<td>0.39</td>
<td>0.23</td>
<td>0.6</td>
<td>0.46</td>
<td>0.16</td>
<td>0.34</td>
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<td>T. purpurea + A. flavus</td>
<td>0.64</td>
<td>0.54</td>
<td>1.11</td>
<td>1.16</td>
<td>0.35</td>
<td>0.64</td>
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<tr>
<td>T. purpurea + A. fumigatus</td>
<td>0.47</td>
<td>0.50</td>
<td>0.61</td>
<td>0.91</td>
<td>0.49</td>
<td>0.42</td>
</tr>
<tr>
<td>T. purpurea + A. niger</td>
<td>0.59</td>
<td>0.44</td>
<td>1.07</td>
<td>1.21</td>
<td>0.42</td>
<td>0.43</td>
</tr>
<tr>
<td>T. purpurea + A. terreus</td>
<td>0.64</td>
<td>0.63</td>
<td>1.25</td>
<td>1.21</td>
<td>0.39</td>
<td>0.56</td>
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<tr>
<td>A. persica + A. flavus</td>
<td>0.67</td>
<td>0.44</td>
<td>0.98</td>
<td>1.03</td>
<td>0.40</td>
<td>0.69</td>
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<tr>
<td>A. persica + A. fumigatus</td>
<td>0.59</td>
<td>0.50</td>
<td>0.62</td>
<td>0.87</td>
<td>0.52</td>
<td>0.68</td>
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<tr>
<td>A. persica + A. niger</td>
<td>0.61</td>
<td>0.65</td>
<td>0.58</td>
<td>1.16</td>
<td>0.52</td>
<td>0.87</td>
</tr>
<tr>
<td>A. persica + A. terreus</td>
<td>0.72</td>
<td>0.60</td>
<td>0.62</td>
<td>1.17</td>
<td>0.56</td>
<td>0.74</td>
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<tr>
<td>C. procera + A. flavus</td>
<td>0.73</td>
<td>0.50</td>
<td>1.50</td>
<td>0.89</td>
<td>0.97</td>
<td>0.69</td>
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<tr>
<td>C. procera + A. fumigatus</td>
<td>0.88</td>
<td>0.47</td>
<td>1.87</td>
<td>1.12</td>
<td>1.19</td>
<td>1.14</td>
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<tr>
<td>C. procera + A. niger</td>
<td>0.91</td>
<td>0.54</td>
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<td>0.90</td>
<td>1.08</td>
<td>0.75</td>
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<td>C. procera + A. terreus</td>
<td>0.97</td>
<td>0.53</td>
<td>1.40</td>
<td>1.38</td>
<td>0.98</td>
<td>0.78</td>
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</tbody>
</table>

AcP = Acid Phosphatase, AlP = Alkaline Phosphatase, BG = Beta-glucosidase, De = Dehydrogenase, Ur = Urease, Am = Amidase. 1 and 2 are 15 and 30th days withdrawal, respectively.

Table 2: Percent temporal deviation in various soil enzymes with Litter types and fungal species.

<table>
<thead>
<tr>
<th>Variables</th>
<th>AcP</th>
<th>AlP</th>
<th>BG</th>
<th>De</th>
<th>Ur</th>
<th>Am</th>
</tr>
</thead>
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<td></td>
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<td>2</td>
<td>1</td>
<td>2</td>
<td>1</td>
<td>2</td>
</tr>
<tr>
<td>T. purpurea + A. flavus</td>
<td>-15.81</td>
<td>5.39</td>
<td>80.16</td>
<td>50.66</td>
<td>48.11</td>
<td>6.34</td>
</tr>
<tr>
<td>T. purpurea + A. fumigatus</td>
<td>5.41</td>
<td>48.77</td>
<td>-14.52</td>
<td>140.64</td>
<td>27.28</td>
<td>-51.16</td>
</tr>
<tr>
<td>T. purpurea + A. niger</td>
<td>-26.40</td>
<td>13.65</td>
<td>2.43</td>
<td>390.87</td>
<td>15.23</td>
<td>-67.68</td>
</tr>
<tr>
<td>T. purpurea + A. terreus</td>
<td>-1.96</td>
<td>-2.95</td>
<td>43.05</td>
<td>580.23</td>
<td>10.49</td>
<td>63.02</td>
</tr>
<tr>
<td>A. persica + A. flavus</td>
<td>-33.30</td>
<td>4.40</td>
<td>75.09</td>
<td>372.13</td>
<td>40.83</td>
<td>10.77</td>
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<tr>
<td>A. persica + A. fumigatus</td>
<td>-15.72</td>
<td>40.42</td>
<td>30.50</td>
<td>390.71</td>
<td>13.23</td>
<td>-65.46</td>
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<tr>
<td>A. persica + A. niger</td>
<td>7.40</td>
<td>100.09</td>
<td>67.42</td>
<td>287.11</td>
<td>75.71</td>
<td>-8.52</td>
</tr>
<tr>
<td>A. persica + A. terreus</td>
<td>-16.57</td>
<td>89.30</td>
<td>31.32</td>
<td>302.87</td>
<td>43.42</td>
<td>11.16</td>
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<tr>
<td>C. procera + A. flavus</td>
<td>-31.22</td>
<td>-40.57</td>
<td>-29.18</td>
<td>228.40</td>
<td>57.09</td>
<td>-53.16</td>
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<tr>
<td>C. procera + A. fumigatus</td>
<td>-46.41</td>
<td>-39.97</td>
<td>-4.75</td>
<td>182.30</td>
<td>40.21</td>
<td>-45.31</td>
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<tr>
<td>C. procera + A. niger</td>
<td>-40.82</td>
<td>-47.05</td>
<td>-30.02</td>
<td>86.68</td>
<td>42.64</td>
<td>-62.56</td>
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<td>C. procera + A. terreus</td>
<td>-45.55</td>
<td>-1.80</td>
<td>-20.83</td>
<td>63.06</td>
<td>46.57</td>
<td>9.48</td>
</tr>
</tbody>
</table>

AcP = Acid Phosphatase, AlP = Alkaline Phosphatase, BG = Beta-glucosidase, De = Dehydrogenase, Ur = Urease, Am = Amidase
days), the concentration of this enzyme was lower with *T. purpurea - A. terreus* (1.54) and with *A. persica-A. flavus, T. purpurea - A. niger*, and *T. purpurea-A. fumigatus* (Table 1). *T. purpuria-A. terreus* had the highest percent positive deviation (580.23) over two sampling periods (Table 2). Such findings imply that the efficacy of this litter-fungi complex is time-dependent.

**Urease (µg urea hydrolyzed released h⁻¹g⁻¹ of soil)**

The effects of litter type and fungal species on this enzyme were more pronounced during the second sampling period (Table 1). The highest concentration (406.45) was obtained with *A. persica-A. niger*, while the complex of *C. procera-A. flavus* after 15 days withdrawal was the least effective (186.87) for this enzyme, and the highest positive deviation (75.82) was obtained with *A. persica-A. niger* (Table 2).

**Amidase (µg NH₄⁺ released h⁻¹g⁻¹ of soil)**

The effects of litter types and soil fungi on the concentration of this enzyme were greatest (0.40) with *T. purpurea - A. terreus*, but the least (0.08) with the same litter type but with *A. niger* (Table 1). Impacts of *C. procera* with different soil fungi were recorded more during 15 days withdrawal compared to 30 days. Both positive and negative temporal deviations were recorded for this enzyme and *T. purpurea - A. flavus, T. purpura-A. terreus, A. persica-A. terreus, C. procera-A. terreus* supports concentration of this enzyme with time advancement. However, effect of *T. purpura-A. niger, T. purpura-A. fumigatus, A. persica-A. flavus, A. persica-A. niger*, and *C. procera with A. flavus, A. fumigatus, A. niger* on enzyme concentration held back with time advancement (Table 2). The results of the analysis of variance (ANOVA) revealed that all of the studied variables, including litter type, fungal species, and sampling time, as well as their interactions, caused significant variations in alkaline phosphatase, dehydrogenase, and urease enzymes. Significant differences in acid phosphatase were brought about by litter and fungal species, as well as sampling time. Their interactions, however, were insignificant. For beta-glucosidase, neither the sampling factor nor its interaction with fungal species were significant. For amidase, litter types and their interactions with sampling time were not significant. The effects of predictors on enzyme activities are clearly visible as we have got many fold increase in concentration of studied enzyme (Table 1) in comparison to control. However, treatments like *T. purpurea-A terreus* (15 days) for dehydrogenase (-10.70%) and *T. purpurea-A. fumigatus* (-25.47%), *T. purpurea – A. niger* (-49.74%), *A. persica-A. fumigatus* (-41.07%) and *C. procera-A. niger* (-16.81%) for amidase (30 days) revealed their negative impacts on these enzymes with compared to control. Bi-plots of all the soil enzymes showed that first two axes (F1 and F2) together accounted 100 per cent variabilities (69.41 and 30.58, 64.06 and 35.79, 86.11 and 13.88, 75.86 and 24.12, 81.96 and 18.03, 51.60 and 48.39 for acid phosphatase, alkaline phosphatase, beta-glucosidase, dehydrogenase, urease and amidase, respectively). Such results indicated the appropriate use of this tool as the cumulative percentage of variance for each enzyme approached >80 per cent. Certain specific trends were visualized that were pertains to litter-fungal complex and with withdrawal time (a) *C. procera with A. fumigatus, A. niger* and with *A. flavus*, was more conducive for acid phosphatase, alkaline phosphatase, beta-glucosidase and for amidase with 15 days period, (b) *A. niger and A. terreus* with litter like *T. purpurea, A. persica and A. fumigatus, A. terreus* with *A. persica* and with *C. procera* were more effective for dehydrogenase enzyme with 30 days, while *C. procera with A. niger* and with *A. terreus* having an intermediate temporal positive impact for this enzyme (c) *T. purpurea with A. fumigatus, A. niger* and with *A. terreus* showed a temporal progressive pace for urease and a decreased temporal pace were showed by *T. purpurea with A. fumigatus, niger* and with *A. persica - A. fumigatus* for amidase enzyme. Further with these PCA analysis, the factorial score (distance from centroid) for each litter-fungi complexes along with control and for enzymes were calculated and are presented in (Table 3). With this data set a bi-plot was further constructed (Figure 1) representing all the studied enzymes and litter-fungi complex along with accommodated withdrawal period.
Figure 1: PCA Bi-plot with Factorial Score of Different Enzymes with different treatments.

For this the per cent variability explained by first four axes were, 50.54, 24.60, 13.15 and 7.7 with their eigen value, 6.5, 3.19, 1.7 and 1.0, respectively. This approach suggested a more cumulative relationship among enzyme-litter-fungi complex and revealed that (a) A. persica with fungi like A. terreus, A. niger, A. flavus are crucial one for the urease (b) T. purpurea with A. fumigatus and A. niger and A. persica- A. fumigatus were more effective for amidase, (c) C. procera with A. fumigatus, A. niger and with A. flavus were more supportive for beta-glucosidase and dehydrogenase, (d) C. procera- A. terreus and T. purpurea- A. flavus found to be effective with alkaline phosphatase and (e) Acid phosphatase having more proximity with T. purpurea -A. terreus . With this tool our biochemical soil quality equation was developed which can be equate as

\[ SQI = \sum_{i=1}^{n} -0.68 \text{ Acid Phosphatase} - 0.46 \text{ Alkaline Phosphatase} + 0.32 \text{ Beta-glucosidase} + 0.18 \text{ Dehydrogenase} + 1.89 \text{ Amidase} - 1.25 \text{ Urease} \quad (3) \]

Here, the numbers represent the distance of each enzyme from the centroid and sign represent their relative position on PCA bi-plot (Figure 1). Use of PCA in the development of a soil quality index are provided by many workers also (Laishram et al., 2012; Cherubin et al., 2016; Guo et al., 2018). Significant correlations among enzymes (studied during first and second withdrawal and between enzymes quantified during these two sampling periods and among enzymes) are depicted in Figure 2. CCoA revealed that urease during the first withdrawal period not related to any other studied enzymes, however, it showed correlation with almost all enzymes during second sampling period except amidase.

Significant intra-relationship was also recorded between BG\textsubscript{1} and BG\textsubscript{2} (r\textsuperscript{2} = 0.72). Interestingly, amidase during the first sampling period was significantly correlated with all studied enzymes except urease, however, during the second sampling period, it was linked with alkaline phosphatase (AIP\textsubscript{2}) only (r\textsuperscript{2} = 0.54). Correlations among weighted enzymes (Figure 3) suggested that both acid phosphatase as well as alkaline phosphatase related to all other enzymes except amidase. Amidase (r\textsuperscript{2} = 0.55) and urease (r\textsuperscript{2} = 0.56) were correlated with beta-glucosidase and with alkaline phosphatase, respectively. The strongest relationships were presented between BG\textsubscript{1}-Am\textsubscript{1} (r\textsuperscript{2} = 0.92) and between BG\textsubscript{w} – De\textsubscript{w} (r\textsuperscript{2} = 0.87).
Interactive effects of non-fodder litter and fungal species

Figure 2: Correlation matrix for different soil enzymes (temporal, and combined)
Significant relationships are indicated by bold lines). Acid P = Acid Phosphatase; Al P = Alkaline Phosphatase; Beta Glu = Beta-glucosidase; Dehydro = Dehydrogenase. 1, 2, represents the 15 and 30th days withdrawals.

Figure 3: Correlation matrix for different soil enzymes (weighted)
Extracellular enzyme activities (EEAs) are indicators of both soil microbial activity and nutrient availability for plants. However, it is unclear how EEAs change in response to litter and micro-organism inputs particularly in the Indian hot desert. The effects of the desertic environment on soil enzymes have been explored with different predictors like plantation regime of Caragana microphylla (Cao et al. 2008), soil properties (Stursova and Sinsabaugh, 2008; Buscardo et al. 2021), biological soil crust (Liu et al. 2014), effects of rainfall treatments (Laura et al. 2015), long term restoration of desertified land (Zhang et al. 2015a), effects of grazing and cultivation (He et al. 2017). The temporal dynamics of enzyme activities should reflect the availability of substrates (Zhang et al. 2015b). Our findings address this discrepancy by indicating that soil enzyme activities vary with litter
and fungal species types, as well as their combined actions. In this study, the temporal increase in enzyme activities such as dehydrogenase and urease could be attributed to litter's dual role in (a) providing suitable substrate for microbial activities and (b) contributing to the formation and stability of soil aggregates. These findings are consistent with those of Acosta-Martinez et al. (2003) and Fang et al. (2013). Using saline plant species like Asteriscus maritimus, Arthrocnemum macrostachyum, Frankenia corymbosa, Halimione portulacoides, Limonium cossonianum, Limonium caesium, Lygeum spartum, and Suaeda vera, Caravaca et al. (2005) reported significant variations in dehydrogenase, urease, phosphatase, and beta-glucosidase. They also came to the conclusion that this alteration may be attributable to various microbial communities connected to the rhizosphere soil or to various quantities of microbial biomass carbon. Effects of monoculture and polyculture practices on rhizosphere enzyme activities were studied by Yang et al. (2007), Fang et al. (2013) and Bogat and Walczak (2022). They concluded that cause-effect relationships between type of cultural practices and modification in rhizosphere enzyme concentration cannot be generalized and they are region and species specific. The seasonal dynamics of soil enzyme activities in response to leaf litter of Cassia siamea, Shorea robusta, Eucalyptus citriodora, Acacia auriculiformis, Anacardium occidentale, Dalbergia sissoo was reported by Venu et al. (2016). Two fundamental lines of thought can be contributed in light of the recent findings: (a) different litter-fungus species complexes functioned differently at the level of the specific enzyme. Particularly for beta-glucosidase, amidase, acid phosphatase, and alkaline phosphatase, their effects evolved with time. The aforementioned factors led to an increase in dehydrogenase and urease activity over time. All four fungi containing C. procera suppress the activities of acid phosphatase, alkaline phosphatase, and beta-glucosidase with temporal backwash (i.e., more at 15 days and lesser after 30 days). Such patterns were seen for amidase except with C. procera-A. terreus. As a result of this study, a litter-fungus species specific complex can be recommended for the studied enzyme, (b) all enzyme predictors: We created a bio-chemical soil quality index using PCA and CCoA, in which each enzyme was weighted with numerical values based on their relationships with litter type (plant species), fungi species types, and withdrawal period (temporal factor). As a result, faith in studied enzymes with litter-fungi-time complex is provided. At specific enzyme levels, our current findings suggest that (a) urease activities were modulated by Aerva persica in collaboration with fungi such as A. terreus, A. niger, and A. flavus (b) amidase concentration was effectively controlled through litter of a legume plant species T. purpurea in collaboration with fungi such as A. fumigatus, A. niger, and A. persica (c) Beta-glucosidase and dehydrogenase were most supportive by combined

<table>
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<tr>
<th>Variables</th>
<th>AcP</th>
<th>AIP</th>
<th>BG</th>
<th>De</th>
<th>Ur</th>
<th>Am</th>
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<tbody>
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<td>-2.27</td>
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<td>A. persica + A. fumigatus</td>
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<td>0.72</td>
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AcP = Acid Phosphatase, AIP = Alkaline Phosphatase, BG = Beta-glucosidase, De = Dehydrogenase, Ur = Urease, Am = Amidase
action of *C. procera* with *A. fumigatus, A. niger, A. flavus* (d) *C. procera-A. terreus* and *T. purpurea-A. flavus* complexes, on the other hand, were more effective for alkaline phosphatase and (e) acid phosphatase having more proximity with *T. purpurea-A. terreus*. Our correlation analysis revealed the complex relationships between enzymes using an enzyme-specific approach. Canonical Correlation Analysis (CCoA) revealed that urease did not correlate with any other enzymes studied during the first withdrawal period, but it did correlate with almost all enzymes except amidase during the second sampling period. Surprisingly, amidase was significantly correlated with all studied enzymes except urease during the first sampling period, but only alkaline phosphatase ($\text{AlP}_2 r^2 = 0.54$) during the second sampling period. Correlations among weighted enzymes revealed that acid phosphatase and alkaline phosphatase were both related to all other enzymes except amidase. These findings are consistent with those of Acosta-Martinez *et al.* (2003) and Mathur (2020). The region's first compressive effort employing locally accessible desert plant species and fungus resulted in the development of a soil quality index. This index considers how time, litter, and different fungal species behave. This index, however, may not be applicable to other scenarios such as cultural practices (addition of fertilizers, irrigation types, soil tillage, fallow land practice, and so on), land use (forest land, agricultural land, orchid, and so on), mining, and rehabilitation practices because the studied data set is restricted to controlled conditions. Such practices unquestionably have a unique interaction with the biochemical properties of soil. Future work will therefore involve validating and improving this index using numerous scenarios.

**Conclusion**

The current study is the first from the hot, arid region of India to look at the interaction and temporal effects of diverse fungus species and wild plant species used as litter sources on soil enzyme concentrations. Using PCA and CCoA, we developed a bio-chemical soil quality index, where each enzyme was weighted with numerical values based on their associations with the kind of litter (plant species), fungal species types, and withdrawal period (temporal factor). The idea of sustainable utilization of arid wild species that are neither cultivated nor have direct market potential can be used to determine the importance of the current study (provisional ecosystem services). They can therefore be utilized as regular litter for a variety of crops to maintain soil fertility.

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**Conflict of interest**

The authors declare that they have no conflict of interest.

**References**


Interactive effects of non-fodder litter and fungal species


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