10 Soil and Crop Management Effects on Soil Microbiology

Ann C. Kennedy, Tami L. Stubbs, and William F. Schillinger

CONTENTS

Introduction .......................................................... 295
Soil Microbial Communities ........................................ 296
Microbial Diversity .................................................. 298
Nutritional Strategies .................................................. 300
Management Effects on Soil Microbial Communities ........ 301
Plant Influences ....................................................... 302
Roots and Microflora ................................................ 302
Plant Competition .................................................... 303
Plant Diversity/Crop Rotation ...................................... 303
Crop Residue ......................................................... 304
Resources .............................................................. 306
Nutrient Status/Cycling ....................................... 306
Plant Growth-Regulating Compounds .................... 306
Amendments ......................................................... 308
Agromicrobials ....................................................... 308
Arbuscular Mycorrhiza (AM) ................................... 308
Biological Control .................................................. 309
Organic/Low-Input Farming ..................................... 309
Genetically Modified Organism (GMOs) ................. 310
Disturbance .......................................................... 310
Tillage ................................................................. 311
Grading ............................................................... 315
Strategies for Managing Microorganisms ................ 315
Conclusions ........................................................ 316
References .......................................................... 316

INTRODUCTION

Life in soil is responsible for a multitude of processes vital to soil function. Microorganisms can have a profound effect on plant growth, soil organic matter (SOM) accumulation, and soil condition or soil quality. For more than 2.5 million years, microorganisms have been a life force on earth, establishing communities well before any other life forms. Since the beginning, natural selection has ever increased the microbial diversity in soils. All life is dependent on microbial processes (Price, 1988), and SOM transformations are due to microbial processes (Alten, 1999). In turn, SOM sustains that life and is crucial to soil function. Strategies that increase SOM tend to enhance soil biological processes and vice versa. Understanding these processes and implementing strategies to enhance SOM improve soil quality and maintain biological diversity will help attain sustainable agriculture.

295
Soil Quality

Soil quality is defined as the capacity of a soil to function within ecosystem boundaries to sustain biological productivity, maintain environmental quality, and promote plant and animal health (Doran and Page, 1994). It is easy to visualize a healthy, rich soil and to remember its smell. Descriptive and analytical measurements of the physical, chemical, and biological properties are sometimes used to characterize soil quality. Indicators of soil quality are needed to measure changes in soil function that occur because of alternation in management. Total organic matter can be an indicator; however, changes in total SOM usually respond very slowly to changes in management and thus lack sensitivity. Soil organisms contribute to the maintenance of soil quality because they control many key processes. Soil microorganisms and their communities are continually changing and adapting to changes in their environment. A high-quality soil is biologically active and contains a balanced population of microorganisms. The dynamic nature of soil microorganisms makes them a sensitive indicator to assess changes in soil quality due to management (Kennedy and Papendick, 1995).

This chapter explores microbiological changes occurring with soil and crop management in farming systems. Our discussion of community structure includes microbial survival strategies and delineation of groups of organisms, such as bacteria and fungi, nutritional-based groups or species, and functional determinations. Our goal is to describe changes in the soil biota with management to help identify soil microbial parameters useful in assessing management practices for conserving and enhancing SOM, soil quality, and crop production.

Soil Microbial Communities

The number of microbial species on earth is estimated to exceed 100,000 and may be more than a million (Hawksworth, 1991b; American Society for Microbiology, 1994). Unfortunately, only 3 to 10% of the earth’s microbial species have been identified or studied in any detail (Hawksworth, 1991a). The full potential of these groups of organisms has not been explored. The diversity of microorganisms is thought to exceed that of any other life form (Torsvik et al., 1990; Ward et al., 1992). It is estimated that several thousand genomes are present in each gram of soil (Torsvik et al., 1990).

Soil microorganisms are responsible for many soil processes, such as SOM turnover, soil humus formation, cycling of nutrients, and building soil tilth and structure (Table 10.1; Lasch, 1983; Wood,

<table>
<thead>
<tr>
<th>TABLE 10.1</th>
</tr>
</thead>
<tbody>
<tr>
<td>Beneficial Functions of Soil Microorganisms in Agricultural Systems</td>
</tr>
<tr>
<td>• Release plant nutrients from insoluble inorganic forms</td>
</tr>
<tr>
<td>• Decompose organic residues and release nutrients</td>
</tr>
<tr>
<td>• Form beneficial soil humus by decomposing organic residues and through synthesis of new compounds</td>
</tr>
<tr>
<td>• Produce plant growth-promoting compounds</td>
</tr>
<tr>
<td>• Improve plant nutrition through symbiotic relationships</td>
</tr>
<tr>
<td>• Transform atmospheric nitrogen into plant-available N</td>
</tr>
<tr>
<td>• Improve soil aggregation, aeration, and water infiltration</td>
</tr>
<tr>
<td>• Have antagonistic action against insects, plant pathogens, and weeds (biological control)</td>
</tr>
<tr>
<td>• Help in pesticide degradation</td>
</tr>
</tbody>
</table>
1991). These functions are performed by many different genera and species. Beneficial soil bacteria enhance plant performance by increasing solubility of minerals (Olson, 1982). N. fixation (Albrecht et al., 1981); producing plant hormones (Brown, 1972; Arshad and Frankenberger, 1998); and suppressing harmful pathogens (Chang and Kommedahl, 1968). Beneficial mycorrhizal fungi can enhance plant growth by increasing nutrient (Fitter, 1977; Hall, 1978; Rovira, 1978; Ocampo, 1986); and water (Tinker, 1976) uptake and soil structure by enhancing aggregate formation and stability (Wright and Uppahaya, 1998; Chapter 6). Conversely, plant-suppressive bacteria impair seed germination and delay plant development by producing phytotoxic substances (Wolz, 1978; Snaulow and Schroth, 1982; Alstrom, 1987; Schippers et al., 1988). Pathogenic fungi greatly reduce the survival, growth, and reproduction of plants (Shippton, 1977; Bruehl, 1987; Scurdon, 1987). Another example of the importance of microorganisms to agriculture is the production of antibiotics by strains of fluorescent 

\textit{Pseudomonas} bacteria that suppress the root disease take-all (\textit{Gaumannomyces graminis var. tritici}) in continuous winter wheat (\textit{Triticum aestivum L.}) cropping systems (Thomashow and Weider, 1988).

Specific microorganisms can be manipulated to produce beneficial effects for agriculture and the environment (Lynch, 1983; e.g., rhizobia to increase plant available N (Speent, 1979), mycorrhizal associations to assist nutrient and water uptake (Sylvia, 1998; Mohammad et al., 1995); or biological control of plant pests to reduce chemical inputs (Cook and Baker, 1993; Kennedy et al., 1991). Bacterial or fungal inoculants can be added to soil to aid in the biocontrol of harmful substances such as petroleum hydrocarbons (Rhykerd et al., 1999; Mohr and Steward, 2000), polycyclic aromatic hydrocarbons (Allen et al., 1999), and a wide range of environmental pollutants (Cameron et al., 2000).

The presence of a large and diverse soil microbial community is crucial to the productivity of any agroecosystem. This diversity is influenced by almost all crop and soil management practices, including the type of crops grown. Plants and their exudates influence soil microorganisms and the soil microbial community found near roots (Dunfield et al., 1998; Ibekwe and Kennedy, 1998; Ohtonen et al., 1999). In turn, the composition of the microbial community influences the rate of residue decomposition and nutrient cycling in agroecosystems (Barea et al., 1993). The basic groups of microorganisms in soil are bacteria (including actinomycetes), fungi, algae, and protozoa. Bacteria and fungi are decomposers involved in nutrient cycling and SOM processes and are critical in the functioning of the soil food web. Ninety-five percent of plant nutrients must pass through these organisms to higher trophic levels (Moore, 1994).

Bacteria are diverse metabolically and perform numerous functions. Bacteria consent SOM into carbon (energy sources) used by others in the soil food web, break down pesticides and pollutants, and immobilize and maintain valuable nutrients such as N in the root zone. Bacteria quickly colonize the substrate-rich rhizosphere (Figure 10.1). Actinomycetes are a specialized group of soil bacteria that degrade plant materials such as cellulose. Actinomycetes are important in mineralization of nutrients and some can produce antibiotics. Actinomycetes can tolerate low soil water potential better than other bacteria, but are not tolerant of low soil pH (Alexander, 1998).

Fungi, like bacteria, are vital members of the food web. Fungi are especially important at lower pH, because many bacteria are adversely affected by acid soils. Fungi are able to withstand unfavorable conditions, such as water stress and extreme temperatures, better than other microorganisms (Pappenfleck and Campbell, 1975). They are critical for residue decomposition and accumulation of stable SOM fractions through breakdown of more complex carbon sources such as cellulose, lignin, and other organic materials. These decomposition products are then available for use by other organisms. Fungal mycelia bind soil particles together to form aggregates that increase water-holding capacity and infiltration and reduce erosion. Fungi can be saprophytes on detrital material or in associations with plant roots (Swift and Boddy, 1984). The more calciferous material left from decomposition then accumulates as SOM. Hyphae of arbuscular mycorrhizal (AM) fungi produce the protein glomalin, which improves soil structure (Chapter 6).
Algae occur in soil at populations of $10^5$ to $10^6$ g$^{-1}$ soil, far fewer than bacteria and fungi. The greatest populations of algae are found in moist soil, but their numbers decrease with increasing soil depth. Some algal species are nitrogen fixers and produce mucigel, which can stabilize soil aggregates. Algae are susceptible to soil disturbance and can be good indicators of soil quality. Their populations increase in agricultural systems with reduced disturbance where the surface soil and residue maintain a higher moisture regime for longer periods (Harris et al., 1995), and as a result foster algal growth on the soil surface.

Protozoa are found at populations of $10^3$ to $10^4$ g$^{-1}$ soil. These single-celled organisms prey on bacteria and other microorganisms, and thus regulate bacterial populations (Opperman et al., 1989) and influence SOM decomposition by regulating decomposer populations. Protozoa are crucial to the functioning of soil and other ecosystems because of their role in nutrient cycling and in providing energy for other microorganisms, plants, and animals (Fonnesen, 1999). Fluctuations in microbial populations with tillage affect protozoan populations because protozoa feed on these organisms. Protozoa can be useful indicators of changes in soils because their populations react rapidly to changes in the environment (Fonnesen, 1999).

MICROBIAL DIVERSITY

There are two primary ways that diversity can be evaluated: species diversity and functional diversity. Functional diversity can be a better parameter than species diversity to learn about soil processes and stable SOM fraction formation (Mikola and Siltälä, 1998). However, it is often difficult to obtain actual measurements of functional diversity, whereas evaluating species diversity, when specific species can be assessed, is easier. The number of organisms in various microbial groups might not be sufficient to illustrate the breadth of diversity found in the soil. Although an increase in microbial populations, such as SOM or CO$_2$, can be an indicator of increased functioning, it might not necessarily be due to higher functional diversity. One of the earliest studies involving soil diversity and soil respiration (Salomaa, 1981) established differences in bacterial and fungal diversity by inoculating soil with varying soil suspensions. Respiration rate was reduced with the lower dilution of the assumed lower microbial diversity. The true extent or dimension of the diversity of soil microorganisms is unknown, although molecular investigations suggest that culturing techniques underestimate population numbers (Holben and Tiedje, 1988; Tosvik et al., 1996). The functioning of a group of organisms is as important as the number of species in regulating ecosystem processes (Grime, 1997; Wardle et al., 1997; Bardgett and Shine, 1999). How much diversity is required to ensure sustainable and efficient SOM turnover, as well as other important functions? Greater use of diversity indices is limited by absence of detailed information on the composition.
of microbial species in soil (Torsvik et al., 1998). Diverse systems are thought to have higher agricultural productivity, resilience to stress, and be more sustainable and provide risk protection (Giller et al., 1997; Wolters, 1997). A diverse system has a wider range of function with more interactions among microorganisms that influence each other to varying degrees. A higher number of different types of organisms present in a system means there are more to perform various processes and fill a niche that might not be filled if a particular group is inhibited by stress (Andren et al., 1995).

Substrate-utilization patterns have been used to obtain fingerprints of community structure (Garland, 1996; Bossio and Scow, 1995; Haack et al., 1995; Wusche et al., 1995; Zak et al., 1994). These measures can also indicate functional diversity, metabolic potential (Dean et al., 1996; Haack et al., 1995; Wusche et al., 1995), and nutritional strategies (Zak et al., 1994). Soil microbial communities as indicated by whole-soil fatty acid methyl ester (FAME) analysis can be differentiated by geographic region (Kennedy and Busacca, 1995) and cropping pattern (Cavigelli et al., 1995). The living microbiological component of soil can be estimated by phospholipid fatty acid (PLFA) analyses (Zelles et al., 1994). Another method for measuring microbial diversity is the DNA hybridization technique, which uses similarity indices. This technique illustrated that extracted bacteria and whole-community DNA had 75% similarity (Griffiths et al., 1996). The DNA microarray technology can be used to rapidly analyze microbial communities based on phylogenetic groupings and increases the ease of molecular analyses (Guschen et al., 1997). These analyses can help further understand the changes occurring among soil communities with various management practices.

Microbial diversity can be linked to susceptibility and resiliency of soils to stress, and thus might affect some soil functions such as SOM decomposition. Partial fumigation of grassland soils produces differing degrees of diversity, with longer fumigation times producing soils with less diversity. There is no direct correlation between the progressive fumigation to reduce diversity and measures of soil function, such as soil microbial biomass, soil respiration, and N mineralization. However, soils with lower diversity initially have more ability to decompose added grass residues (Griffiths et al., 2000). There is greater susceptibility to copper toxicity with decreasing diversity. Soils that contained the most diverse populations showed the greatest resilience to copper-induced stress by quickly rebounding, as shown by an increase in grass residue decomposition rates. In a similar study, no differences were seen in decomposition of Medicago sativaL even though the residues were added to both organic and conventionally farmed soils with different SOM levels (Gunapala et al., 1998). Organically farmed soils initially contained a more abundant microbial population as measured by microbial biomass C and N. When organic amendments were added, soil from the conventionally farmed system increased in microbial biomass C to a level that was comparable to the soil in the organic system. The biotic community in the conventionally farmed soil was sufficient and could respond to added substrate as well as the organic soils did. The microbial communities in this study functioned adequately whether from conventional or organic farming systems (Gunapala et al., 1998).

A reduction in functional diversity does not necessarily impede a soil's ability to decompose residue. Degens (1998) used fumigation to alter functional diversity in a grassland and measured in situ catalytic potential (Degens and Harris, 1997) to characterize the ability of the soil community to mineralize C substrates, with substrate added to the soil directly. The functional indices were different among fumigated, unfumigated, and fumigated and inoculated with untreated soil. There was no relationship between functional diversity and decomposition of wheat straw added into these systems. Water potential might have been the overriding factor controlling decomposition rate, because soils with reduced functional diversity continued decomposing the wheat straw under optimum moisture conditions.

Diversity of soil microorganisms can impact pathogens and pathogens, thus influencing their impact on plant growth. Decreased diversity of actinomycetes, some of which are antagonists of pathogens, correlated with an increase in pathogens of tomato (Workneh and van
Bruggen, 1995). *Cochliobolus sativus*, a pathogen causing a serious disease in wheat, was found in higher numbers, and individual isolates exhibited greater pathogenicity in a continuous wheat rotation than in wheat in a 3-year rotation. This increased pathogenicity was attributed to a reduced number in microbial diversity (El Nashaw and Stack, 1989). Take-all decline of wheat occurs after several years of monoculture and is correlated with the appearance of several different types of organisms and a lack of certain soil organisms in the soil (McCoy et al. 2001). The impact of the microbial community on pathogen load and pathogenicity is complex and changes with the make-up and diversity of the community.

Assuming, all functional groups are present, more microbial diversity might not necessarily be crucial to ecosystem functioning. Soil biodiversity and nutrient cycling are linked in a study of Nigerian tropical soils (Swift et al., 1998). A study comparing native bush soils with those under cultivation showed greater abundance and diversity of soil fauna in the former, but little difference in decomposition of surface residues. Although variation in species richness might not be discernible in many environments, differences can be important in stressed systems or when conditions are altered (Yachi and Loreau, 1999). Organic matter accumulation and rate of decomposition can be important, although slowly changing indicators of ecosystem functioning in less-stressed systems.

The quality and quantity of substrate can affect community structure. Griffiths et al. (1999) used synthetic root exudates to study community structure. Microbial community changes occurred with continual substrate loading increases, and fungi dominated over bacteria in high-substrate conditions. Different organisms have the ability to be a dominant portion of the community when changes in efficiency occur because of changes in optimal growth factors, substrate quality, or substrate concentration. This study is important when considering additions of organic amendments to agricultural soils.

**NUTRITIONAL STRATEGIES**

The concept of r- and K-strategies is an ecological classification system based on the ability of an organism to survive in different environments (MacArthur and Wilson, 1967). To indicate two contrasting methods of selection in animals, K refers to the carrying capacity and r to the maximum intrinsic rate of natural increase (rmax). Although most microorganisms are considered r-strategists and plants and animals K-strategists, there are differences in growth strategies among microorganisms (Andrews and Harris, 1986; Table 10.2). K-strategists favor competition at carrying capacity, whereas r-strategists take advantage of easily available substrates with fast growth rates to facilitate colonization of new habitats in response to a flush of nutrients or other fluxes. Organisms can be both r- and K-strategists, depending on circumstances. An organism can exhibit an r-strategy when faced with fresh resources and an unstable environment, i.e., when organic amendments are applied, but become a K-strategist after resources are depleted and only more recalcitrant substrate is available. Age of plant roots and plant type can also influence the dominant strategy. K-strategists were found in higher numbers on older wheat roots than in young roots (De Leij, 1993). The root surface of ryegrass had more K-strategists than that of white clover (Sarathchandra et al., 1997). Spore formation is a tactic of r-strategists to survive during low nutrient availability. Although the initial colonists of a residue might be r-strategists, organisms involved in humus degradation or lignin and cellulose degradation are K-strategists. Most soil bacteria are generally considered r-strategists, whereas fungi and actinomycetes are usually K-strategists (Bottomley, 1998).

The type of strategy used and various processes influence soil and plant functioning. For example, when root exudates are added to soils contaminated with heavy metals, certain bacterial populations increased, the dominance of various strategy organisms depending on availability of substrate and soil conditions (Koordj and van Elsas, 2000). Exudates added to these polluted soils decreased the overall diversity in favor of r-strategists, whereas K-strategists dominated soils not amended with exudates. In another study, organisms with the same community structure exhibited
<table>
<thead>
<tr>
<th>Characteristic</th>
<th>r-strategist</th>
<th>K-strategist</th>
</tr>
</thead>
<tbody>
<tr>
<td>General</td>
<td>Rapid reproductive rate, extreme reproduction</td>
<td>Adapt to environment, stable and permanent</td>
</tr>
<tr>
<td>Growth rate</td>
<td>Rapid</td>
<td>Moderate</td>
</tr>
<tr>
<td>Substrate-utilization efficiency</td>
<td>Low efficiency</td>
<td>Higher efficiency</td>
</tr>
<tr>
<td>Diversity of substrates utilized</td>
<td>Simple, readily available, not resource limited</td>
<td>Complex, diverse, may be resource limited</td>
</tr>
<tr>
<td>Phenotype</td>
<td>Polymorphic or monomorphic</td>
<td>Mono-specific</td>
</tr>
<tr>
<td>Morphology</td>
<td>Smaller cells, mycelium not highly differentiated</td>
<td>Larger cells, well-developed mycelium</td>
</tr>
<tr>
<td>Reproduction</td>
<td>Simple genetic exchange, rapid rate</td>
<td>Complex genetic exchange, slow rate</td>
</tr>
<tr>
<td>Population dynamics</td>
<td>Explosive, density-dependent</td>
<td>Stable, density dependent by competition or gauging, equilibrium dynamics at or near carrying capacity</td>
</tr>
<tr>
<td>Tolerance to niche overlap</td>
<td>High tolerance</td>
<td>Low tolerance</td>
</tr>
<tr>
<td>Residual colonists</td>
<td>Early</td>
<td>Late</td>
</tr>
<tr>
<td>Competitive adaptations</td>
<td>Few</td>
<td>Many</td>
</tr>
<tr>
<td>Microbial types</td>
<td>Cyanobacteria, dinoflagellates, blooms; Aspergillus, Penicillium; Pseudomonas, Bacillus; hemeophiles, spore formers</td>
<td>Humin, lignin and cellulose degraders, spores, vibrios, symbiotic, Corynebacterium and bacteria</td>
</tr>
</tbody>
</table>


different catalytic response profiles when grown in different soil environments, illustrating the effect of management on the community's functional diversity (Degenz, 1999).

In addition to r- and K-strategies, oligotrophic response can be used to characterize organisms in an ecosystem. Organisms are grouped based on their nutritional strategies. Oligotrophs are organisms that grow under low nutrient supply and subsist on more resistant SOM, whereas copiotrophs flourish in nutrient-rich environments. Bacteria with enhanced growth under high nutrient concentrations are described as copiotrophic. Oligotrophs are more prevalent than copiotrophs in low-substrate concentrations. The proportion of copiotrophs to oligotrophs varies over time; the ratio of copiotrophic to oligotrophic increased immediately after cover crop residue incorporation but decreased 26 d later when readily available C declined (Hu et al., 1999). High quantities of readily available C early in the experiment might have inhibited oligotrophic growth (Hu et al., 1999).

Crop selection, region of the root system, and proximity to plant roots influence the number of oligotrophs and copiotrophs as well as their ratio (Maloney et al., 1997). It is important to understand the response of the microbial community to varying levels of C inputs to better manage for residue decomposition, competition with crop pathogens, and to improve the survival of introduced microorganisms (Hu and van Bruggen, 1997). Analysis of microbial community survival and nutritional strategies can aid in investigations of changes with management.

**MANAGEMENT EFFECTS ON SOIL MICROBIAL COMMUNITIES**

Throughout each season, crop management, resource additions, or soil disturbance influence the microbial community (Figure 10.2). Each crop or soil management practice affects the microbial community and formation or degradation of SOM.
The rhizosphere is a dynamic zone of soil under the influence of plant roots (Bown and Rovira, 1999; Pinton et al., 2001) and has high microbial numbers (Grayston et al., 1998), activity, and diversity (Kennedy, 1998). The rhizosphere is a region of intense microbial activity because of its proximity to plant root exudates, making rhizosphere microbial communities distinct from those of bulk soil (Carl and Treslove, 1986; Whipp and Lynch, 1986). Nutrients exuded by the root or germinating seed stimulate microbial activity (Renaat and Katznelon, 1961). Interactions between plants and rhizosphere microorganisms can play a critical role in plant competition. Competitive interaction among plants can also be important to develop rhizosphere soil communities. Free-living bacteria and fungi from rhizospheres of different pairs of plant species in two fields utilized different substrates and grew differently in the presence of antibiotics, osmotic stresses, and zinc (Westover et al., 1997). Results from these two fields suggest that adjacent plant species influence populations of rhizosphere bacteria and fungi, creating local microscale heterogeneity in rhizosphere soil (Westover et al., 1997). Similar results have been obtained for AM communities associated with certain grass species (McGonigle and Fitter, 1990; Johnson et al., 1992), rhizosphere bacterial populations associated with particular wheat genotypes (Neal et al., 1973), and root bacterial communities following bacterial inoculation (Gilbert et al., 1993).

Composition of plant species can influence the microbial community because of differences in chemical composition of root exudates (Christensen, 1989). Peas and oats exude different amounts of amino acids (Rovira, 1956). Environmental factors regulating plant growth can affect root exudation, including temperature (Rovira, 1959; Vanceur, 1967; Martin and Kemp, 1980), light (Rovira, 1956), and soil water (Martin, 1977). Plants significantly influence the make-up of their own rhizosphere microbial communities (Miller et al., 1989). This is the result of different plant species and cultivar transporting varying amounts of C to the rhizosphere (Liljeroth et al., 1990) as well as different compositions of exudates. Bekwe and Kennedy (1999) showed that wheat

PLANT INFLUENCES

Roots and Rhizosphere

The rhizosphere is a dynamic zone of soil under the influence of plant roots (Bown and Rovira, 1999; Pinton et al., 2001) and has high microbial numbers (Grayston et al., 1998), activity, and diversity (Kennedy, 1998). The rhizosphere is a region of intense microbial activity because of its proximity to plant root exudates, making rhizosphere microbial communities distinct from those of bulk soil (Carl and Treslove, 1986; Whipp and Lynch, 1986). Nutrients exuded by the root or germinating seed stimulate microbial activity (Renaat and Katznelon, 1961). Interactions between plants and rhizosphere microorganisms can play a critical role in plant competition. Competitive interaction among plants can also be important to develop rhizosphere soil communities. Free-living bacteria and fungi from rhizospheres of different pairs of plant species in two fields utilized different substrates and grew differently in the presence of antibiotics, osmotic stresses, and zinc (Westover et al., 1997). Results from these two fields suggest that adjacent plant species influence populations of rhizosphere bacteria and fungi, creating local microscale heterogeneity in rhizosphere soil (Westover et al., 1997). Similar results have been obtained for AM communities associated with certain grass species (McGonigle and Fitter, 1990; Johnson et al., 1992), rhizosphere bacterial populations associated with particular wheat genotypes (Neal et al., 1973), and root bacterial communities following bacterial inoculation (Gilbert et al., 1993).

Composition of plant species can influence the microbial community because of differences in chemical composition of root exudates (Christensen, 1989). Peas and oats exude different amounts of amino acids (Rovira, 1956). Environmental factors regulating plant growth can affect root exudation, including temperature (Rovira, 1959; Vanceur, 1967; Martin and Kemp, 1980), light (Rovira, 1956), and soil water (Martin, 1977). Plants significantly influence the make-up of their own rhizosphere microbial communities (Miller et al., 1989). This is the result of different plant species and cultivar transporting varying amounts of C to the rhizosphere (Liljeroth et al., 1990) as well as different compositions of exudates. Bekwe and Kennedy (1999) showed that wheat

PLANT INFLUENCES

Roots and Rhizosphere

The rhizosphere is a dynamic zone of soil under the influence of plant roots (Bown and Rovira, 1999; Pinton et al., 2001) and has high microbial numbers (Grayston et al., 1998), activity, and diversity (Kennedy, 1998). The rhizosphere is a region of intense microbial activity because of its proximity to plant root exudates, making rhizosphere microbial communities distinct from those of bulk soil (Carl and Treslove, 1986; Whipp and Lynch, 1986). Nutrients exuded by the root or germinating seed stimulate microbial activity (Renaat and Katznelon, 1961). Interactions between plants and rhizosphere microorganisms can play a critical role in plant competition. Competitive interaction among plants can also be important to develop rhizosphere soil communities. Free-living bacteria and fungi from rhizospheres of different pairs of plant species in two fields utilized different substrates and grew differently in the presence of antibiotics, osmotic stresses, and zinc (Westover et al., 1997). Results from these two fields suggest that adjacent plant species influence populations of rhizosphere bacteria and fungi, creating local microscale heterogeneity in rhizosphere soil (Westover et al., 1997). Similar results have been obtained for AM communities associated with certain grass species (McGonigle and Fitter, 1990; Johnson et al., 1992), rhizosphere bacterial populations associated with particular wheat genotypes (Neal et al., 1973), and root bacterial communities following bacterial inoculation (Gilbert et al., 1993).

Composition of plant species can influence the microbial community because of differences in chemical composition of root exudates (Christensen, 1989). Peas and oats exude different amounts of amino acids (Rovira, 1956). Environmental factors regulating plant growth can affect root exudation, including temperature (Rovira, 1959; Vanceur, 1967; Martin and Kemp, 1980), light (Rovira, 1956), and soil water (Martin, 1977). Plants significantly influence the make-up of their own rhizosphere microbial communities (Miller et al., 1989). This is the result of different plant species and cultivar transporting varying amounts of C to the rhizosphere (Liljeroth et al., 1990) as well as different compositions of exudates. Bekwe and Kennedy (1999) showed that wheat
barley (Hordeum vulgare L.), pea (Pisum sativum L.), joined goastrum (Acetosyringanum arborescens) and downy brome (Bromus tectorum L.) grown in two soil types had different rhizosphere microbial communities. Barley cultivars differed in the abundance of fungi and bacteria present in their rhizospheres and rhizosphere, and these differences were sustained over different stages of plant growth (Ljierhorst and Balph, 1988). Two corn (Zea mays) cultivars (Fasiumus susceptible and resistant) and grass (Poa pratensis) lines (disease susceptible and resis-
tant) differed in the numbers of rhizosphere bacteria, with susceptible lines having the highest numbers (Müller et al., 1989). These results were obtained even with no known presence of the pathogens. The rhizosphere microbial communities as determined by Biolog®Biolog® GN microtiter plates, Hayward, CA differed with plant species of wheat, ryegrass (Lolium perenne), bentgrass (Agrostis capillaris), and clover (Trifolium repens). Plant species affected C- utilization profiles of the rhizosphere microbial communities of wheat, ryegrass, clover, and bentgrass. Microorganisms in the rhizospheres of wheat, ryegrass, and clover had higher utilization of C sources than in the bentgrass rhizosphere. Soil type, however, did not affect the rhizosphere soil microbial community profiles (Grayston et al., 1998). In natural plant communities, different plant combinations exhibited unique rhizosphere populations of free-living bacteria and fungi with differing abilities to utilize C substrates and withstand stresses (Westcove et al., 1997). Unique C-source utilization patterns among rhizosphere communities of hydroponically grown wheat, white potato (Solanum tuberosum), soybean (Glycine max), and sweet potato (Ipomoea batatas) were found by using Biolog plates (Garland, 1996). C-source utilization patterns could distinguish among soils from six plant communities (Zak et al., 1994).

Substrate-utilization patterns have been used successfully to differentiate bacteria associated with different cropping and management practices (Garland, 1996; Zak et al., 1994). Crop effects can be associated with plant exudates as a result of the enhanced utilization or inhibition of substrates, similar to the organic content of root hairs, mucilage, or root cell lysates of the particular crop (Garland, 1996). Bossio and Scow (1995) found pattern differences associated with rice straw treatments and flooding. These systems are highly reactive to changes in their environment and can thus serve as easily attained, reliable fingerprints of community shifts as a function of substrate use.

Plant Competition

Competitive interactions of the plant can influence plant productivity and are affected by soil microorganisms, such as mycorrhizal fungi (Crowell and Boerner, 1988; Herrick and Wijson, 1989; Allen and Allen, 1990) and Rhizobials (Turkington et al., 1988; Turkington and Klein, 1991; Chanway and Holm, 1993). Evidence suggests that soilborne pathogens affect plant competitiveness and plant succession (Van der Pette and Peters, 1997). A plant-resistant species, sand fescue (Festuca rubra subsp. arvense), outcompeted the susceptible species, marram grass (Ammophila arenaria), when both coastal grasses were exposed to pathogens (Van der Pette and Peters, 1997).

Plant Diversity/Crop Rotation

Plant species and numbers can drive the make-up of the microbial community and the diversity of rhizosphere microbial populations. The above- and below-ground plant community can influence rhizomicrobial spatial heterogeneity in soil. Aboveground shoot material contributes organic material to the surface layers of soil. Decaying root systems also function as a source of nutrients for the surrounding microorganisms (Swinnen et al., 1995). Compared with monocropping, crop rotation can improve conditions for diversity in soil organisms because of variability in type and amount of organic inputs, and allow for time periods, or breaks, when there is no host available for a particular pest (Alilieri, 1999). Diversity in crop rotation can allow for higher C inputs and diversity of plant material added to soils, depending on the residue level and carbon quality of the crops in
rotation. Crop rotation enhances beneficial microorganisms, increases microbial diversity, interrupts the cycle of pathogens, and reduces weed and insect populations. Legumes in a crop rotation supply symbiotically fixed nitrogen to the system, use less water than many other crops, and reduce pathogen load. Studies have long shown the positive effects of crop rotation on crop growth, attributing these to changes in composition of microbial community (Shipston, 1977; Cook, 1981; Johnson et al., 1992).

Crop rotation and plant cover affected soil microbial biomass C and N of long-term field experiments in Iowa, with the highest values found in the longer rotations (4 years vs. 2 years) and monocropping systems, and the lowest in the continuous corn–rye–barley system. The varied diversity and quality of crop residues, amount of readily decomposable organic material, and root density led to increased soil microbial biomass under crop rotation. N fertilization did not affect microbial biomass in these studies (Moore et al., 2000).

Alienophotic interactions can occur between crops and weeds, between two crops, from desiccation stress and weed redss, and from crop and weed exudates (Anaya, 1999). Neopatogenic allelopathic bacteria can produce plant-inhibiting compounds (Barazani and Friedman, 1999). Crop rotation can be used to alleviate the allelopathic or autopathic effects a crop plant might have on itself. Monocropping encourages proliferation of allelopathic bacteria (Barazani and Friedman, 1999).

By rotating crops, it is possible to lessen the negative effects a crop might have on itself and on subsequent crops (Rice, 1995). The populations and aggressiveness of pathogens can be altered with crop rotation, illustrating changes in microbial diversity and function due to management (El Nashaar and Stack, 1989). In a long-term study, Cochliobolus sativus, a pathogen of spring wheat, was found in higher numbers and individual isolates exhibited greater aggressiveness or ability to cause severe disease in continuous wheat, when compared with wheat in a 3-year rotation. Continuous monocropping led to changes in the soil community, which increased pathogen load and reduced barley growth compared with that by grains in multiple-crop rotation (Olsson and Gerhardtson, 1992). Continuous monocropping of wheat, however, can lead to suppression of the take-all pathogen or take-all decline. This natural defense occurs in soils in the presence of fluorescent pseudomonas bacteria that produce the antibiotics phenazine and phthorococinol (Mazzola et al., 1995). Barley plants produce compounds that can help protect it from fungus (Drechslera arenaria) and armyworm (Mythimna conogeta) larvae (Lover and Housh, 1995).

Crop rotation can influence root colonization by mycorrhizae. In years following spinach (Spinacia oleracea) and bell pepper (Capsicum annum), spore populations of most species of AM were depressed and had lower infectivity compared with those in years following wheat, rice, or corn (Douds et al., 1997). Cover crops, such as autumn-sown cereals or vetches, increase the AM inoculum potential for subsequent crops (Boswell et al., 1998; Galvez et al., 1995). Cover crops aid in maintaining a viable mycelial network. A cover crop of winter wheat inoculated with AM increased AM infection rate, and it turned increased the growth and yield of a subsequent corn crop (Boswell et al., 1998). Soil from no-till, low-input fields with a hairy vetch cover crop maintained higher levels of colonization by indigenous AM than soils that had been tilled or received high-input management (Galvez et al., 1995). Use of cover crops can maintain AM when inoculum levels might otherwise be low and enhance infection of the subsequent crop.

Crop Residue

Additions of crop residue are critical to maintain or increase SOM levels in agricultural soils (Figure 10.3). Cropping systems vary in residue quality and quantity, the microbial community supported, contributions to SOM, and ability to withstand the effects of disturbance. The residue decomposition process depends on the organism present, type of SOM, and environmental conditions (Martin, 1933). Residue decomposition can also be affected by availability of carbon for microbial growth, physical separation because of landscape position, soil horizonation, or encapsulation of SOM in
soil aggregates and clay peds, and low N availability (Paul, 1984). Water, temperature, soil pH, aeration and oxygen supply, nutrient availability, crop residue composition, C:N ratio of crop residue, and microfactors are critical factors in residue degradation (Parth and Pappas, 1978). Residue quality and management influences the composition of the microbial community. Higher numbers of culturable bacteria, including actinomycetes, were observed from decomposing soybean residues in buried bags than from wheat or corn. Fungal populations were highest on corn and lowest on wheat (Breder and Wagner, 1981). Sorghum (Sorghum bicolor (L.) Moench) residues buried by conventional tillage contained greater fungal hypyal length but fewer actual fungal propagules than with no-till, whereas no-till mineral soil had greater fungal hyphal length but no difference in propagule counts compared with conventionally tilled soils (Beare et al., 1993). In their study of sorghum residues, Beare et al. (1993) identified genera of fungi that were specialized for surface residue, whereas buried residues contained no specialized fungal community.

Crop species (Cookson et al., 1998) and cultivar (Chalaux et al., 1995) vary in their decomposition rate as well as the microbial population to support the amount of C mineralized from crop residues depends on the type of residue and residue composition (Herfisens and Breeland, 1999). Amino acids and simple sugars, which are metabolized most rapidly in the residue-decomposition process, support populations of bacteria. More complex compounds such as cellulose and lignin are broken down by xylanases or oligotrophs. Lupine (Lupinus albus) residue decomposes more rapidly and supports higher populations of bacteria and fungi than does wheat or barley residue (Cookson et al., 1998).

Surface management (undisturbed on surface, incorporation, burning, or mechanical removal) of wheat or barley stubble also affects decomposition and microbial populations. However, wheat straw incorporated into soil had a higher decomposition rate, mass lost, and substrate-induced respiration than where stubble was burned or removed (Cookson et al., 1998). Residue management did not affect residue decomposition or microbial activity of barley or lupine (Cookson et al., 1998).

The authors hypothesized that the higher lignin:N ratio of wheat caused the response to incorporation.
Decomposition of residue by microorganisms is dependent on the presence of mineral N supply or high N residues (low C:N ratios; Waggett and Wolff, 1998). Decay rate is better correlated with initial N content of residue than with lignin content of low soil N. To adequately meet the needs of the microorganisms involved in decomposition without requiring either added fertilizer or mineral N sources from the soil, residues must contain at least 1.5 to 1.7% N, corresponding to a C:N ratio of ca. 25 or 30 (Parr and Papendick, 1978). The effects of adding inorganic N fertilizer to haeven decomposition of low N residue occur quickly, and after several months the effects of added N on decay cannot be detected (Parr and Papendick, 1978). Knowledge of changes in C and N availability is required to manage crop fertility needs throughout the growing season. Active C and N pools in SOM in agricultural fields vary seasonally, and are dependent on crop rotation, tillage depth, and N fertilization (Frankenberger et al., 1994). Each of these factors affects the type and amount of substrate available for microbial utilization.

**Resource**

**Nutrient Status/Cycling**

Nutrient availability and the role of microorganisms in nutrient immobilization are important concerns in agriculture. Manipulations of food webs to maintain plant nutrition while minimizing N losses are worthwhile goals of sustainable agricultural systems (Altieri, 1999). Bacterial and fungal abundance in the rhizosphere is influenced by the nutrient status of both plant and soil. The percent mycorrhizal cover of roots of Plantago lanceolata was positively correlated with leaf N and P; whereas root colonization by bacteria and other fungi was negatively correlated with plant P (Newman et al., 1981). It might be difficult to separate the effects of soil nutrients on rhizosphere populations from effects involved with increased or altered root exudation of organic compounds. Gram-negative bacteria in monoculture can modify N availability (Wolin and Tilton, 1990); and it has been hypothesized that changes in soil N availability influenced by plant species affect composition of AM fungal communities (Johnson et al., 1992). Microbial population changes occur with added fertilizer and tillage. Nitrogen fertilization increased numbers of fungi and Gram-negative bacteria in rhizosphere of rice (Emmimath and Rangawarki, 1971). Kircher et al. (1993) found that in no-till treatments receiving N fertilizer, fungal populations were higher than under no-till conditions with no fertilizer added. Higher fungal populations in the fertilized treatment were due to increased corn crop growth and higher amounts of residue to serve as substrate for microbial populations, as well as increased root growth and higher amounts of root exudates, which, in turn, increase microbial biomass. Soils that were conventionally tilled, planted with a crimson clover cover crop, and rotated with corn had more actinomycetes, Bacillus spp., and total culturable bacteria than other growe under conventional tillage with fertilizer added, whereas fungi and Gram-negative bacteria were different.

**Plant Growth-Regulating Compounds**

Plant growth-regulating compounds are substances produced by plants and microorganisms in the rhizosphere that enhance seed germination and plant growth (Arshad and Frankenberger, 1998). Soil microorganisms synthesize plant growth regulators, such as auxins, abscisic acid, cytokinins, ethylene, and gibberellins (Frankenberger and Arshad, 1995). These compounds and the organisms that produce them can protect against plant pathogens and stimulate biofertilization (fixation of atmospheric N, or solubilization of nutrients) and plant growth (Figure 10-6). The mechanisms of action are often not readily apparent. Initially, it was thought the N fixation by Azotobacter and Azospirillum was the major reason for plant growth promotion; however, other substances, such as auxins, cytokinins and gibberellins, can stimulate growth (Hausch and Vancura, 1970; Barbier et al., 1993; Jansen et al., 1992). Bacillus and Rhizobium also produce plant growth-stimulating compounds (Frankenberger and Arshad, 1993). Plant growth promotion can also be an indirect
effect of siderophore or antibiotic production, which leads to the reduction in pathogen colonization and infection of the seed and root (Kloepper et al., 1989; Glick 1995).

Microbially derived auxins, ethylene, and other compounds have also been implicated in plant growth inhibition. Plant growth inhibition can be correlated with elevated mobile acetic acid levels produced by rhizobacteria in sugarbeet (Beta vulgaris; Loper and Schroth, 1986), sour cherry (Prunus cerasus L.; Dubickovsky et al., 1993), maize (S.zea and Frankenberger, 1994), lettuce (Lactuca sativa; Barazani and Friedman, 1999), and several weed species (Sarwar and Kremer, 1995). Ethylene, produced by plants, soil fungi, yeasts, and bacteria, can affect plant development from seed germination to senescence (Beyer et al., 1994). Microbial synthesis of ethylene can be affected by the availability of organic substrates and crop residues (Goodlass and Smith, 1978; Lynch and Harper, 1980; Arshad and Frankenberger, 1990).
Amendments

Throughout the history of agriculture, farmers have added amendments to soil to increase crop yield. These amendments have the potential to increase SOM accumulation and plant productivity. Soil amendments can also cause alterations to the soil microbial community. These changes can be quantified by microbial population structure and soil enzyme techniques. Soil microbial populations and activity increase, in general, with manure additions (Altieri, 1999). Manure also increases populations of Collembola and earthworms (Altieri, 1999).

Bio-solids (sewage sludge) is a material that is sometimes applied for agricultural benefit. Long-term application of bio-solids with low and high metal contents added (Cd, Cu, Ni, Zn) showed few differences in the effects on culturable bacteria, but had a dramatic effect on the whole bacterial community, where numbers of the subdivision Proteobacteria increased with high metal concentration (Sanda et al., 2001). Caution must be exercised before applying large amounts of bio-solids to agricultural lands, and bio-solids should be tested for their heavy-metal content before application. Heavy-metal accumulations from bio-solids application can negatively affect microbial communities. Zinc-contaminated agricultural soils (from bio-solids) were tested for microbial diversity and catalytic versatility (Wenderoth and Reber, 1999). Microbial diversity was reduced, and the microbial community experienced a shift to less Gram-positive bacteria and more Gram-negative bacteria compared with the non-contaminated system. The diversity of the Gram-negative bacteria declined under high zinc stress. Stress or heavy-metal contamination can affect overall populations, specific groups, and also the diversity within various groups, as the individual species have different ways to adapt to stress.

Agromicrobials

Numerous agromicrobial products have been touted to increase soil fertility, microbial diversity, and crop yields. Microbial inoculants, such as effective microorganisms (EM) containing yeasts, fungi, bacteria, and actinomycetes increase yields of onion (Allium cepa L.) and pea, and increase cob weights of sweet corn (Daly and Stewart, 1999). The consistency of plant response to these types of products has not yet been demonstrated, and further critical study is needed.

Arbuscular Mycorrhiza (AM)

Mycorrhiza are nonpathogenic fungi that form symbiotic associations with plant roots (Chapter 6). Mycorrhiza are involved in the nutrient cycling process, especially in stressed environments (e.g., P-deficient soils) and can play an active role in SOM accumulation by increasing plant growth by solubilization of nutrients and by producing recalitrus compounds (glomalin). Frogal hyphal threads allow roots to expand the volume of soil that can be explored for nutrients and water that otherwise might be inaccessible to the plant. Mycorrhizal associations enhance nutrient uptake in the rhizosphere and expand the volume of soil the root can explore (Sylvia, 1998). This relationship is especially beneficial under moisture-limiting conditions. Wheat plants inoculated with AM and subjected to water stress at three different times had higher grain yield and biomass than plants that were not inoculated with AM (Ellis et al., 1985). In the Palouse region of eastern Washington, mycorrhizal fungi lessened the severity of water stress in winter wheat (Mohammed et al., 1995). AM species, abundance, and spore distribution are affected by tillage and crop inputs (Douds et al., 1995). Glomus occultum numbers were higher under no-till, whereas other Glomus species were more abundant under conventional tillage in a corn-soybean rotation (Douds et al., 1995).

The interactions of AM and other microorganisms often benefit plants, although the relationships are not always readily evident (Edwards et al., 1998). Presence of AM can enhance relationships with introduced organisms in the rhizosphere of crops. Edwards et al. (1998) found that
biological control agents for *P. fluorescens* did not affect AM function in the rhizospheres of tomato (*Lycopersicon esculentum*) and leek (*Allium porrum*). AM plants had higher shoot weights than non-AM plants, and *P. fluorescens* populations were higher in the presence of AM.

**Biological Control**

Biological control is the use of pathogens, parasites, or other predators to reduce the population or activity of pest organisms (*DeBach*, 1964). Another broader definition includes all forms of intervention, such as genes and gene products, to reduce the impact of pests on crops and beneficial organisms (*Cook*, 1977). The three major strategies for biological control are classical, inundative, and integrated management (*DeBach*, 1964; *TeBeest*, 1991). The classical approach involves the importation of exotics or the use of natural enemies for release, dissemination, and self-perpetuation on target pests. The addition of a virulent strain to suppress pests is the inundative approach. The biocontrol agent is not self-sustaining and must be applied to the target host every season. Integrated management is a broad approach that involves management practices to conserve or enhance native enemies of pests. Biological control is an alternative to pesticides and is part of sustainable agriculture management.

Biological control agents have been investigated for their control of diseases, such as take-all root disease in wheat. The phenomenon known as take-all decline (the reduction in severity of take-all disease) is attributed to naturally occurring strains of fluorescent pseudomonads that produce the antibiotics phenazine and phthiocadino in annually monocropped wheat (*Thosnash* and *Weller*, 1988). Deleterious rhizobacteria have been shown to inhibit such weeds as downy brome (*Bromus tectorum* L.; *Kennedy* et al., 1991), jointed goatgrass (*Aegilops cylindrica* Hat.); *Kennedy* et al., 1992), and velvetleaf (*Abutilon theophrasti* Medik.; *Kremer*, 1987). Several fungal isolates have been investigated for use in weed biological control, such as *Fusarium spp.* against leafy spurge (*Euphorbia* spp.) in the rangelands of the U.S. and Canada (*Caesar* et al., 1990), *Exserohilum rostratum* for grass weeds (*Echinocloa* spp.) in rice production (*Zhang* and *Watts*, 1997), and *Colletotrichum gloeosporioides* f. sp. aeschynomenae to control northern jointvetch (*Aeschynomene virginica*; *Luo* and *TeBeest*, 1999). Several conditions must be met before a biological control agent can be widely used in a crop or rangeland situation. The agent must have adequate shelf life (*Cross* and *Polonienko*, 1996), the ability to be mass-produced (*Oleskevich* et al., 1998), the ability to survive and compete in a field situation, and a simple method of application of the organism and subsequent delivery of the plant-inhibitory compound (*Kremer* and *Kennedy*, 1996).

**Organic/low-input farming**

Organic farming does not allow use of synthetic pesticides of fertilizers and is intended to reduce the detrimental effects of agriculture on soils, animals, food, and the environment. Organic matter and microbial biomass are higher in organic farming systems than in conventional systems (*FileBuch* and *Mäder*, 1997; *Reganold* et al., 1993; *Mueh* and *Goh*, 1997; *Wander* et al., 1994). AM fungi were 30 to 60% higher in root of plants from low-input practices in a long-term field trial that compared organic and conventional systems (*Mader* et al., 2000). In this study, AM was highest in the control soils, lowest in the conventional system, and intermediate in the organic system. The control soils were not fertilized, whereas the pesticide use, disturbance, and high fertility in the conventional systems reduced AM infection. Soils under animal-based organic management had higher levels of the light fraction of particulate SOM than crop-based organic systems or conventional systems (*Wander* et al., 1994). This might be the result of a more biologically active substrate pool due to a lower C:N ratio and higher respiration rate, higher amounts of organic residue added, and less soil disturbance in the animal-based system. Microbial biomass C and dissolved organic C increased as organic inputs increased, and microbial communities as determined by PLFA were
different in organic farms and conventional farms (Landquist et al., 1999). Organic management systems that employed animal manure and legumes for N supply were equally profitable as higher-input conventional systems after 15 years in a study in Pennsylvania (Drinkwater et al., 1998). The organic management systems had lower leaching losses of N and higher levels of soil organic C and N.

Biodynamic agriculture is an organic farming system that uses specific fermented preparations as either field sprays or compost inoculants (Koepf et al., 1976). Soil quality parameters were not different among biodynamic, organic, and conventional management systems, but differed with fertilization level (Penfold et al., 1995). Field-applied biodynamic sprays and compost did not alter soil microbial characteristics compared with conventional practices in a cereal-legume cropping system in the state of Washington (Carpenter-Boggs et al., 2000a). In other studies in the state of Washington, however, biodynamic management resulted in higher microbial biomass, respiration, and SOM than organically managed or conventionally managed systems (Goldstein, 1986). Bio-dynamic preparations for compost development altered compost microbial community and increased compost temperature (Carpenter-Boggs et al., 2000b).

Genetically Modified Organisms (GMOS)

The impact of the addition of genetically modified organisms (GMOS) on soil populations and plant productivity is of interest as more GMOS are introduced into agricultural systems. In a microcosm study of soils from Canada and the U.S., assessment of nontarget effects of two GMOS indicated that there were functional and community differences as long as GMOS persisted in soil; however, effects differed with the GMOS used (Gugliardi et al., 2001). Inoculation of transgenic potatoes with two bacterial biological control agents did not reduce survival of bacterial biological control agents compared with nontransgenic potatoes, nor was the indigenous bacterial community impacted by the introduced bacteria (Loffman et al., 2000). When a GMOS and wild-type Pseudomonas fluorescens was inoculated into the rhizosphere of wheat, both bacterial strains caused shifts in the native microbial populations in the rhizosphere and phylloplane of wheat; however, there were no changes in nonrhizosphere soil and no negative effects on plant health (De Liq et al., 1995). Addition of a genetically modified P. fluorescens in the rhizosphere of pea affected soil enzymes activities and microbial communities (Naseby and Lynch, 1998). Differences are evident with the introduction of some GMOS, but the impact of these differences on soil microbial community, plant productivity, soil quality, and SOM accumulation is case specific, and long-term impacts are not clear.

Disturbance

Agroecosystem function and SOM dynamics are greatly influenced by anthropogenic activities. Soil erosion caused by excessive tillage is the most visual example of humankind’s influence on soil function. Microorganisms are highly sensitive to physical soil disturbance (Elliott and Lynch, 1994), and their population dynamics can serve as early warning indicators of changes in soil quality. Fluxes in microbial diversity and functional diversity can contribute greatly to the understanding of soil quality, and the development of sustainable agroecosystems (Thomas and Kevan, 1993; di Castri and Soanes, 1990; Hawksworth, 1991a). Soil organisms are useful in classifying disturbed or contaminated systems, because diversity can be affected by minute changes in the ecosystem. Severe disturbances, such as those caused by heavy tillage with a moldboard plow (which completely inverts the surface soil), overgrazing, and pollutants, can reduce aboveground plant diversity and growth. This reduction in plant biomass and lack of a varied carbon source decreased microbial growth and functioning (Christensen, 1989; Zak, 1992).
100 Years of Dryland Farming in the Inland Pacific Northwest, U.S.A.

Wheat–fallow, in which only one crop is grown every 2 years, is the dominant cropping system in the low-precipitation dryland cropping region of the inland Pacific Northwest. Early settlers grew a wheat crop every year after they first broke the land out of native bunch grass and sagebrush in the 1880s. Grain yields were frequently low; however, it soon became apparent that farmers could better stabilize yields by having a fallow year to store soil moisture between wheat crops. During the fallow year, essential nutrients, mainly N, were released through mineralization of SOM. The native grass prairie provided a reserve of readily decomposable SOM that supplied nitrogen for crop use for many years. A rapid reduction in SOM content occurred when prairie soils were brought under cultivation, especially when alternated with fallow. In some soils, more than 25% of the organic matter was lost in the first 20 years of farming. In undisturbed native soil, SOM in the top 10 cm of soil is 4% at Pullman, WA (350 mm precipitation), 3% at Pendleton, OR (410 mm precipitation), and 1.5% at Lind, WA (240 mm precipitation). Distinct decreases in SOM have been observed on farmland compared to undisturbed native soil in all three areas. Organic matter content in the top 10 cm of cropland soil at Lind, for example, is at present less than 1%. Data are available on the long-term fate of SOM in a continuous winter wheat–fallow rotation from an ongoing 70-year-old study at Pendleton, OR (Rasmussen et al., 1989; Figure 10.5). Since 1931, SOM has continuously declined under all residue management methods except when 22 mt (fresh weight) ha⁻¹ of cattle manure was applied every other year. SOM decline has been highest when stubble was burned in the fall and when no nitrogen was applied. Maintaining an adequate nitrogen supply and returning all residue to the soil has reduced, but not arrested, SOM decline (Figure 10.5).

Tillage

Up to 50% of the SOM in some soils has been lost after years of intensive tillage, clearing vegetation, and draining wetlands for farming (Caubelardella and Elliott, 1993). Carbon is sequestered in soils through “humification of organic residues, building of organonomineral complexes to form aggregates, positioning of SOM beneath the tillage layer, use of deep-rooting crops, and calcification” (Bruce et al., 1999). Emissions of CO₂ from agricultural soils can be reduced by minimizing or eliminating tillage and by growing perennial crops (Lal et al., 1999). In the U.S., many farmers have 10-year contracts under the Conservation Reserve Program (CRP) to grow perennial grasses and shrubs for environmental and soil-conserving benefits (USDA-FSA, 2000). Improved farming techniques, higher productivity from farmland, and government programs that pay landowners to plant permanent vegetation on highly erodable lands have combined to increase levels of SOM in many soils (Lal et al., 1999). No-till farming results in less CO₂ released to the atmosphere than do intensive tillage (moldboard plowing) and minimum tillage (disk harrowing; Reicosky and Lindstrom, 1993). Incorporated straw emits more CO₂ than does surface straw or soil with no straw applied (Curtin et al., 1998). Gale and Cambardella (2000) compared contributions of root and surface residues to soil organic C in no-till soils, and found that the greatest increases were due to maintenance of root-derived C, illustrating the importance of leaving root biomass intact (no disturbance) for maximum SOM accumulation.

Conservation-tillage systems maintain at least 30% of the crop residue on the soil surface (Stroox et al., 1989) and help prevent soil erosion by wind and water (Figure 10.6; Papendick, 1996). Positive attributes of retaining crop residue on the soil surface are improved soil quality (Karlen et al., 1994; Dalal et al., 1991; Doran et al., 1996) through increased biological activity, leading to improved soil aggregation and more SOM content (Elliott and Papendick, 1986).
FIGURE 10.5 Soil organic matter decline in a winter wheat–summer fallow rotation at Pendleton, OR. Rapid decline in SOM occurred after the onset of dryland farming in the early 1880s. A long-term experiment was initiated in 1931 to test effects of soil amendments (cattle manure and pea vines), N fertility (0 and 45 kg ha⁻¹), and burning of residue. SOM steadily declined in all treatments except on addition of 22 mt manure ha⁻¹ every other year with no burning of residue. (Modified from Rasmussen, P. E., H.P. Collins, and R.W. Smiley, 1989. Long-term Management Effects on Soil Productivity and Crop Yields in Semi-arid Regions of Eastern Oregon. Oregon State University Bulletin 675. Corvallis, OR.)

FIGURE 10.6 No-till planting in the dryland wheat production region of the inland Pacific Northwest. No-till preserves plant residue on the soil surface for erosion control, promotes microbial populations, and provides other environmental benefits.

Conservation tillage, increased cropping intensity (e.g., rotation in fallow), crop rotation, and use of cover crops improve soil quality (Karlen et al., 1992). Crop residues on the soil surface, however, can negatively affect crop yield by impairing seedling emergence, serving as hosts for pathogens, or nutrient immobilization (Elliot and PapENDICK, 1986). Even distribution of crop residue at harvest and selection of a no-till planter for specific soil and residue conditions reduce the possibility of yield loss.
No-till increases microbial biomass in surface soils (0 to 15 cm; Drijber et al., 2003), increases the ratio of fungi to bacteria, and provides for a more diverse population of residue decomposers and a slower release of nutrients than does conventional tillage (Alkire, 1999). The changes in the physical and chemical properties of soil resulting from tillage greatly alter the matrix supporting growth of the microbial population. Within a given soil, there is considerable variation in the composition of the microbial community and diversity with depth in the profile. In agricultural systems, microbial activities differ drastically with depth, with the highest microbial activity occurring near the surface in no-till, and more evenly distributed activity throughout the plow layer of silled soil (Doran, 1980).

Composition of the microbial community influences the rate of residue decomposition and nutrient cycling in both no-till and tillage-based systems (Beare et al., 1993). Fungi dominate decomposition in a no-till system, whereas the bacterial component is responsible for a greater portion of the decomposition of residue with tillage. In a study of the diversity of native prairie and cultivated soils, diversity indices were higher with tillage than with grassland (Kennedy and Smith, 1995). With the substrate exposed by tillage, more surface area was available for colonization and more activity occurred. Increase in diversity seen early on with disturbance indicates a change in the microbial community to one that exhibits a greater range of substrate utilization and stress resistance. When comparing microbial numbers among burned, tilled, or no-tilled fields of double-cropped wheat and soybean in Georgia, there were no change in total bacteria or nitrifiers with burning or tillage. Ploes that were not burned or tilled initially had higher numbers of alget, actinomycetes, and fungi; however, there were no treatment differences later in the growing season (Harris et al., 1995). Preemergent herbicides had no effect on microbial numbers in that study. Buriedorghum residue under conventional tillage contained more fungal hyphae and CFP than surface residue in no-till soil; however, in bulk soil there were no differences in fungal CFP between treatments, and higher numbers of fungal hyphae were found in no-till soil (Beare et al., 1993). These studies illustrate the alteration of the make-up of microbial communities and possibly the diversity of basic microbial groups with disturbance.

Studies have shown varying results with regards to N immobilization in reduced-tillage systems. When farmers first convert to minimum- or no-till cropping, they often encounter lower N availability for the first several years because of reduced mineralizable N. SOM (and N) accumulates under no-till, however, and a new equilibrium is established in which mineralized N and microbial biomass C are higher than under intensive tillage (Simard et al., 1994). Higher tillage intensity under conventional tillage decreased the amount of N mineralized per unit of biomass C, which could lead to a decline in SOM quality. In a long-term (11-year) study in Canada, soil C and mineralizable N were highest with no till compared with conventional tillage, regardless of cropping sequence or cropping frequency (Campbell et al., 1996). Conversely, decomposing surface residues in some no-till systems can immobilize enough N to cause N deficiency in succeeding crops (Knowles et al., 1993), and increased tillage intensity reduces the potential for N immobilization (Fujiit and Schimel, 1999). In these instances, cropping intensity and rotation (Kolberg et al., 1999; Knowles et al., 1995) and fertilizer placement (Knowles et al., 1993) must be managed to ensure success of no-till farming systems. Carbon gain in the soil is a function of both residue input and clay content of the soil. After 20 years of wheat and barley residue maintenance by using no-till and high N fertilization in Australia, organic C, total N, and microbial biomass were higher and pH was lower at the soil surface than under conventional tillage in which residue was burned and lower levels of N were applied (Dudal et al., 1991).

SOM increases when crop residue is retained on the soil surface (as in no-till systems), when erosion is reduced, and crops are adequately fertilized (Campbell and Zentner, 1993). In the Canadian prairies, potential exists for C sequestration under a long-term (12- to 15-year) no-till continuous wheat farming system compared with a system using conventional tillage wheat-fallow
because of lower CO₂ flux from the soil as more organic matter accumulation with reduced residue disturbance and continuous cropping (Curtis et al. 2000). Four years of no-till increased SOM in the top 2.5 cm of soil compared with conventional tillage in a Mississippi study with grain sorghum-corn, cotton and soybean-wheat rotations (Rhodon, 2000). No-till in the sorghum-corn rotation showed the maximum accumulations of soil organic content, especially in the cusp 2 cm of soil, compared with conventional tillage and rotations that included soybean in the Great Plains (McCullister and Chen. 2000).

Although numerous studies indicate higher SOM under no till vs. conventional tillage, some studies show little or no difference in SOM between the two systems, especially in low-precipitation regions where residue production from crops is minimal (A.C. Kennelly, unpublished data). Macromineralic SOM (>50 um) and microbial biomass-C can be good indicators of change in residue management; however, effects of tillage might be limited to vertical distribution without influencing SOM turnover (Angers et al. 1995). Needelman et al. (1999) found that no-till fields in a corn-soybean rotation in Illinois had higher SOM in the top 0 to 5 cm of soil than that in conventionally tilled fields, but SOM was not different between the two tillage treatments when the entire sampling depth (0 to 30 cm) was considered. SOM levels did not differ from conventional tillage levels in the top 15 cm of soil after 30 years of a no-till wheat-sorghum-fallow rotation in Kansas (Thompson and Whitney, 2000). Maintaining or increasing SOM is critical to crop production to improve soil water-holding capacity and aeration, provide nutrients for plants and microbes, and maintain soil physical properties such as friability and low bulk density. The amounts of SOM that accumulate in different systems vary greatly with geographic location (soil type, precipitation, and climate), length of time for which a particular management scheme is used, tillage intensity, and crop residue inputs.

Population and diversity of geomic patterns of the N₂-fixing bacteria Bradyrhizobium increased with no-till compared to conventional tillage in southern Brazil (Ferreiras et al. 2000), even though the field was last inoculated 15 years before the study. Along with no-till, crop rotations containing soybean increased populations of Bradyrhizobium. Treatments that did not include soybeans in rotation for 17 years and were in conventional tillage contained the least amount of Bradyrhizobium.

Wardie et al. (1999) studied three methods of controlling weeds (mulching, herbicides, tillage) and found that mulching (adding residue) increased soil C, microbial biomass, and activity in surface soils (1 to 10 cm) over the course of a 7-year study in New Zealand; however, some immobilization of N might have occurred late in the study. Herbicide application did not adversely affect microbial biomass and activity. Where less weed biomass was present, microbial respiration was reduced, probably because of more decomposition of weeds than crop plants. Tillage for weed control was not detrimental to substrate-induced respiration, CO₂-C released from chloroform fumigation, or soil organic C in the study. The authors emphasized the need for long-term studies, as many of their results were not apparent until after 6 years.

Although most of the effects of reduced tillage are positive, there are some instances wherein more physical soil disturbance is advantageous. Direct seeding wheat into cereal or grass residue increases the risk of infection by pathogens causing the diseases take-all, Rhizoctonia (Rhizoctonia solani) root rot, Cephalosporium (Cephalosporium gramineum) stripe, and Pythium root rot (Pythium spp.; Cook and Haglund, 1991). Crop residue can serve as a host for the pathogens (Cook, 1986). Crop rotation and tillage are suggested to alleviate disease pressure in wheat (Cook and Haglund, 1991). Abawi and Wilmer (2000) cite numerous examples in which yield of bean was increased because of less disease with intensive tillage compared with reduced tillage or no-till. The increase in yield with tillage was attributed to reduced compaction, improved drainage, and higher soil temperature, which led to improved bean root competition against pathogens.
Grazing

Careful management of grazing lands is needed to protect soils from the negative effects of overgrazing and to maintain benefits of permanent plant cover. Livestock grazing is thought to be less damaging to soil quality than is conventional crop management; however, soil quality can be impacted by compaction and continual removal of plant cover (Southorn and Cattle, 2000). Cattle grazing can also affect the biomass and biodiversity of plants by causing patches that differ in size and plant species (Cad and Britzuela, 1998). Also, because of overgrazing, species composition in grazing lands can shift from perennial species to annual grasses (DiTomasso, 2000). This land then becomes more susceptible to invasion by broad-leaved weed species, which degrades soil productivity (DiTomasso, 2000). Although many weed species have deep taproots, they produce less aboveground biomass than do most crop plants, often leaving surface soil vulnerable to erosion. A high infestation of spotted knapweed (Centauraea maculosa) reduced water infiltration rates (DiTomasso, 2000). Additionally, overgrazing and recolonization with weed species led to less soil moisture available to grass species and less contribution to SOM than did fibrous root systems of grass species (DiTomasso, 2000). Abell and Bucher (1999) showed the negative effects of overgrazing of native rangelands in Argentina, where the overgrazed site had the lowest SOM and microbial activity. In a comparison of restored, partially restored, and overgrazed rangelands, they found that soil water, SOM, N content, and microbial activity were highest at the restored site. In another study on the soil quality of grasslands in New Mexico, Liu et al. (2000) found no negative effects on microbial diversity (substrate utilization) or microbial activity (enzyme analysis) from intense grazing; however, burning reduced microbial diversity.

Integrated systems combining crop production and livestock production with perennials are suggested as a means to improve soil quality and combat the decline in organic C of soil from the Great Plains of the U.S. after decades of cultivation (Krahl and Schuman, 1996). Well-managed grasslands used for livestock grazing adjacent to streams can protect or enhance soil quality by stabilizing stream banks and reducing erosion (Lyons et al. 2000). These managed riparian areas can reduce the impact of livestock grazing and help restore degraded stream banks.

STRATEGIES FOR MANAGING MICROORGANISMS

Although the technology for managing microorganisms for sustainable agricultural production systems has not yet been developed, several strategies have been used for centuries to optimize soil life (see Chapter 2 for a discussion on SOM management). First, management practices that increase SOM should be used, especially in SOM-depleted systems. Organic matter is responsible for providing substrate for microorganisms, but also improves microbial habitat. Organic matter in various forms of decay improves soil physical properties, increases water-holding capacity and nutrient availability, and acts as a cementing agent for holding soil particles together. SOM can be maintained or increased by incorporating crop residues, crop rotation, cover crops, permanent plantings, maintaining soil fertility, and adding animal manures or biosolids. Addition of plentiful amounts of organic residues helps ensure a productive soil and stimulates plant growth by providing food for microorganisms. The movement toward sustainable farming systems with diverse, healthy soil microbial communities that closely imitate the processes of native, undisturbed systems, can be realized by using these practices or adopting a combination of several practices.

A second strategy is to ensure a diverse plant community through crop rotation or grazing management. Minimizing fallow or increasing root growth in soil will provide substrate additions and adequate nutrition for a healthy soil and large, diverse populations of microorganisms. Tillage and burning of crop residues often negatively and dramatically affect the chemical and physical properties of soil, which alter growth of microorganisms and processes for which they are responsible. Minimum tillage or no till helps prevent erosion of valuable topsoil.
Options for Farmers

The following management principles will help maximize soil quality in low-precipitation areas, such as in the inland Pacific Northwest:

- Minimize tillage to the degree feasible to leave as much residue as possible on or near the soil surface.
- Maintain adequate nitrogen inputs. Because of the linkage between soil N and organic matter, adequate (but not excessive) nitrogen inputs are a requisite for optimum crop growth and residue return.
- Minimize the use of summer fallow, if possible. Consider recropping to spring wheat or barley after wet winters. Use a no-till drill, if feasible, to plant seed and fertilizer in one pass through the standing residue of the previous crop.
- Emphasize wind and water erosion control, because any loss of topsoil increases loss of SOM.

Applying large quantities of organic materials, such as cattle manure, can increase SOM. This, however, is not a realistic option for most farmers because of the large quantities of manure required, as the size of the average farm exceeds 1000 ha.

CONCLUSIONS

Microorganisms are responsible for a multitude of soil processes, such as SOM dynamics, nutrient cycling, and changes in soil structure. In agroecosystems, microorganisms can affect all levels within the ecosystem through functions such as N and C cycling, plant growth promotion and inhibition, and natural biological control. Microorganisms have more diversity than does any other group of organisms on earth, but our knowledge of these organisms is still limited. We need to increase our understanding of microbial communities and their function in agroecosystems. Several strategies have been suggested to optimize soil life. The most critical to sustainability is to use management practices that increase SOM, reduce disturbance, and maintain a diverse plant community. There is a wealth of genetic potential in the soil, but we do not presently have the means or understanding to use the full potential of the earth’s oldest inhabitants. With a better understanding of soil ecology and changes in soil biota with management, best management practices can be developed to conserve and enhance SOM, soil quality, and crop productivity for sustainable agricultural systems.

References


Thompson, C. A., and D. A. Whitney. 2000. Effects of 30 years of cropping and tillage systems on soil 


Turkington, R., F. B. Holt, C. P. Chaawaya, and J. D. Thompson. 1988. The influence of microorganisms, 

Turkington, R., and E. Klein. 1991. Competitive outcome among four pasture species in sterilized and 

USDA-FSAA. 2000. The conservation reserve program 28th signup. U.S. Department of Agriculture, Farm 
Service Agency, Washington, D.C.


Vascara, V. 1967. Root exudates of plants. III. Effects of temperature and “cold shock” on the exudation 

In Sylvia, D. M., J. J. Fulmshin, R. G. Hartel, and D. A. Zauberer (Eds.). *Principles and Applications 
of Soil Microbiology.* Prentice-Hall, New York.


biomass dynamics, activity, and plant litter decomposition to agricultural intensification over a seven-


European Communities, Luxembourg.


Wojciech, F., and A. H. C. van Bruggen. 1995. Bacterial density, composition and diversity in organically and 
conventionally managed rhizosphere soil in relation to suppression of corky root of tomatoes. *Appl. 


Wright, S. F., and A. Upadhyaya. 1998. A survey of soils for aggregate stability and glomalin, a 


Yacki, S., and M. Loreau. 1999. Biodiversity and ecosystem productivity in a fluctuating environment. The 


