Seasonal Dynamics of Mycorrhizal Colonization in the Rhizosphere of Some Dominant Halophytes

A. FŰZY, T. TÓTH and B. BIRÓ

Research Institute for Soil Science and Agricultural Chemistry (RISSAC) of the Hungarian Academy of Sciences, Budapest

Introduction

Attention has already been called to the importance of mycorrhizal symbiosis in plant health and fitness in the literature (BIRÓ et al., 2000, 2005b; CANTRELL & LINDERMAN, 2001; HIRREL & GERDEMANN, 1980). The symbiosis, however, is a sensitive and dynamic connection between the fungi and the host plants. Other microbiological counterparts therefore can reflect the soil–environmental conditions and its functioning, especially in the case of environments under stress (BIRÓ, 2001; KÁTAI, 1994). During the vegetation period interactions have site-specific spatial and temporal patterns due to the dynamic plasticity of the mycorrhizal symbiosis. Several authors described the annual variability of the symbiotic structures on the same host plants, grown in the same environments (MILLER, 1999; CARVALHO et al., 2001).

The rate of fungal intensity and function can be greatly influenced by biotic and abiotic environmental factors and versatile environmental conditions (BIRÓ et al., 2005a,b). The importance of climatic zonality, thus the value of macro- and microclimatic conditions in the vegetation and in the symbiotic relationship was also reported (ALLEN et al., 1998; BRAUNBERGER et al., 1997; SZILI-KOVÁCS et al., 2000). The geographical site of a certain species can determine the seasonal variability of the soil- and rhizobiological status even on the same soil types (BIRÓ, 2001; VAN DUIN et al., 1990; BIRÓ et al., 2002).

Mycorrhizal fungi are obligate endosymbionts of about 80% of the higher plants. The ratio of arbuscular mycorrhizal (AM) colonization is greatly affected by the host plants and their physiological status, providing a dynamic fluctuation during the vegetation periods (BIRÓ et al., 2005a). Relatively few studies have been carried out, however in this respect on the Hungarian steppe (HILDEBRANDT et al., 2001; LANDWEHR et al., 2002).

The present work’s aim was the regular monitoring of mycorrhizal colonization on the most dominant halophytes to understand the: 1. mechanism of seasonal or annual variability; 2. the interrelations with the main soil characteristics and water
fluctuations; 3. the role of zonality in the species-specific characteristics between the macro- and the microsymbionts; and 4. the site-climatic differences at two selected saline sites in Hungary.

**Materials and Methods**

**Site description and sampling procedures**

Soil and root samples were collected from the rhizosphere of the most typical halophytes, such as the *Plantago maritima, Aster tripolium, Artemisia santonicum, Puccinellia limosa, Festuca pseudovina* and *Lepidium crassifolium*, grown on the

<table>
<thead>
<tr>
<th>Table 1</th>
<th>Soil characteristics of the selected sampling sites (Hungary, 2001)</th>
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</thead>
<tbody>
<tr>
<td>Measured parameters</td>
<td>Apajpuszta (Site A)</td>
</tr>
<tr>
<td>Coordinates</td>
<td>47°5.2’ N; 19°5.8’ E</td>
</tr>
<tr>
<td>Main features</td>
<td>grassland, occasionally pastured</td>
</tr>
<tr>
<td>Soil type</td>
<td>solonetz</td>
</tr>
<tr>
<td>Specific salts</td>
<td>Na₂CO₃–Na₂SO₄</td>
</tr>
<tr>
<td>pH</td>
<td>8.51–9.64</td>
</tr>
<tr>
<td>EC of 1:2.5 soil:water suspension (dS·m⁻¹)</td>
<td>0.38–0.82</td>
</tr>
<tr>
<td>OM (%)</td>
<td>0.86–1.78</td>
</tr>
<tr>
<td>Clay content (%)</td>
<td>9–26</td>
</tr>
</tbody>
</table>

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<tr>
<th>Table 2</th>
<th>Meteorological data of the Kiskunság region during the two sampling periods</th>
</tr>
</thead>
<tbody>
<tr>
<td>Months</td>
<td>2001</td>
</tr>
<tr>
<td>January</td>
<td>69 (-)*</td>
</tr>
<tr>
<td>February</td>
<td>2 (-)*</td>
</tr>
<tr>
<td>March</td>
<td>64 (43)*</td>
</tr>
<tr>
<td>April</td>
<td>37 (20)*</td>
</tr>
<tr>
<td>May</td>
<td>17 (11)*</td>
</tr>
<tr>
<td>June</td>
<td>94 (84)*</td>
</tr>
<tr>
<td>July</td>
<td>127 (55)*</td>
</tr>
<tr>
<td>August</td>
<td>21 (17)*</td>
</tr>
<tr>
<td>September</td>
<td>98 (85)*</td>
</tr>
<tr>
<td>October</td>
<td>14 (1)*</td>
</tr>
<tr>
<td>November</td>
<td>39 (41)*</td>
</tr>
<tr>
<td>December</td>
<td>13 (15)*</td>
</tr>
</tbody>
</table>

*data of a 20-day period before the root and soil sampling
saline soils of two characteristic sites (site A: Apajpuszta, site Z: Zabszék) in Hungary. The soil and root samples contained the rhizosphere of at least five plants grown on the same spot (the similar salt concentration was measured by a portable field conductometer).

The most important soil physical and chemical characteristics are given in Table 1. Data on the weather conditions, such as the temperature and the precipitation at the sampling sites for the two vegetation periods are summarized in Table 2.

**Soil and root analyses**

Samples of 1 g fresh lateral roots of the main halophytes were randomly removed, cleansed in tap water and stored in 70% alcohol before examining the AMF colonization. After storage root samples were cut to approximately 1 cm segments, boiled in a 15% KOH solution for 40 minutes, stained with aniline blue and fixed in 40% lactic acid, as it was suggested by PHILLIPS and HAYMAN (1970). After these procedures, the colonization characteristics were studied according to the five-class system of TROUVELOT et al. (1986). A 100×, 200× or 400× magnification was used by an Olympus microscope to assess the accurate AMF structures in the root segments.

The presence and abundance of hyphae, vesicles and arbuscules were determined in 30 root segments for each sample. Frequency and intensity of mycorrhizal colonization (F%, M%) and the absolute and relative arbusculum richness (A%, a%) were estimated and calculated after the assessments.

**Fig. 1**

Relative location of the 6 main halophyte species on the toposequence and the concomittant salt concentration (EC, dS·m⁻¹) at the study areas (site A: Apajpuszta and site Z: Zabszék)
Statistical analyses. – One-way variance analysis was performed to follow the seasonal changes. Results are shown in figures, including the least significant differences (LSD5%).

Results and Discussion

Host specificity in mycorrhizal colonization

Fig. 1 shows the location of the investigated plant species along the toposequence in Apajpuszta. *Plantago maritima* is a common plant and can be found at all points of the toposequence, except in the bare spots. *Artemisia santonicum* and *Festuca pseudovina* originate from an association of the higher altitudes, while *Aster tripolium* and *Puccinellia limosa* are the dominant species in wetland elevation zones (the right side of Fig. 1). In the bare spots the only plant is the Brassicaceae *Lepidium crassifolium*.

The colonization values of the five investigated plant species are presented in Fig. 2. In the case of higher altitude with less salt in the soil more than half of the root system is colonized (*Plantago maritima, Artemisia santonicum* and *Festuca pseudovina*), at lower altitude (5–20 cm lower) with more salt in the soil the colonization rate is lower, but remarkable (*Puccinellia limosa*), or seasonally can be quite high (*Aster tripolium*). During summer periods the symbiosis is detectable also on *Lepidium crassifolium*, which is generally known as a non-mycotrophic halophyte. Species specificity is recognizable in the mycorrhizal functioning of the selected investigated plants, as well. Arbusculum richness was especially high in the case of *Plantago maritima*, while it was almost zero for *Puccinellia limosa*. According to these results it can be stated that the colonization ratio of the arbuscular mycorrhizal fungi in the rhizosphere of the most dominant halophytes is greatly determined by a host-specificity, as it was also established by Carvalho et al. (2001). In addition to these plant physiology driven properties, the physical and chemical characteristics of the saline soils and the weather conditions can dramatically affect the temporal and spatial variability of the host-mycorrhizal symbiosis, as shown in the following.

Site variability of the mycorrhizal colonization

Two characteristic saline sites were selected for the study. Site A (Apajpuszta) has meadow solonetz soil and is occasionally pastured by sheep. Site Z (Zabszék) has a solonchak soil in the vicinity of the Zabszék lake, and therefore, it has a periodic water flooding situation. The main soil characteristics of the sites are summarized in Table 1.

Fig. 3 shows the arbuscular mycorrhizal colonization parameters of two selected halophytes. Apparently a much higher mycorrhizal colonization (M%) occurred on both plant species at the Apaj study area. The intensity of mycorrhizal colonization at this site was between 40–100% and only 20–60% as an average at the Zabszék site on *Aster tripolium*. This plant had a higher value of arbusculum richness (A%)
Fig. 2
Mycorrhizal colonization of dominant halophytes at Site A (Apajpuszta) in 2001 from March to December. The whole column represents the colonization intensity (M%), the white part of the column stands for the arbusculated part of the root system (A%).
Fig. 3

Mycorrhizal colonization of two selected halophytes at the two salt affected sites in 2002 from March to November. The whole column represents the colonization intensity (M%), the white part of the column stands for the arbusculated part of the root system (A%).

In comparison with the *Puccinellia limosa*, which had almost zero arbuscularity at both sites. A quite high A% value was recorded only in July on the *Aster tripolium* grown in the Zabszék area. This intensive colonization reflects an improved functioning of the AM fungi under the highly stressed environmental conditions (i.e. in summer), similarly to earlier findings (FÜZY et al., 2003; BIRÓ et al., 2000). At the Zabszék site the water content of the topsoil was found to be near zero, and the host plant, *Aster tripolium* has a greater inductive capacity for the symbiosis, to survive the natural stress conditions. In other periods of the year water fluctuation is intensive at this site, therefore the mycorrhizal colonization value rather depends on the soil’s water content and the parallel severity of other environmental stresses (like high or low temperatures, salt content etc.). Such importance of the soil water conditions in the environment is already a well-known phenomenon (VARALLYAY, 1999). On the long-term, however, characteristic site-specificity can be recorded on saline soils, which can be reflected on the basis of the plant–microbe interactions, similarly to earlier findings on heavy metals (KÁDAR et al., 2001).

**Temporal changes in mycorrhizal colonization**

Root samples of *Plantago maritima* were collected at four different points on the Apajpuszta site. It was possible, therefore to compare the colonization variability on the same host plant at a particular site during the vegetation period. Data presented on Fig. 4 show the mycorrhizal values from March 2001 to February 2002 with monthly frequency. An especially characteristic dynamism of the colonization...
pattern was found on the *Plantago maritima*, known to be a good-mycotroph plant species. Mycorrhizal colonization increased significantly during the spring period up to the early summer (to June). A concomitant decrease was observed, however, from June to July, and an enhanced mycorrhizal intensity (M%) was recorded between July and November. A similar tendency was also observed for the arbusculem richness (A%) values, with two characteristic maximum levels in May–June and August–September. The importance of the AM fungi in the nutrient supply is more pronounced in these periods, which may intensify the functioning of the microsymbiont partner.

The decrease in mycorrhizal colonization during the summer period is correlatable by the negative effect of high temperature and drought, which are known stressors in environmental studies (HILDEBRANDT et al., 2001; LANDWEHR et al., 2002). In those stressed soils the stress-buffering effect of the arbuscular mycorrhizal fungi can be developed (RUIZ-LOZANO et al., 1996; BÍRÓ et al., 2005b). On a long-term level, however, the species diversity and the specific functioning of the fungi may also be reduced (HILDEBRANDT et al., 2001; LANDWEHR et al., 2002). According to this finding, it is assumed that the highly salt-tolerant *Glomus geosporum* is the main mycorrhiza type in saline soils that can be involved in the stress reduction of the most dominant halophytes, shown in this study.

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**Fig. 4**

The mycorrhizal colonization of *Plantago maritima* at site A (Apajpuszta) in 2001

A. Mycorrhizal intensity (M%). B. Arbusculem richness (A%). The figures show the mean data and LSD_{5%} values, as a result of one-way variance analysis.
Summary

Dominant halophytes, such as Plantago maritima, Aster tripolium, Artemisia santonicum, Puccinellia limosa, Festuca pseudovina and Lepidium crassifolium were monitored for their colonization by arbuscular mycorrhizal fungi (AMF) for two vegetation periods, sampled by monthly frequency. Two saline sites (A: Apajpusza and Z: Zabszek) were selected for the survey due to their similar physical and chemical soil characteristics (including the salt level and salt specific anions) and the distinct site use (pasturing or not) or water regime (as drought or temporally flooded at the vicinity of the Zabszek lake).

Colonization of arbuscular mycorrhizal fungi greatly depended on the plant species and their physiological status, such as the mycotroph or non-mycotroph character. In this respect the highest mycorrhizal intensity (M%) and arbusculum richness (A%) was found on the Plantago maritima, while Puccinellia limosa was the less dependent species on mycorrhiza fungi.

A characteristic seasonal dynamism was found at both saline sites, for all sampled halophytes. A maximum mycorrhizal colonization was recorded in late spring and early summer, when there was an increased plant-physiological initiative for the helpful symbiosis (i.e. intensive vegetative and/or generative phases of the hosts).

The rate of fungal intensity in the root system (M%), and especially the arbusculum richness (A%) on the other hand was found to be dependent on the site use and the water fluctuation in the soils. A more versatile dynamism of the mycorrhizal colonization was found therefore at the vicinity of the Zabszek lake with the permanent water fluctuations. Under more drought stressed conditions an increased mycorrhizal colonization and functioning (arbusculum richness) was found, generally preceeding the high environmental stress, which was deleterious for both symbiont partners. Such mycorrhizal dynamism in the rhizosphere, however, seems to be a common strategy between the hosts and the microsymbionts in the “struggle for life” process in the Hungarian steppe.

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Key words: arbuscular mycorrhiza, halophytes, seasonal dynamism, drought stress, salt stress

References


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