Arbuscular mycorrhizal fungi (AMF) are associated with ca. 80% of living terrestrial plants (Pirozynski, 1981). They play a multifunctional role in ecosystems such as plant nutrient uptake facilitation and the improvement of plant-soil water relations for a wide range of species (Newsham et al., 1995a), affecting both their productivity and relative abundance, and ultimately influencing plant species diversity (Grime et al., 1987; Gange et al., 1990, 1993). Mycorrhizae are capable of influencing plant community structure within the ecosystems (Wilson and Hartnett, 1997, 1998; Van der Heijden et al., 1998). Moreover, mycorrhizal fungi have a high physiological diversity (Allen et al., 1995) due to a high genetic diversity (Sanders et al., 1996); consequently, mycorrhizal fungi have a high functional diversity within ecosystems (e.g. accumulation of toxic materials, decomposition of organic matter, protection from root pathogens, pro-
duction of environmental biochemicals -antibiotics and enzymes-) (Miller, 1995; Zak and Visser, 1996).

In terms of nutrition, AMF require carbon from a host plant to survive. They can obtain carbon from the specialists living on limited plant species in particular habitats with long survival times (Warcup, 1981; Allen, 1991) to generalists found in swards or associated with early and later successional plant groups (Fitter, 1990; Titus and del Moral, 1998a, 1998b; Turner and Friese, 1998; Hartnett and Wilson, 1999).

Arbuscular mycorrhizal fungi (AMF) form a wide range of asexual spores or colonization structures (Zak and Visser, 1996), which allow them an extraordinary capacity for growing, dispersing and surviving (Allen, 1991); however, AMF life history is not yet well understood. Therefore, the aim of this paper is to review and assess scientific literature relevant to arbuscular mycorrhizal fungi (AMF) population dynamics: reproduction and dispersal mechanisms, and establishment and distribution in natural ecosystems. Mycorrhiza are important in both plant establishment and ecosystem restoration, and recently their importance in maintaining plant diversity and ecosystem functioning has been recognized.

**Mycorrhizal fungi life cycle**

As in many other fungi, the generalized life cycle of the mycorrhizal fungi (MF) (figure 1) responds in a highly plastic adaptation to its surrounding environment. An individual spore (sexual or asexual in the case of AMF), whether surviving a stress period or dispersing to a new habitat, will germinate producing a germ tube, which is a slender hypha initially having one growing tip. The hypha will expand and branch as long as conditions for growth are adequate to form a mycelium or hyphal network. This mycelium can be either haploid or diploid with each cell containing either one or more nuclei; it continues to expand until reaching an unfavorable habitat or contacting another compatible mycelium. Following this contact, anastomosis may occur between hyphae, followed by exchange of genetic material. The genetic material may either remain intact as separate nuclei (if the nuclei are the same, it is referred to as a homokaryon cell and if the nuclei are different, as a heterokaryon cell), or fuse to become diploid. Rarely, the fungus continues to grow with a 2n nucleus. More often, it undergoes sexual recombination to form new, recombined haploid nuclei. All four possibilities occur in mycorrhizal fungi, depending on the classification and genetic potential of each species (Allen, 1991; Zak and Visser, 1996).

Arbuscular mycorrhizal fungi (AMF) are thought to reproduce exclusively by asexual clamydospores and hyphal fragmentation, while ectomycorrhizal fungi (EMF) produce sexual spores (Allen et al., 1995). Due to hyphal fusion, gene turnover, and/or molecular drive, AMF individual clamydospores and hyphae may have hundreds to thousands of genetically different nuclei (heterokaryon), which can ex-
Plain the high genetic diversity this kind of fungi possesses without any exchange of genetic material among individuals (Sanders et al., 1996). Consequently, a high functional diversity is found in AMF (Allen et al., 1995; Wildman, 1995; Streitwolf-Engel et al., 1997; Van der Heijden et al., 1998; Hartnett and Wilson, 1999).

A great part of the ecological role of an organism, and the strength and variety of population and community interactions are determined by growth form (Zak and Visser, 1996). Fungal mycelium presents a modular growth form, exhibiting a high plasticity in shape, size and reproductive potential, and the same genetic individual (genet) can simultaneously be exposed to different environments and selection pressures (Andrews, 1992; cited in Zak & Visser, 1996). Modular growth provides fungi several specific attributes: indeterminate and iterative growth, maximum reproductive potential early in adult life, and an indefinite life span; fungi senescence and death may thus only be local, environmental effects on development are major, resources are obtained through growth across nutrient patches, and local disturbances are relatively unimportant (Zak and Visser, 1996). Fitness in fungi is therefore most readily defined in terms of population survival through time without enumerating offspring. This survival can take one of three forms: (1) the ability to retain a patch of soil through time, (2) the ability to tolerate stress conditions and reestablishment following the stress period, and (3) the ability to migrate and establish new colonies in new habitats (Allen, 1991).

It has been shown that MF can retain a patch of soil for long time periods; for example, some basidiomycetous mycorrhizal fungi form fairy rings that can last up to hundreds of years (Allen, 1991). Christensen (1989) registered the occurrence of many tropical fungi in temperate grassland soils, whose ancestors were originally deposited during the Pleistocene. In theory, mycelium and spores of many fungi have the capacity to survive for long time periods or maybe indefinitely (Allen, 1991).

Examples of AMF inoculum survivorship to stress conditions and their reestablishment through time are shown by several studies carried out to determine the role played by AMF and other factors (e.g. soil pH and texture) upon plant re-colonization after a disturbance process such as an early plant community succession (Nicolson, 1960; Allen and Allen, 1980; Gange et al., 1990, 1993) and Mount St. Helens eruption (Allen et al., 1984; Allen, 1987; Carpenter et al. 1987; Titus and del Moral, 1998a, 1998b).

**Dispersal of mycorrhizal fungi**

**Disturbance.** In order for mycorrhizal fungi to expand their range (figure 2), there must be a source of new habitats. Disturbances may represent the major source of these new habi-
tats available to mycorrhizal fungi. Small or gradual disturbances such as those created by animals (e.g. soil removal - mounds- while creating their burrows) still retain some existing inoculum, allowing those fungi a better chance for establishment based solely on inoculum density. Severe disturbances, natural (e.g. landslides, fire or volcano eruptions) or man-induced (e.g. mineral extraction or agriculture), create open habitats susceptible of invasion by new plants and mycorrhizal fungi (Allen, 1991). These disturbances may, or may not, lead to marked changes in the mycorrhizal formation (e.g. AM fungal species, AM fungal inoculum density, and AM fungal percentage of root colonization) (Abbot and Robson, 1991).

A well documented example of severe disturbance such as the eruption of Mount St. Helens, USA, in 1980, gave way to the study of several aspects concerning the mycorrhizal association: the process of AM fungal inoculum re-establishment (Allen et al., 1984; Allen, 1987), the role of the mycelia as part of the substrate for the colonization by non-vascular plants (Carpenter et al., 1987), and the relation between AMF presence/absence and the plant species composition in primary succession (Titus and del Moral, 1998a, 1998b). These studies concluded that in an early successional ecosystem, plants initially establish in distinct patches and that most of these plants are mycorrhizal. Dispersal, survival, and establishment (mycorrhizal colonization) of AM fungal association are thus important to the long term growth and survival of many plants (Allen and MacMahon, 1988), and of the AMF-plant association (Allen, 1991). Arbuscular mycorrhizal fungi (AMF) are thus one of the interacting factors influencing the competitive interaction among plant species that leads to the change of the plant species over successional time (Titus and del Moral, 1998a, 1998b). Allen et al. (1984) considered that the reconstruction of a given ecosystem depends, in part, on the ability of a particular fungus, such as the AMF or the EMF, to reach, survive at, and increase on a given site.

**Spore density and dormancy.** Spores of AMF are abundant in both nature and agricultural soils. However, results on spore density are related to the different methods employed (e.g. wet sieving and decanting method -Gerdemann and Nicolson (1963)-, including or not centrifugation, or different soil samples weight: 50 g or 100 g of dry soil). For instance, spore density ranged from 0 to 137 per 50 g of soil (Abbot and Robson, 1977), and from 2 to 1952 spores per 100 g of dry soil (Hayman and Stovold, 1979) in different Australian soil samples, and in a sand dune in USA, total spore density ranged from 0 to 677 per 100 g of sand (Sylvia, 1986). In the three studies, most of the samples contained at least two spore types. Hayman and Stovold (1979) demonstrated that there was no correlation between the size of an AM fungal spore population and the mycorrhizal infectivity, and that spore numbers were frequently lower in natural than in adjacent cultivated soils. In addition, Hayman (1982) showed that AM fungal spores can survive at least for one year in the field.

Furthermore, there are seasonal fluctuations in spore density, which varies greatly in relation to AM fungal and plant phenology; sometimes, both associated are phenologically synchronized. Sylvia (1986) and Gemma and Koske (1988) showed that both flowering and sporulation might occur almost simultaneously, and that newly-formed spores were dormant, apparently serving to maintain a high population of inoculum in the soil in the non-growing season of the hosts; for instance, dormancy in Gigaspora gigantea spores had two sequential limitations that inhibit their immediate germination: 2 to 9 week long period of dormancy followed by a temperature-imposed exogenous dormancy when temperature is below 15°C (Gemma and Koske, 1988).

**Plant root to root contact and arbuscular mycorrhizal fungi (AMF) dispersal.** Read et al. (1976) suggested that most AMF spread via root to root contact. Powell (1979) estimated that AMF could spread at rates of 6 to 43 cm/year by root expansion in a greenhouse experiment; while Warner and Mosse (1980) found that AM fungal hyphae expanded a few dm/year (ca. 24 cm) within sterile plots in the field. Natural spread of mycorrhizal fungi through plant root to root contact within the soil is thus slow (Powell, 1979) and greatly influenced by plant species presence (Warner and Mosse, 1980; Miller et al., 1989); most of the hyphae are located within 0.1 to 0.2 cm of the root surface (“nutrient depletion zone”) having an effect similar to the one exercised by the root hairs (Owusu-Bennoah and Wild, 1979).

Ecologically, mycorrhizal growth leads to interconnections between plants, which favors the nutrient transfer among them. It is likely to be of great significance for survival of seedlings in circumstances where both, light and nutrients, are already extensively exploited by established plants. Nutrients move into the fungus from the source root and are then translocated through interconnecting hyphae to sink plants without entering into the soil solution (Whittingham and Read, 1982).

**Mycorrhizal fungi (MF) dispersal by animals and erosion agents.** Arbuscular mycorrhizal fungi (AMF) and some EMF have been observed to spread via small animals such as ants, grasshoppers, termites, wasps, birds, and rodents (MclVeen and Cole Jr., 1976; Maser et al., 1978; Allen et al., 1984; Allen, 1987; Warner et al., 1987; Koide and Mooney, 1987; Allen and MacMahon, 1988; Zoberi and Grace, 1990; Herrera et al., 1997), big mammals as ungulates, and bears (Cázares and Trappe, 1994), and via some erosion agents such as wind and water (Allen et al., 1984; Carpenter et al., 1987; Trappe, 1988).

Earthworms, termites and ants turn over large quantities of soil (ca. 2 to 5 cm of soil to the surface in a decade), playing an important role in both horizontal and vertical distribu-
tion of AMF and EMF spores through dispersal of their casting material and soil removal (McIlveen and Cole Jr., 1976; Zoberi and Grace, 1990). Mycorrhizal inoculum has also been observed in gopher mounds in both serpentine grasslands (Koide and Mooney, 1987), and forest clearings (Allen et al., 1984; Allen and MacMahon, 1988), while communities of microfungi have been reported in rodent food stores (Herrera et al., 1997). Birds and mud dauber wasps are important in dispersal of mycorrhizal fungi spores in view of the distance and speed of dispersal, and not by the amount of inoculum they can disperse (McIlveen and Cole Jr., 1976).

On the other hand, most EMF, which fruit above ground, can also be dispersed across long distances if their spores are entrained in the upper air flows above the canopy; for example, the wind can move spores up to 2 km (Warner et al., 1987). Hypogeous fungi, as well as many forest floor mycorrhizal fungi, which produce brightly colored fruiting bodies, are eaten by squirrels and other small (Trappe and Maser, 1976; Maser et al., 1978) and big mammals (Cázares and Trappe, 1994), dispersing the spores through fecal pellets.

Thus, plant community re-colonization and/or permanence also depends upon dispersal of viable MF propagules, fungal inoculum establishment, presence of “adequate” plant propagules, and appropriate environmental conditions.

Factors influencing arbuscular mycorrhizal fungi (AMF) distribution

Arbuscular mycorrhizal fungi (AMF) are believed to be among the most abundant fungi in soil (Gerdemann and Nicolson, 1963). They are abundant in grasslands, savannas, scrub and open woodlands, rain forests, deserts and sand dunes; on a global scale, they are virtually ubiquitous (Hayman, 1982), and they differ greatly in their effects on plants, ranging from mutualistic to neutral to parasitic (Hayman, 1982; Fitter, 1991; Johnson et al., 1997). Though AMF are obligated root symbionts, very little about their population dynamics is known; however, some insights have been gained by studying the natural distributions of their spores (Johnson et al., 1992; Hartnett and Wilson, 1999).

Distribution, activity and survival of AMF have been shown to be influenced by several factors (table 1) such as soil fertility, soil moisture, soil compaction, soil depth, soil water saturation, pH, topography, burning frequency, temperature, light intensity, altitude, latitude, plant susceptibility-phenology, AMF phenological variations and environmental disturbance, as well as by the physical movement of the water within the soil, and by the earthworms and the soil microfauna activities (Hayman, 1982).

Distribution of AMF changes across a soil moisture-nutrient gradient (Anderson et al., 1984), soil pH can influence AMF infection, sporulation, and spore germination (Green et al., 1976; Porter et al., 1987a, b), and low levels of phosphorus within the soil can regulate AMF spore germination and root colonization (Hettick and Wilson, 1989).

Soil compaction can deplete AM fungal growth. Nadian et al. (1997, 1998) considered that the absence of any observable mycorrhizal growth response in highly compacted soil was attributed to the significant decrease in the oxygen (O₂) content of the soil atmosphere, change in soil pore size distribution, and probably to ethylene production from impeded roots or to any combination of these factors.

The analysis of the influence of soil depth on mycorrhizal performance shows certain controversy. Jakobsen and Nielsen (1983) found that the proportion of root length infected decreased in relation to soil depth (> 40 cm) and that the susceptibility to infection was independent of host species, while Zajicek and Hettick (1986) observed that the degree of colonization varied with plant species and soil depth, and suggested that the deep-rooted growth habit of certain plants (e.g. forbs) probably is a survival mechanism by which competition for water and nutrients with shallower-rooted and fast-growing plants (e.g. grasses) is avoided, since soil fertility generally decreases as soil depth increases. Another example is the woody legume Prosopis glandulosa that develops functional root symbiotic associations with nitrogen (N)fixing bacteria and AMF at depths greater than 4 m, and population densities of both symbiotic organisms are substantially greater at depth than near the surface (Virginia et al., 1986).

In a marsh, Cooke et al. (1993) determined that the presence of infected roots at 42.5 cm depth and the lack of detectable oxygen at this depth suggested that sufficient oxygen transport occurs in host plants to sustain the growth of the AMF in the roots. Miller (2000) found that AMF colonization was strongly negatively correlated with water depth and concluded that flooding was partially but not totally inhibitory to AM fungal colonization, ratifying Søndergaard and Laegaard (1977), Koske et al. (1985), and Cooke et al. (1993) previous conclusions.

Other factors, such as topography and burning frequency, affect gradients of variation in AMF species. Nevertheless, much of the AMF species compositional gradients may be also an indirect consequence of topographic and fire effects on plant species distributions (Gibson and Hettick, 1988).

Temperature can also influence AM fungal species richness distribution along a latitudinal (north-south) gradient, although temperature effects on the AMF communities may be separated into two components: a direct effect on the fungi and an indirect effect mediated through the host plant (Koske, 1987). Allen et al. (1995) found that AMF exhibited a latitudinal gradient of occurrence. There were three distinct communities of fungi within two sites indicating that AMF formed their own specific community patterns regardless of the host plant.

Phenological variations occur in AM fungal populations, as well as in saprophytic and pathogenic fungi populations.
Effect AM fungal species richness was positively correlated with percentage soil organic matter and negatively correlated with Ca, Mg, and P content of soil, while total spore number was positively correlated with N and organic matter content and negatively correlated with Ca, Mg and P content of soil.

Total P uptake was significantly greater in mycorrhizal plants than in non-mycorrhizal ones; the response was smaller as soil compaction was increased. Soil compaction to a bulk density 1.6 Mg m\(^{-3}\) had no effect on the percentage of root length colonized, but total root length colonized decreased as soil compaction was increased. Soil compaction, which increased bulk density from 1.20 to 1.75 Mg m\(^{-3}\), reduced O\(_2\) content of the soil atmosphere from 0.16 to 0.05 m\(^3\) m\(^{-3}\).

The proportion of root length infected decreased markedly below 40 cm soil depth; root density varied greatly between crops, whereas the absolute length of infected roots was similar. The degree of colonization varied with plant species and soil depth; spores of *Glomus fasciculatum* were found to a depth of 220 cm. *Prosopis glandulosa* developed functional root symbiotic associations with N\(_2\)-fixing nodules and AMF at depths greater than 4 m in moist soil above a seasonally stable water table.

Roots were colonized by AMF to a depth of 42.5 cm, but none arbuscule was observed below 37.5 cm. AM fungal colonization was strongly, negatively correlated with water depth; colonization was lowest in plots that were consistently wet but rose as some plots underwent seasonal drying; soils that were wet for > 1 year had the same ability to form mycorrhizas in bait plants as those that had remained dry.

AMF spores germination was influenced by pH: *Glomus mosseae* germinated best at pH 7, *Gigaspora coralloidea* at pH 5, and *G. heterogama* at pH 6. pH influenced the ability of infection, sporulation, and spore germination: *Acaulospora leavis* was distributed in soil samples ranging from pH 4.5-4.9, two species of *Gigaspora* from pH 4.5-6.4 and three different strains of *Glomus* at pH 5.5-8.4.

Even closely related hosts (five grasses) may cause divergence in AM fungal communities on initially identical soils; fungal communities in the sandy end of the soil gradient diverged predictably from the fungal communities in the black soil end of the gradient. AMF isolates exist which are effective in promoting plant growth over a range of edaphic and host conditions.

Gradients of variation in AMF species were related primarily to topography and burning frequency and secondarily to original plot position with experimental plot rows.

There were distinct northern, central and southern communities of fungi. *Glomus aggregatum* and *G. fasciculatum* occurred throughout the range of sagebrush, *G. deserticola* was restricted to the central and southern portions of its range, *Scutelllospora calospora* occurred along the western and northern portions of the range, and *G. mosseae* was southern and central in distribution. AMF form their own specific community patterns regardless of the host plant (*Artemisia tridentata*).

| Source | Factor | Research Site | Effect |
|--------|--------|---------------|--------|
| Anderson et al. (1984) | Soil fertility and moisture | Tallgrass prairie | AM fungal species richness was positively correlated with percentage soil organic matter and negatively correlated with Ca, Mg, and P content of soil. |
| Nadian et al. (1997, 1998) | Soil compaction | Greenhouse: pots | Total P uptake was significantly greater in mycorrhizal plants than in non-mycorrhizal ones; the response was smaller as soil compaction was increased. Soil compaction to a bulk density 1.6 Mg m\(^{-3}\) had no effect on the percentage of root length colonized, but total root length colonized decreased as soil compaction was increased. Soil compaction, which increased bulk density from 1.20 to 1.75 Mg m\(^{-3}\), reduced O\(_2\) content of the soil atmosphere from 0.16 to 0.05 m\(^3\) m\(^{-3}\). |
| Jakobsen and Nielsen (1983), Zajicek and Hetrick (1986), Virginia et al. (1986) | Soil depth | Field grown crops | The proportion of root length infected decreased markedly below 40 cm soil depth; root density varied greatly between crops, whereas the absolute length of infected roots was similar. The degree of colonization varied with plant species and soil depth; spores of *Glomus fasciculatum* were found to a depth of 220 cm. *Prosopis glandulosa* developed functional root symbiotic associations with N\(_2\)-fixing nodules and AMF at depths greater than 4 m in moist soil above a seasonally stable water table. |
| Cooke et al. (1993), Miller (2000) | Soil H\(_2\)O saturation | Salt marsh grasses | Roots were colonized by AMF to a depth of 42.5 cm, but none arbuscule was observed below 37.5 cm. AM fungal colonization was strongly, negatively correlated with water depth; colonization was lowest in plots that were consistently wet but rose as some plots underwent seasonal drying; soils that were wet for > 1 year had the same ability to form mycorrhizas in bait plants as those that had remained dry. |
| Green et al. (1976), Porter et al. (1987a, b) | Soil pH | Greenhouse: pot cultures | AMF spores germination was influenced by pH: *Glomus mosseae* germinated best at pH 7, *Gigaspora coralloidea* at pH 5, and *G. heterogama* at pH 6. pH influenced the ability of infection, sporulation, and spore germination: *Acaulospora leavis* was distributed in soil samples ranging from pH 4.5-4.9, two species of *Gigaspora* from pH 4.5-6.4 and three different strains of *Glomus* at pH 5.5-8.4. |
| Johnson et al. (1992), Sylvia et al. (1993) | Diverse soil factors (composition, texture, structure, nutrients, pH, etc.) | Plots of monocultures of five successional grass species | Even closely related hosts (five grasses) may cause divergence in AM fungal communities on initially identical soils; fungal communities in the sandy end of the soil gradient diverged predictably from the fungal communities in the black soil end of the gradient. AMF isolates exist which are effective in promoting plant growth over a range of edaphic and host conditions. |
| Gibson and Hetrick (1988) | Topography and burning frequency | Plots in a tallgrass prairie | Gradients of variation in AMF species were related primarily to topography and burning frequency and secondarily to original plot position with experimental plot rows. |
| Koske (1987) | Temperature | Plots on a latitudinal gradient along a barrier dunes | Average AM fungal species richness was positively correlated with distance south along the gradient and with temperature parameters. |
| Allen et al. (1995) | Latitudinal gradient of occurrence | Plots in the Mojave Desert | There were distinct northern, central and southern communities of fungi. *Glomus aggregatum* and *G. fasciculatum* occurred throughout the range of sagebrush, *G. deserticola* was restricted to the central and southern portions of its range, *Scutelllospora calospora* occurred along the western and northern portions of the range, and *G. mosseae* was southern and central in distribution. AMF form their own specific community patterns regardless of the host plant (*Artemisia tridentata*). |
Arbuscular mycorrhizal fungal communities present differences in relative density, spore production and differential activities, correlated to the environment (e.g. rainy season vs. dry season), to the host plant species (e.g. host specificity) and to other fungal species (e.g. competitive interactions) (Dhillion and Anderson, 1993; Gange et al., 1993; Abbot and Gazey, 1994; Miller, 1995; Merryweather and Fitter, 1998a, 1998b; Titus and del Moral, 1998a, 1998b; Turner and Friese, 1998). The competitive ability of AMF can be affected by many factors, such as the host plant species, particular environmental conditions, AMF phenology, inoculum frequency and its differential incubation time, the spatial distribution of the interacting propagules (Koske, 1981; Wilson, 1984), and the specific AMF genetic traits (Sanders et al., 1996), as well as by the presence of ectomycorrhizae (Moyersoen et al., 1998). These findings illustrate the importance of selecting an efficient AM fungal strain, native or introduced, to inoculate into disturbed soils for ecological restoration or for agricultural purposes (Lambert et al., 1980; Porter et al., 1987b; Stahl et al., 1988).

The hypothesis that environment, soil, and plant communities influence more AMF distribution than the specific AMF-plant association, is partially supported by these studies. However, other studies have demonstrated that there is a functional relationship between the species of AMF selected by a particular plant species (Hayman, 1982; Carey et al., 1992; Johnson et al., 1992; Dhillion and Anderson, 1993; Hartnett et al., 1993; Sylvia et al., 1993; West et al., 1993a, b; Allen et al., 1995; Clapp et al., 1995; Miller, 1995; Newsham et al., 1995b; Titus and del Moral, 1998a, b). Particular AMF species are able to provoke specific effects on plant performance (Stahl et al., 1990; Streitwolf-Engel et al., 1997; Van der Heijden et al., 1998; Hartnett and Wilson, 1999). For instance, AMF influence the clonal growth of two species of Prunella (Lamiaceae), P. vulgaris and P. grandiflora (Streitwolf-Engel et al., 1997). Therefore, AM fungal species and/or communities influence plant communities through either stabilizing or destabilizing feedback mechanisms (Van der Heijden et al., 1998).

Establishment of mycorrhizal fungi

E. Stahl (1900, Jahrbücher für Wissenschaftliche Botanik 34:539-668, cited in Allen, 1991) divided plants into nonmycotrophic, facultatively mycotrophic and obligately mycotrophic in relation to plant families; however, it would be unwise to assume that a plant would be non-mycorrhizal or mycorrhizal because it belongs to a particular family (Newman and Reddell, 1987), since it has been established that mycotrophy or non-mycotrophy is an expression of ecological adaptation (Bellgard, 1991). It was believed that within a community, the lack of mycorrhizae was a characteristic of early successional habitats (Nicolson, 1960; Pendleton and Smith, 1983). Nowadays, it is known that ruderal plant species, which frequently colonize disturbed sites, are often facultative mycotrophs (Francis and Read, 1994); Corkidi and Rincón (1997) observed the mycotrophy of seven plant species distributed in different successional stages in a sand dune ecosystem along the Gulf of Mexico shore; however, it looks like early successional annuals are predominantly non-mycotrophic.

For their establishment, AMF must invade and colonize roots (figure 3), either if AMF arise from a localized contact or from a dispersed inoculum. Allen (1991) considered that

**Figure 3.** Factors affecting mycorrhizal fungi (MF) establishment.
inoculum density, host and fungal genetic compatibility, edaphic factors, and plant-microbial activity influence the formation of the mycorrhizal association.

According to Morton (1990), once genes for specialized processes interacting with both plants and AMF were widely distributed among the symbionts, only those characters responsible for co-adaptational changes in mycorrhizal phenotypes continued to evolve in clonal subunits. These processes resulted in coadaptation between arbuscular clones and those hosts having compatible “symbiosis genes”. Recently, it has been strongly suggested that the degree of compatibility or “specificity” between AMF and host plant might rely on specific biochemical regulatory processes initiated in the host as a result of the attempts of colonization by the fungus; the induction of compatibility or incompatibility reactions must be thus determined by the particular host-fungus combination (Douds et al., 1998). Furthermore, Hayman (1982), Fitter (1991), and Johnson et al. (1997) described the AMF-host plant association as a mutualism-parasitism continuum related to the cost-benefit received by the plant. Mycorrhizal associations are beneficial (mutualistic) to plants when net costs are less than net benefits (e.g. improved access to limiting soil resources), and detrimental (parasitic) when costs exceed benefits (e.g. reduction of root growth).

In the case of edaphic factors (table 1), they affect the ability of both plant and AMF to disperse and survive. High available nutrient levels and saturated soils often reduced mycorrhizal associations (Powell, 1980; Anderson et al., 1984). Allen (1991) formulated two assumptions in order to explain this process: (1) if the plant has access to all necessary resources, the association will not form or will be strongly reduced as the fungus then becomes a carbon drain, and (2) the fungus will invade any root encountered. Both hypotheses assume that the plant is the regulating agent.

The activity of neighboring plants is also known to affect mycorrhizal formation. Competition between plants determines the available resources. If a high density exists, resources will be lower and, just as low soil nutrient concentrations can stimulate colonization, increasing competition can result in resource depletion and increase AMF colonization (Allen and Allen, 1980; Grime et al., 1987; Gange et al., 1993; Van der Heijden et al., 1998; Hartnett and Wilson, 1999).

On the other hand, AMF do not colonize regions infected by endoparasitic nematodes, and nematodes rarely infect regions colonized by AMF; both organisms are often mutually exclusive, each reducing the population of the other (Ingham, 1988). Hence, antagonistic interactions take place between soil organisms and AMF, as well as predation on hyphae by nematodes and microarthropods (St. John and Coleman, 1982; Fitter, 1985).

The rate and intensity of spreading and colonization differs among AM fungal species, as well as the type of colonization structures (root length occupied by hyphae, arbuscules or vesicles) (Duke et al., 1994). It has been suggested that the type and rate of colonization of the different structures may be more important than the absolute amount of fungal biomass in the root, indicating that aspects of the AM fungus-plant relationship may be different in the same plant and/ or in different plant species through time in relation to the exploitation of temporary nutrient pulses or patches, and to phenological changes; it ratifies that several AM fungal species can be involved in a single association (Miller et al., 1989; Sanders and Fitter, 1992; Duke et al., 1994).

Mycorrhizal fungi have a wide range of specificity responses (Warcup, 1981; Molina and Trappe, 1982). Some, such as many species of Glomus, show almost no specificity within plant species capable of forming arbuscular mycorrhizae (AM) (Van der Heijden et al., 1998; Hartnett and Wilson, 1999). In some Australian orchid associations (Warcup, 1981), EM fungal specificity was highly variable: some associations were highly specific (e.g. Caladenia and Sebacina vermifera) and others were not. Allen (1991) pointed out that many ericoid mycorrhizal fungi can associate with a wide diversity of heathland shrubs, which tend to be highly stable through time; these shrubs do not form mycorrhizae with other plants; this ensures these fungi a host, despite changes in plant community.

Nevertheless, even if host-specific EMF are able to form ectomycorrhizas with a wider range of hosts, sexual reproduction via sporocarp production may still be mediated through specific host association (Molina and Trappe, 1982). On the other hand, many of the fungi that form arbuscular mycorrhizae show little specificity (Clapp et al., 1995). Arbuscular mycorrhizae (AM) can be formed on herbs as well as shrubs and trees (Allen, 1991).

However, other studies have shown that particular AM fungal species can be highly specific; in this case, AM fungal specificity is related to specialized habitats. For instance, Dhillon (1992) demonstrated that indigenous mycorrhizal fungal isolates showed considerable amount of host preference in a tall grass prairie (e.g. Glomus geosporum, Schizachyrium scoparium, and Andropogon gerardii), and Hartnett and Wilson (1999) concluded that there is a differential host species response to fungal specific colonization that can explain the dominance of C4 perennial grasses in a tallgrass prairie.

Finally, it is important to point out that both AMF and EMF often grow together in the same root (Bellgard, 1991). Both fungal groups are equally able to colonize the same niche, although competitive exclusion can take place; EMF synthesize enzymes that can inhibit AMF growth (Moyersoen et al., 1998).

Final comments

In part, the limited studies of the AM association as a function of both organisms is due to the different scales needed
and the complexity of interactions between levels of organization within the system: AM fungal species-host plant. For a better understanding on the mechanisms and interactions related to AMF evolution and ecology through their life cycle, further research is needed (figure 4). For instance, research on mycorrhizal fungi considers the effects of the association on the physiology or production of individual plants; the two mutualists are thus rarely studied as individual organisms which interact enhancing the survival of each other.

A recent approach to the arbuscular mycorrhizal association is focused on the importance of AMF in the ecological restoration of disturbed areas. For example, some studies carried out in the arid and semi-arid regions of Mexico have shown that most of the established perennial plant species are mycorrhizal (Carrillo-García et al., 1999; Camargo-Ricalde et al., in press), and that AM fungal spores are important components of the resources islands created by different shrubs and trees (e.g. *Bursera fagaroides, Mimosa luisana*) (Carrillo-García et al., 1999; Camargo-Ricalde and Dhillion, in press); however, AM fungal population dynamics remain not well understood.

Finally, to completely understand the AM fungal population dynamics, an integrative research including both, the above- and below-ground organisms interactions and environmental factors, is needed.

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**Figure 4.** Arbuscular mycorrhizal fungi (AMF) evolution and ecology must be understood in terms of AMF contribution to both AMF and plant fitness. AMF life cycle responds to surrounding environment; thus, it expresses how these fungi have adapted to natural selection pressures through time. In ecological terms, AMF life cycle points out the factors and interactions involved in their reproduction (e.g. specific AM fungal species-host plant), dispersal (e.g. new habitats and root to root contact), distribution (e.g. environment, soil and plant community), and establishment (e.g. specificity, competition, mutualistic-parasitic continuum association).

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