Adaptation strategies of rare plant species to heterogeneous soil conditions on a coast of a lagoon lake as revealed by analysis of mycorrhizal symbiosis and mineral constituent dynamics

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Abstract

Soil chemical heterogeneity in sea-affected wetlands and its effect on mineral nutrition is a seldom studied aspect in ecophysiology of wild plants. The aim of the present study was to analyze seasonal changes in concentration of different mineral nutrients in leaves of several salt marsh species (Aster tripolium, Glaux maritima, Plantago maritima, Trifolium fragiferum, Triglochin maritima) in comparison to changes of plant-available nutrients in soil, with emphasis on mycorrhizal symbiosis. Extreme heterogeneity was noted for soil mineral nutrient concentrations, varying both spatially and temporarily. A pronounced correlation between the concentrations of different soil mineral nutrients was observed. Early season flooding at the end of May coincided with a peak of summary precipitation and high sea level, and eventually resulted in significant increase in soil salinity in June. Peaks of Na and Cl accumulation in plant leaves in June were clearly related to the corresponding peak of soil concentrations of the respective ions. Seasonal trends of soil mineral concentrations were compared with respective leaf mineral concentration dynamics of the five species studied within the season. In general, a trend of leaf N concentration followed that for soil N content. Minimum intensity and frequency of mycorrhizal symbiosis in July corresponded to the peak of soil salinity in June for roots of A. tripolium, G. maritima, P. maritima, and, possibly, T. fragiferum. Mycorrhizal symbiosis of G. maritima was the most sensitive to these conditions, with the intensity close to zero in July and August. Both intensity and frequency of mycorrhizal symbiosis in roots of T. maritima was highest in July, with significant decrease in August and September. The presence of arbuscules was evident in roots of all species, but with different intensity at different time points. The results of this study support the hypothesis that wild plants native to habitats with heterogeneous soil conditions have developed effective physiological adaptations allowing to grow normally and reproduce within a wide range of mineral concentration, as well as drastic changes in soil salinity.

Key words: chlorophyll a fluorescence, mineral nutrients, mycorrhizal symbiosis, physiological adaptations, rare coastal plant species, soil chemical heterogeneity.

Abbreviations: EC, electrical conductivity; PI, Performance Index.

Introduction

Heterogeneous soil conditions affect plant localization and growth in salt marshes. In sea-affected marshes, fluctuations in water level lead to high heterogeneity in soil characteristics, including salinity and availability of mineral nutrients (Karlsons et al. 2011). In particular, shortage of nitrogen together with a high abundance of P, Mg, Fe, Zn, Cu on the background of significantly increased NaCl concentration are the most important characteristics of coastal wetlands. However, mineral nutrition of coastal marsh species is a less-studies aspect of coastal plant ecophysiology. In general, while the soil salinity-mineral nutrient relationship has been thoroughly studied with crop species (Grattan, Grieve 1999), the same problem has been less frequently investigated for wild plants in natural conditions.

It can be suggested that plant species composition of moderately saline temperate wetlands largely reflects a multitude of edaphic conditions. A number of relatively rare coastal marsh species are of special interest for ecophysiological studies. Several characteristic coastal marsh species (Plantago maritima, Triglochin maritima) represent typical leaf-succulent euhalophytes, characterized by almost cylindrical or globular leaves (Breckle 2002). Trifolium fragiferum, on the other hand, is a clonal species and typical legume, possessing rhizobial symbiosis enabling to use nitrogen fixed from the air, and it is designated as facultative halophyte (Rumbaugh et al. 1993; Nienartowicz, Wilkon-Michalska 1993). Aster tripolium is a salt-recreting

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halophyte, having rapid turnover of leaves, replacing older leaves that have a high concentration of salt (Brekcke 2002). *Glaux maritima* is a species capable for vegetative propagation characterized by the presence of salt glands with a relatively high efficiency of salt removal (Rozema et al. 1978).

Usually it is assumed that mycorrhizal symbiosis is an important adaptation of wild plants to nutrient-limited conditions (Ruiz-Lozano, Azcón 2000) and it can also contribute to stress tolerance (Feng et al. 2002). In a complex study of putative mycorrhizal structures in roots of 29 coastal plant species it was revealed that coastal salt marsh species *Aster tripolium*, *Glaux maritima*, *Plantago maritima*, *Trifolium fragiferum*, *Triglochin maritima* display mycorrhiza-specific structures in their roots, including hyphae, vesicles and arbuscules (Druva-Lusite, Levinsh 2010), pointing to existence of functionally active symbiosis in these plants. However, the intensity of symbiosis was variable and at relatively low level, reaching above 10% only for *Aster tripolium*, *Plantago maritima* and *Trifolium fragiferum*. It was argued that increased soil salinity is among the main environmental factors resulting in decreased intensity of mycorrhizal symbiosis of salt marsh species (Druva-Lusite, Levinsh 2010).

It is not completely understood how different plant species adapt to highly heterogeneous soil conditions of saline wetland. The most important problem concerns physiological mechanisms in saline wetland plants with possible adaptive importance, including mycorrhizal symbiosis and adaptivity of mineral nutrition. It can be suggested that specific physiological adaptation mechanisms exist to maintain mineral equilibrium in plant tissues at optimal levels through acquisition, transport and efficiency of use of mineral nutrients. The aim of the present study was to analyze seasonal changes in concentration of different mineral nutrients in leaves of number of salt marsh species in comparison to changes of plant-available nutrients in soil, with emphasis on mycorrhizal symbiosis.

**Materials and methods**

Wet meadows on the coast of Lake Liepaja are periodically flooded by water from the lake during heavy and prolonged rains through changes in water level and are affected by the Baltic Sea. Lake Liepaja is a typical lagoon-type lake, in the northern part connected with the Baltic Sea by a channel which results in relatively high water salinity at the northern (8 to 10 dS m⁻¹) and central part (2 to 4 dS m⁻¹) due to the action of prevailing northwest winds (Latvian Environmental Protection Fund 2008). This forms characteristic coastal salt marsh vegetation on the coasts of the Lake with several rare or especially protected species [*Aster tripolium* L., *Blysmus rufus* (Huds.) Link, *Glaux maritima* L., *Juncus gerardii* Loisel., *Plantago maritima* L., *Spergularia salina* J. et C. Presl, *Triglochin maritima* L. etc.]. However, characteristic halophyte vegetation has established only in places with significant anthropogenic pressure, around paths and near the coast, or in the form of small islands surrounded by relatively high herbaceous vegetation composed mostly of *Phragmites communis* L., *Typha* spp. and various characteristic dicotyledonous wet meadow species.

The particular study site along Eduarda Tīsē Street in Liepaja was localized in the part of the lake with water salinity about 7 dS m⁻¹ (Latvian Environmental Protection Fund 2008). The water in this area has a high concentration of dissolved P (> 0.08 mg L⁻¹). As model species for this study, the especially protected coastal marsh species *Aster tripolium*, *Glaux maritima*, *Plantago maritima*, *Trifolium fragiferum*, and *Triglochin maritima* were chosen (Cabinet of Ministers of Latvia 2000). Three subsites identically affected by water level changes were selected within a different distance from the coast (3 to 15 m, 120 m, 170 m) in places where the model plant species were located. Small-scale spatial heterogeneity in soil properties were eliminated by sampling in different locations within the particular subsite and by using plant material from various locations for mineral constituent analysis. Every month three to five samples were taken from each subsite and analyzed separately for concentration of plant-available mineral nutrients, pH and EC as described previously (Karlsons et al. 2008). Average monthly values from each subsite were used to calculate average values for the site.

Plant material for chemical analysis was collected monthly from May till September within the period between the dates 10th and 15th. To ensure non-disturbing plant sampling, three leaf samples were collected for each

**Table 1.** Variation of soil mineral nutrient, Na and Cl concentration (mg L⁻¹), pH and EC during the vegetation season. *n* = 38

<table>
<thead>
<tr>
<th>Mean value ± SE</th>
<th>Range</th>
<th>Coefficient of variation (%)</th>
</tr>
</thead>
<tbody>
<tr>
<td>Nitrogen (N)</td>
<td>45 ± 4</td>
<td>14 ÷ 108</td>
</tr>
<tr>
<td>Phosphorus (P)</td>
<td>224 ± 27</td>
<td>92 ÷ 916</td>
</tr>
<tr>
<td>Potassium (K)</td>
<td>98 ± 11</td>
<td>17 ÷ 270</td>
</tr>
<tr>
<td>Calcium (Ca)</td>
<td>8564 ± 800</td>
<td>1313 ÷ 22938</td>
</tr>
<tr>
<td>Magnesium (Mg)</td>
<td>1799 ± 06</td>
<td>650 ÷ 2875</td>
</tr>
<tr>
<td>Sulphur (S)</td>
<td>510 ± 99</td>
<td>40 ÷ 2400</td>
</tr>
<tr>
<td>Iron (Fe)</td>
<td>1740 ± 252</td>
<td>300 ÷ 8350</td>
</tr>
<tr>
<td>Manganese (Mn)</td>
<td>94 ± 16</td>
<td>8 ÷ 400</td>
</tr>
<tr>
<td>Zinc (Zn)</td>
<td>29 ± 4</td>
<td>4 ÷ 90</td>
</tr>
<tr>
<td>Copper (Cu)</td>
<td>3.0 ± 0.3</td>
<td>0.5 ÷ 7.5</td>
</tr>
<tr>
<td>Molybdenum (Mo)</td>
<td>0.039 ± 0.003</td>
<td>0.011 ÷ 0.09</td>
</tr>
<tr>
<td>Boron (B)</td>
<td>4.6 ± 0.6</td>
<td>0.2 ÷ 13.2</td>
</tr>
<tr>
<td>Sodium (Na)</td>
<td>1200 ± 184</td>
<td>225 ÷ 5150</td>
</tr>
<tr>
<td>Chlorine (Cl)</td>
<td>1357 ± 202</td>
<td>275 ÷ 6300</td>
</tr>
<tr>
<td>pH</td>
<td>7.0 ± 0.2</td>
<td>4.2 ÷ 8.5</td>
</tr>
<tr>
<td>EC</td>
<td>6.4 ± 0.9</td>
<td>1.3 ÷ 25.5</td>
</tr>
</tbody>
</table>
plant species, each consisting of one fully grown leaf from a number (at least, 10) of individual plants. Concentration of Na, Cl and mineral nutrients in leaf tissues was analyzed as described previously (Karlsons et al. 2008). Root samples for analysis of mycorrhizal symbiosis were obtained from a rhizosphere of intact plants (Andersone et al. 2011), prepared and measured as described previously (Druva-Lūsīte et al. 2008).

Chlorophyll a fluorescence was measured monthly using a Handy PEA continuous fluorescence meter (Hansatech Instruments, UK) as described previously (Druva-Lūsīte et al. 2008). For each time point, at least 10 plants per species were analyzed with five individual measurements each.

Climate data (summary precipitation and sea level changes) were obtained from the Latvian Environment, Geology and Meteorology Centre (http://www.meteo.lv) and recalculated according to the time of analysis separately for the first and the second half of the month.

**Results**

Extreme heterogeneity was noted for soil mineral nutrient concentrations, varying both spatially and temporarily (Table 1). The lowest variability in soil samples during the season (as indicated by the respective coefficients of variation) was found for Mg (33%), followed by Mo (46%). Moderate variability (55 to 63%) was characteristic for N, Ca, Cu and increased (71 to 76%) variability for K, P, B, Zn. High variability (90 to 105%) was found for Fe, Cl, Na, Mn, and the highest was for S (121%). Soil salinity was also highly variable (81%), to a large extent reflecting high variation in Na (96%) and Cl (92%) concentration.

Early season flooding at the end of May coincided with a peak of cumulative precipitation and high sea level, and eventually resulted in significant increase in soil salinity in June (Fig. 1). Peaks of Na and Cl accumulation in plant leaves in June (Fig. 2B, 3B) were clearly related to the corresponding peak of soil concentrations of respective ions (Fig. 2A, 3A). Interestingly, all species accumulated comparable concentrations of Na in June, while at the end of the vegetation season, in conditions of lowered soil Na concentration, *T. maritima* and *P. maritima* had significantly higher leaf tissues levels of Na in comparison to *A. tripolium*, *G. maritima* and *T. fragiferum* (Fig. 2B). The same phenomenon was evident for Cl ions (Fig. 3B). In addition, *G. maritima* plants accumulated relatively less Cl ions in June in comparison to Na ions.

Seasonal trends of respective soil mineral concentrations were compared with leaf mineral concentration dynamics of the five species studied within a season. In general, a trend of leaf N concentration followed that for soil N content (Fig. 4). Both *T. fragiferum* and *T. maritima* accumulated higher concentration of N, which was especially pronounced at the end of the vegetation season, with more than three-fold N concentration in comparison to the rest of the species. Another notable characteristic was high N concentration in developing leaves in May for all species analyzed except *A. tripolium*.

Plant-available concentration of soil P was high in May and June, significantly lower in July and August, with some increase again in September (Fig. 5A). A trend in tissue concentration of P was similar to that in soil for all species except *T. maritima*, but there was no increase in P concentration in September (Fig. 5B). Plant-available concentration of soil K was high in May and September with significantly lower level in June, July and August (Fig. 6A). Concentration of K in leaf tissues was at the highest level in May followed by a significant decrease in July (Fig. 6B).

In spite of the relatively stable soil Ca (Fig. 7A) and Mg (Fig. 8A) concentration during the season, some species (*P. maritima* and *A. tripolium* for Ca, and *P. maritima* for Mg) exhibited significantly higher concentration of Ca and Mg ions in May and June (Fig. 7B and 8B). Changes in soil concentration of plant available S were similar to those of P (Fig. 9A), but this was not reflected by S concentration in plant tissues (Fig. 9B). Similarly, significantly higher concentration of plant-available Fe in soil (Fig. 10A)

![Fig. 1. Changes of soil electrical conductivity (EC; A), cumulative precipitation (B) and sea level (C) in Liepaja within the vegetation season.](image-url)
Fig. 2. Changes of Na concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 3. Changes of Cl concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 4. Changes of N concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.
Fig. 5. Changes of P concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 6. Changes of K concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 7. Changes of Ca concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.
Fig. 8. Changes of Mg concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 9. Changes of S concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 10. Changes of Fe concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.
Fig. 11. Changes of Mn concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 12. Changes of Zn concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.

Fig. 13. Changes of Cu concentration in soil (A) and plant leaves (B) during the vegetation season. Data are means ± SE from 12 samples for soil concentration and from three samples for each plant species for each time point.
was not reflected by corresponding changes in leaf Fe concentration (Fig. 10B).

Plant-available concentration of soil Mn (Fig. 11A), Zn (Fig. 12A) and Cu (Fig. 13A) did not change significantly during the season, but their seasonal pattern of accumulation in plant leaves was clearly species-specific (Fig. 11B, 12B, 13B). A highly variable accumulation trend in leaves was evident also for Mo (Fig. 14B), which showed the same pattern in soil as that of Na and soil EC (Fig. 14A). Soil B concentration was high in May and September (Fig. 15A) but tissue concentration of B was stable or slowly rising (Fig. 15B).

In July and August *G. maritima* plants showed relatively high increase of Fe (7-fold) and Mn (6-fold) concentration in comparison to average values of other plants. In May and June *P. maritima* plants exhibited significantly higher tissue concentration of Ca, Mg and S in comparison with other plants, coinciding with the highest level of soil salinity (Fig. 7, 8, 9). For S, increased level in *P. maritima* leaves was maintained also in July and August, diminishing only in September (Fig. 9).

Minimum intensity and frequency of mycorrhizal symbiosis in July corresponded to the peak of soil salinity in June for roots of *A. tripolium, G. maritima, P. maritima*, and, possibly, *T. fragiferum* (Fig. 16). Mycorrhizal symbiosis of *G. maritima* was the most sensitive to these conditions, with mycorrhiza intensity close to zero in July and August (Fig. 16B). Arbuscules in roots of *G. maritima* were evident only in September with relatively high intensity (51.1%, data not shown). In contrast, both intensity and frequency of mycorrhizal symbiosis in roots of *T. maritima* was highest in July, with significant decrease in August and September (Fig. 16C). The presence of arbuscules was evident in roots of *T. maritima* plants from July till September, with increasing intensity (14.3, 20.6, 23.6%, in July, August, September, respectively; Fig. 16E). For *P. maritima*, intensity
Fig. 16. Seasonal dynamics of intensity of mycorrhizal symbiosis (M%), frequency of mycorrhizal symbiosis (F%) and intensity of arbuscules of mycorrhizal symbiosis (A%) in roots of Aster tripolium (A), Glaux maritima (B), Plantago maritima (C), Trifolium fragiferum (D), Triglochin maritima (E). Data are means ± SE from 3 measurements for each time point. M%, intensity of mycorrhiza; A%, intensity of arbuscules; F%, frequency of mycorrhiza.

and frequency of mycorrhizal symbiosis had different character. While intensity was relative low in June and July and increased in August and September, frequency was high during the season (Fig. 16C). Arbuscules in roots of P. maritima plants occurred with increasing intensity during the season (Fig. 16C). In roots of A. tripolium there was a significant decrease in both mycorrhizal intensity and frequency in July followed by a recovery later in the season (Fig. 16A). In contrast, arbuscules were at low level in July (6.2%), August (4.0%) and September (2.2%) in roots of A. tripolium. Mycorrhiza in roots of T. fragiferum showed an increasing trend from July to September (Fig. 16D), and arbuscules were found only in August (9.3%, data not shown).
Fig. 17. Seasonal changes of chlorophyll a fluorescence parameters in leaves of *Aster tripolium*, *Glaux maritima*, and *Triglochin maritima*. Data are means ± SE from 5 measurements from 10 plants for each bar.

Photochemical activity of photosystem II as indicated by chlorophyll a fluorescence parameters showed significant changes during the season for *A. tripolium* and *G. maritima* plants (Fig. 17). Decrease of $F_v/F_m$ and PI in leaves of *A. tripolium* plants in July coincided with the lowest level of mycorrhization intensity in roots. In contrast, *G. maritima* plants had low intensity of chlorophyll a fluorescence parameters in June followed by a significant increase in July. In contrast, no significant changes in photosystem II activity were evident for *T. maritima* plants, with a tendency to decrease slightly in July.

As it was found that many soil mineral nutrients had similar concentration trends during the vegetation season, relationship between concentrations of different plant-available mineral nutrients in soil was analyzed (Fig. 18). The most significant correlation ($R^2$ 0.90 to 1.00) was found between Na and Cl ion concentration, and both were closely related to soil EC. Soil N concentration had highly significant correlation with Cu, Zn, Fe, K, and S concentration ($R^2$ 0.80 to 0.89), and less pronounced with that of Na and B ($R^2$ 0.70 to 0.79). K concentration correlated with that of S, Ca with Mg, Zn with Fe, Zn with Cu and B (all 0.80 to 0.89).

Negative relationship was found between soil pH and soil EC ($R^2 = -0.678$) and the majority of soil minerals (except P, Ca, Mg, Mn and B) with $R^2$ ranging from -0.481 to -0.748. In addition, soil Mg was negatively correlated with soil Cu, Cl, Na, S, N and K concentration (with $R^2$ values from -0.306 to -0.489), and soil Ca was negatively correlated with soil Na ($R^2 = -0.403$) and Cl ($R^2 = -0.392$). At the level of plant mineral constituents, the most pronounced were correlation between Na and Cl, and Cu
Rosette-forming salt marsh plants *A. tripolium*, *P. maritima* and *T. maritima* are characterized as species with limited importance of succulence in salt tolerance, suggesting programmed senescence and removal of old leaves and continuous formation of new ones as the main strategy of salt avoidance (Albert 1975). *G. maritima* is a species with an ability to replicate vegetatively, characterized by the presence of salt glands on the surface of the shoot as an effective way of removal of salts from the tissues, as well as intracellular osmotic adaptation (Rozema, 1975). *T. fragiferum*, in turn, is a typical legume characterized by rhizobial symbiosis, which makes it possible to use air nitrogen, and it is facultative halophyte (Nienartowicz, Wilkon-Michalska 1993), with unknown salt-tolerance mechanisms. It can be expected that the studied species have different adaptive mechanisms to dramatically changing salinity and conditions of mineral heterogeneity.

Model plants in the present study can be divided in two groups in respect to accumulation of Na ions ( *P. maritima* and *T. maritima* in the first group, and *A. tripolium*, *G. maritima* and *T. fragiferum* in the second), with only slight differences between them in relation to ion accumulation at the end of the vegetation season. The first group of plants had higher tissue concentrations of Na in August and September. Differences in Cl dynamics was higher for *G. maritima*, which accumulated lower levels of Cl in the first half of the season, but later the differences between the groups diminished. Thus, in heterogeneous environment halophytes of different systematic groups accumulated similar concentrations of salts, in spite of the dramatically different potential mechanisms of salt resistance.

Changes in mineral constituent concentrations in plant tissues are often analyzed only from the point of view of optimality of the particular nutrients, without further analysis of possible functional consequences. Thus, foliar nutrient analysis has been use to predict mineral requirements of crop species (Osvalde 2011). On the other hand, when wild plants grow in native soils, they certainly possess mechanisms of adaptation to particular mineral nutrient composition and concentrations. However, it is obvious that particular changes as a result of changing environmental conditions should reflect dynamics of certain physiologically important chemical substances that contain the elements of interest, as well as reflect dynamics of equilibrium between the acquisition, active use and storage of them. Significantly large differences were found for tissue contents of mineral nutrients in the various wetland model species.

If some important biochemical mechanism is using the particular ion, it’s concentration in particular tissues can increase well above toxic threshold level. Also, detoxification of ions by means of compartmentation and complex formation is a strategy against toxicity of high tissue concentrations of particular ions. In the present study, tissue concentrations of several ions were well above
toxicity levels reported for the majority of crop plants; for example, Fe (above 100 mg kg$^{-1}$) for all species, Mn (above 300 mg kg$^{-1}$) for G. maritima. It has been shown that moderate to high soil salinity induces Mn accumulation in leaf tissues of *A. tripolium* (Karlsons et al. 2008). Therefore, accumulation of Fe and Mn in leaf tissues of *G. maritima* can be regarded as an effect provoked by a high soil salinity. Similarly, accumulation of K ions in leaf tissues of *A. tripolium* can be associated with increased soil salinity, as in controlled conditions high soil salinity resulted in significant increase in tissue K concentration (Karlsons et al. 2008).

Similar increase of Cu and Zn content seen in June in leaves of *A. tripolium* and *P. maritima* (Fig. 12, 13) coinciding with the highest soil salinity (Fig. 1) points to the involvement of both ions in some resistance mechanism against high salinity. A particular type of superoxide scavenging enzyme, superoxide dismutase, contains both Cu and Zn in the active centre of the molecule and is involved in antioxidative defense (Müller et al. 2004). It is possible, that the increase of leaf Cu and Zn concentration revealed in *A. tripolium* and *P. maritima* during an episode of high soil salinity is associated with a corresponding increase in Cu-Zn superoxide dismutase protein level and activity as a part of resistance to salinity-induced endogenous oxidative stress. It is widely accepted that salinity increases production of reactive oxygen species and that induction of a sufficient level of enzymatic antioxidative capacity is one of the internal factors of salinity tolerance in both glycophytes and halophytes (Miller et al. 2010). Indeed, superoxide dismutase activity increased in *A. tripolium* plants grown under 75% seawater salinity (Geissler et al. 2009). In addition, Cu accumulation is enhanced in *A. tripolium* at increased soil salinity in controlled conditions (Karlsons et al. 2008).

Exclusion of the excess amount of ions from photosynthetic tissues with accumulation in roots is a typical adaptive mechanism of plants allowing not to reach toxic level (Karlsons et al. 2011). This is especially important for transitional metals, accumulation of which can lead to formation of hydroxyl radicals in photosynthetic tissues (Müller et al. 2004). Thus, *T. maritima* plants grown in a substrate resembling natural soil accumulated 5-fold level of Fe, 10-fold level of Mn, 2-fold level of Zn and 2-fold level of Cu in roots in comparison to that in leaves (Karlsons et al. 2011). In the present study, however, *G. maritima* plants accumulated relatively high tissue concentration in leaves of both Fe and Mn, reaching 1500 and 600 mg kg$^{-1}$, respectively, in August (Fig. 10B, 11B). This concentration is far above critical toxic levels of Fe (100 mg kg$^{-1}$) and Mn (300 mg kg$^{-1}$) in plant tissues reported (Macnicol, Beckett 1985). However, no physiological signs of toxicity were noticed. In addition, *P. maritima* was accumulating high concentration of Ca, Mg and S in leaves, especially, at the beginning of the season (Ca, Mg) or even later (S). Concentration of S in leaves was four-fold the mean concentration of S usually found in cultivated plants (Osvalde 2011). In controlled conditions, moderate to high soil salinity induced up to two-fold Mn accumulation in *A. tripolium* tissues (Karlsons et al. 2008). In contrast, S accumulation in both leaf and root tissues of *A. tripolium* were significantly inhibited by high salinity (Karlsons et al. 2008). It seems that accumulation of extremely high concentration of particular ions in photosynthetic tissues is a typical physiological characteristic of plants native to chemically heterogeneous soils as in sea-affected wetlands. Possible adaptive significance of the phenomenon needs to be proven experimentally, but chemical sequestration of particular ions in leaf cell vacuoles can be suggested as a possible mechanism of tolerance, as already shown for heavy metals (Sharma et al. 2016).

The problem of mycorrhizal vs. non-mycorrhizal plants is especially important for plants in highly heterogeneous environments. While one of the model species used in the present study, *T. maritima*, is considered to be only occasionally mycorrhizal (Harley, Harley 1987) or even non-mycorrhizal (Hildebrandt et al. 2001), it did show significant change of mycorrhizal pattern in roots during the vegetation season. The intensity of colonization was not high, but the presence of arbuscules indicated significant functional activity of the symbiosis. In a previous study it was noted that roots of *T. maritima* contained linear hyphae, vesicles with storage lipid inclusions, as well as arbuscules (Druva-Lūsīte, Levīnš 2010). In conditions of significantly decreased soil salinity in July, intensity of mycorrhizal symbiosis increased up to 17%, while frequency reached 70% (Fig. 16E). It has been argued that low intensity of mycorrhizal symbiosis does not suggest its ineffectiveness (Füzy et al. 2008).

Changes in intensity of mycorrhizal symbiosis during the vegetation season have been associated with differences in growth and development of particular species, rather than with changes in abiotic factors (Bohrer et al. 2004). Direct effect of changes in abiotic conditions (temperature, soil moisture) on mycorrhizal symbiosis can not be excluded (Hildebrandt et al. 2001; Tibbett, Cairney 2007). In the present study, seasonal trends in mycorrhizal intensity in roots indeed followed a species-specific pattern, but some similarities were noticed. Firstly, no direct relationship between soil P concentration and intensity in mycorrhizal symbiosis was found. It is usually assumed that P availability is directly negatively related to mycorrhizal symbiosis, as investment of reduced carbon substances for mycorrhizal growth can be beneficial only in the case of low nutrient availability (Jones et al. 1991). Similarly, fungal colonization parameters did not correlate with soil P in a study with low-alpine herb species (Ruotsalainen et al. 2002). A more reliable indicator of P translocation through mycorrhizal structures to plant could be the presence of arbuscules, but their intensity increased during the season for *P. maritima* and *T. maritima* plants showing no relationship with soil P concentration. Secondly, increased
soil salinity seemed to be related to low level of mycorrhizal colonization, at least, for some species. A. *tripolium*, *G. maritima* and *T. fragiferum* had a low level of mycorrhizal intensity in July, following increased soil salinity in June. In conditions of moderately high soil salinity (EC 10 to 15 dS m$^{-1}$) *G. maritima* plants had constantly very low level of mycorrhization, less than 4% (Druva-Lūsīte et al. 2008), but in the present study mycorrhization intensity increased up to 52% in August when soil salinity decreased to 4 dS m$^{-1}$. Decrease in mycorrhizal intensity can be related to the fact that the respective mycorrhizal fungi may have lower salinity tolerance than their halophytic host plant (Johnson-Green et al. 2001). It seems that mycorrhizal symbiosis in plants from coastal wetlands serves only a minor role in mineral nutrient acquisition. In addition to improvement of general stress tolerance of mycorrhizal plants (Garg et al. 1006) more specific function of mycorrhizal symbiosis can be related to accumulation of excess levels of metal ions, especially, heavy metals (Leyval et al. 1997; Hildebrandt et al. 2007).

The present study was not aimed at complete characterization of changes in photochemistry of photosynthesis of different salt marsh species due to soil temporal heterogeneity. However, the data presented here clearly show that edaphic conditions related to soil inundation with saline water had significant effect on photochemistry of photosystem II of typical coastal wetland species *A. tripolium* and *G. maritima*. Decreased $F_v/F_m$ and $PI$ in leaves of *A. tripolium* in July coincided with the lowest intensity of mycorrhizal colonization, possibly related to increased soil salinity in June. In contrast, *G. maritima* had the lowest fluorescence parameters in June, followed by significant increase in July, when mycorrhizal intensity was lowest. In another study, photosynthetic performance of *G. maritima* in natural conditions was significantly inhibited due to direct damage to photosystem II only when soil salinity increased above 10 dS m$^{-1}$ (Druva-Lūsīte et al. 2008).

In conclusion, the results of this study support the hypothesis that wild plants that are native to habitats with drastically heterogeneous soil conditions have developed effective adaptations as a result of evolution, allowing to grow normally and reproduce within a wide range of mineral concentration conditions, as well as under drastic changes in soil salinity.

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**References**


