Effects of genetic modifications to flax (*Linum usitatissimum*) on arbuscular mycorrhiza and plant performance

Magdalena Wróbel-Kwiatkowska · Katarzyna Turnau · Katarzyna Góralska · Teresa Anielska · Jan Szopa

Abstract

Although arbuscular mycorrhizal fungi (AMF) are known for their positive effect on flax growth, the impact of genetic manipulation in this crop on arbuscular mycorrhiza and plant performance was assessed for the first time. Five types of transgenic flax that were generated to improve fiber quality and resistance to pathogens, through increased levels of either phenylpropanoids (W92.40), glycosyltransferase (GT4, GT5), or PR2 beta-1,3-glucanase (B14) or produce polyhydroxybutyrate (M50), were used. Introduced genetic modifications did not change the degree of mycorrhizal colonization as compared to parent cultivars Linola and Nike. Arbuscules were well developed in each tested transgenic type (except M50). In two lines (W92.40 and B14), a higher abundance of arbuscules was observed when compared to control, untransformed flax plants. However, in some cases (W92.40, GT4, GT5, and B14 Md), the mycorrhizal dependency for biomass production of transgenic plants was slightly lower when compared to the original cultivars. No significant influence of mycorrhiza on the photosynthetic activity of transformed lines was found, but in most cases P concentration in mycorrhizal plants remained higher than in nonmycorrhizal ones. The transformed flax lines meet the demands for better quality of fiber and higher resistance to pathogens, without significantly influencing the interaction with AMF.

Keywords

Mycorrhiza · *Linum usitatissimum* · Transgenic flax · Plant performance

Introduction

Although genetically modified plants (GMPs) have already been accepted by several countries for use in the open field (Liu 2010), there is still a paucity of knowledge on the potential hazards of releasing such organisms into nature, including their potential impact on non-target microbiota (Giovannetti et al. 2005; Liu 2010; Stefani and Hamelin 2010). Arbuscular mycorrhizal fungi (AMF) are among the most common microbiota in soils, and plant dependency on AMF communities is essential and extensive in both natural and disturbed habitats (Van der Heijden et al. 1998; Smith and Read 2008). AMF can be crucial in low-input agriculture, while in strongly fertilized soil they can be totally ineffective or absent (Boyetchko 1996; He and Nara 2007).

As recently reviewed by Stefani and Hamelin (2010), GMPs can alter or not affect mycorrhizal development or colonization rates. Negative effects have been observed in Bt transgenic rice (Ren 2006), corn, and defensin-expressing aubergine (Turrini et al. 2004a). This involved reduced presymbiotic hyphal growth and development of appressoria (Turrini et al. 2004a) and reduced root colonization and spore density in rhizosphere soil (Ren 2006). Over half of the studies showed no effect of Bt transgenic plants on AMF, but to avoid potential risks it nevertheless seems crucial to test every newly developed GMP for its...
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effect on arbuscular mycorrhizal association before released into the field.

Genetically modified (GM) flax lines have been created as a model for studying the role of antioxidants and to improve antifungal resistance and retting efficiency for the production of fiber-rich material (Szopa et al. 2009; Gredes et al. 2010; Skórkowska-Telichowska et al. 2010, Czemplik et al. 2011). Studies on the effect of genetic modifications of flax on plant–AMF interactions are important as GM flax starts to be cultivated in the field and the commercial demand for this crop is large. GM flax lines pose little environmental risk because their seed production and adventitious presence among subsequent crops and yield are relatively easy to control (Jhala et al. 2010). Nevertheless, mechanisms that the plant uses against pathogens can have deleterious effects also on mutualistic symbionts (Whipps 2004). Thus, the aims of the present study were to investigate the effect of different genetic modifications of flax (Linum usitatissimum) on root colonization by Glomus intraradices and compare mycorrhizal dependency of individual lines.

Materials and methods

Five types of transgenic flax (Table 1) were used in the present study. The transgenic lines have been generated from Linola and Nike cultivars and characterized previously (Wróbel et al. 2004; Wróbel-Kwiatkowska et al. 2004; from Linola and Nike cultivars and characterized previously). The transgenic lines have been generated with Ralstonia eutropha genes coding for β-ketothiolase (phbA), acetoacetyl-CoA reductase (phbB), and PHB synthase (phbC) for poly-β-hydroxybutyrate (PHB) synthesis. Two other plant lines (GT4, GT5) derived from Linola cultivar overproduce glycosyltransferase, resulting in the accumulation of proanthocyanin, lignan, phenolic acid, and unsaturated fatty acids in seeds. Plants of an MB type were also obtained, by classical crossing of lines M and B. All these transgenic flax lines show improved resistance to common flax pathogens such as Fusarium oxysporum and/or Fusarium culmorum, which cause diseases such as vascular wilt, corm rot, root rot, or damping-off (Armstrong and Armstrong 1981; Jones 1991).

Culture conditions of mycorrhizal and nonmycorrhizal plants

Seeds of the GM flax lines were germinated on wet filter paper in Petri dishes at approximately 20°C. Five-day old seedlings (n=5 to 9 depending on the line) were transferred into 800 ml Ø 13-cm pots with sterile substratum composed of a mixture of garden soil (Rolex, containing peat, sand, pine bark, compost enriched with organic fertilizers and dolomite), sand, and expanded clay in the ratio 3:3:2 (v/v/v). The substratum was amended with 25 g per kg of rock phosphate (SIARKOPOL, Tarnobrzeg). Prior to seedling transfer, ca. 15 g of AM fungal inoculum containing propagules of G. intraradices UNIJAG.PL24-1 isolated from a grassland close to Inowroclaw (Poland) was introduced below the seedlings. Inoculum was produced in 3-L pots of sterile substratum (sand/expanded clay 2:1, v/v) with Zea mays and Plantago lanceolata as host plants. After 4 months of cultivation, the shoots were harvested and the inoculum was dried for 30 days. Inoculum quality was assessed on P. lanceolata

Table 1 List of transgenic plants investigated

| Transgenic line | Origin of gene | Type of gene and gene construct | Method of gene introduction and selection | References |
|-----------------|----------------|---------------------------------|------------------------------------------|------------|
| W92.40          | Petunia hybrida | CHS, CHI, DFR, triple gene construct, each under 35S promoter | Agrobacterium mediated transformation, PCR, Northern blot | Lorenc-Kukula et al. 2005 |
| GT4, GT5        | Solanum sagarandimum | SsGT1 gene under nap promoter | Agrobacterium mediated transformation, PCR, Northern blot | Lorenc-Kukula et al. 2009 |
| M50             | Ralstonia eutropha | phbA, phbB, phbC, triple gene construct, phbB and C genes under 35S, phbA under 14-3-3 promoter | Agrobacterium mediated transformation, PCR, Northern blot | Wróbel et al. 2004 |
| B14             | Solanum tuberosum | beta-1,3-glucanase gene under 35S promoter | Agrobacterium mediated transformation, PCR, Northern blot | Wróbel-Kwiatkowska et al. 2004 |
seedlings grown for 6 weeks; inoculation resulted in over 90% AMF colonized root length.

The pots were placed in Sunbags (Sigma) and flax plants were grown under greenhouse conditions: 25/18°C maximum/minimum average air temperatures, 12 h light supplementation if needed (200 μmol·m⁻²·s⁻¹ PAR). Plants were irrigated weekly with distilled water and once per 3 weeks with Long Ashton solution (Davies et al. 1992). After 3-month growth, when plants started to form flowers, photosynthesis was measured and plants were harvested. Length of roots and shoots and dry weight of shoots were estimated (roots were used for estimation of mycorrhizal parameters).

Mycorrhizal colonization

Roots were washed, cleared in 10% KOH for 24 h at room temperature and, after rinsing in tap water, acidified for 1 h in 5% lactic acid then stained 24 h at room temperature in 0.05% aniline blue in lactic acid, in order to visualize the fungal structures inside the roots. One-centimeter root pieces were mounted on slides in lactoglycerol (approx. 45 root pieces per plant) and mycorrhizal parameters in root systems were assessed according to Trouvelot et al. (1986; http://www2.dijon.inra.fr/mychintec/Mycocalc-prg/download.html): frequency of mycorrhiza presence (F%), intensity of mycorrhizal colonization (M%), and arbuscule abundance (A%). Images were acquired using a NIKON Eclipse E800 microscope equipped with DIC, a digital image capture system and a computer image analysis system.

Plant growth assessment

Shoot length was measured, then material was dried for 2 days at 40°C and weighed (Radwag analytical balance, WPA 60/c/1). Mycorrhizal dependency (Md) of plants for growth (mass, shoot length) as well as performance index (see below) was determined according to the equation given by van der Heijden and Sanders (2002)

P and Zn concentrations in plants

Dried, grinded, and homogenized flax shoots were digested in 10-mL concentrated (65%) nitric and perchloric acids (Ultranal, POCh) according to Pilegaard (1979). Residues were dissolved in 10 mL 0.2% nitric acid, filtered, and diluted if necessary. Zinc concentrations were analyzed by flame atomic absorption spectrometry using AAAnalyst 200, Perkin-Elmer. Nitric acid was used as a blank and Ulva lactuca BCR-279 as a standard reference material (Promochem GmbH; Zn=51.3 mg kg⁻¹). Concentrations of phosphorus were determined by flow injection analysis system (FIA) coupled with flow colorimeter (FIA compact, MLE GmbH).

Fluorescence measurements

Chlorophyll a fluorescence transients of intact leaves were measured with a plant efficiency analyzer fluorometer (Hansatech Instruments, UK) and recorded as described by Strasser et al. (1995). Each transient was analyzed according to the OJIP test (Strasser et al. 2004, 2007) and the following biophysical parameters (referring to time point zero) were calculated to characterize PSII behavior: (1) specific energy fluxes per reaction center for absorption (ABS/RC); (2) electron trapping (TRo/RC), electron dissipation (Do/RC), and electron transport (To/RC); (3) the flux ratios or yields, i.e., the maximum quantum yield of primary photochemistry (ϕPo = TRo/ABS); (4) the efficiency with which a trapped exciton can move an electron into the electron transport chain further than Qψ (ψo = ETo/TRo), the quantum yield of electron transport (ϕEo = ETo/ABS); (5) and the performance index (PI), which provides quantitative information about the state of plants and their vitality. The latter incorporates three independent expressions: (1) ratio of reaction center chlorophylls and the total chlorophyll of PSII; (2) an expression related to primary photochemistry ϕoPo/(1−ϕoPo); and (3) an expression related to electron transport ψo/(1−ψo) (Oukarroum et al. 2007).

Data analysis

Data from mycorrhizal and nonmycorrhizal plants were compared using the t test or were transformed with decimal logarithms prior to analysis to meet statistical assumptions and compared using ANOVA, STATISTICA (ver. 7.0) software (significance level P<0.05).

Results

AMF colonization

Arbuscular mycorrhiza of *Arum* type with well-developed arbuscules were observed in all inoculated flax plants. These structures were not found in any of the non-inoculated samples. The frequency of mycorrhiza (F%), reflecting propagule density in the samples, ranged from 93% to 100% and showed the smallest differences between root samples among studied parameters. The intensity of mycorrhizal colonization (M%) ranged from 40% to 65%. Significantly higher M% values were noted in the case of lines B14 and W9240, while other transformed lines from the cultivars Linola and Nike were similar (Fig. 1). The largest differences between samples involved arbuscule abundance (A%) in roots which was highest in the two transgenic lines W9240 and B14. The other lines derived from cv. Linola showed similar values for arbuscule abundance. In contrast,
the GM lines derived from cv. Nike either had higher (B14) or significantly lower (M50) A% (Fig. 1). Roots were free of parasites or nematodes.

Plant growth and performance

Mycorrhizal plants of cv. Nike and Linola were characterized by increased total biomass and root weight in comparison to nonmycorrhizal controls (Fig. 2). Smaller but still significant differences ($P=0.049$) were found in root or shoot weight between mycorrhizal and nonmycorrhizal GM lines. There were no significant differences between shoot weight of mycorrhizal transgenic lines and of their parent cultivars. In the case of root weight, statistically significant differences were found only between GT4 and cv. Linola (Fig. 2). Among nonmycorrhizal plants, higher shoot weight was found in GT4 than in cv. Linola and lower weight in B14 than in cv. Nike, while the root weight of W92.40 and GT4 were 3 and 2 times higher than in cv. Linola, respectively, and in M50 and GT5 were over 1.5 and 2 times higher than in cv. Nike. Md for growth was higher in cv. Nike (30%) than in cv. Linola (18%). Transformation of these two cultivars resulted in decreased values of growth Md in comparison to the parent cultivars. Negative values were found in the lines W92.40 (−18%), GT4 (−14%), and GT5 (−12%), while the Md value of B14 was ca. 10% lower than in cv. Nike.

P concentration in flax shoots was significantly higher in mycorrhizal plants than in nonmycorrhizal ones, both in GM lines and parent cultivars, except for line M50 (Fig. 3a). On the contrary, Zn concentration depended on the flax cultivar, being significantly increased in cv. Linola, but not in cv. Nike. No significant differences were found in two modified lines, B14 and W92.40 (Fig 3b). No significant differences in Md concerning P and Zn were found between GM and non-GM lines. No differences were found in flowering time of the GM and non-GM lines or between mycorrhizal and nonmycorrhizal plants.

The measure of plant performance, Pl, revealed differences between transgenic lines and the parent cultivars. PI of mycorrhizal plants of cultivars Nike and Linola were significantly higher than in nonmycorrhizal ones (Fig. 3). Md estimated on the basis of plant performances (PI) in these two cultivars were 12% and 26%, respectively. Among transformed lines, a similar situation was found only in the case of W92.04 (Md=21%); a negative effect of mycorrhization on PI was not significant in the case of GT4 (Md=−17%), while in other cases the values were similar in mycorrhizal and nonmycorrhizal plants (Fig. 4). In addition, PI increase in non-transformed cultivars following mycorrhization was associated with statistically significant differences in several PSII parameters, such as ABS/RC, TRo/RC, DIo/RC, and To/RC, the maximum quantum yield of primary photochemistry ($\varphi_{Po}=TRo/ABS$), and the quantum yield of electron transport ($\varphi_{Eo}=ETO/ABS$) (data not shown). This was not the case for transgenic lines where PI in mycorrhizal plants was not significantly different from nonmycorrhizal ones (GT4, M50, B14, and GT5).
Discussion

AMF root colonization in flax has been previously described (Dickson et al. 2003; Vierheilig et al. 2008) and effects of the symbiosis on non-modified flax have been the subject of both field (Thingstrup et al. 1998; Olsson et al. 1999) and laboratory (Püschel et al. 2008; Rydlova et al. 2010) studies. Mycorrhizal effects have been shown on flax growth, due to increased uptake of P and Zn by fungal mycelium (Thompson 1994; Thingstrup et al. 1998; Smith et al. 2004), on fiber weight, seed production, and fatty acid composition of seed oils (Rydlova et al. 2010), on drought resistance (Drue and Schonbeck 1992; von Reichenbach and Schonbeck 1995) and on resistance to plant pathogens (Dugassa et al. 1996). The main aim of the present study was to determine whether genetic transformation of flax by different resistance-related genes affects AMF and/or their effects on host performance. Transformation did not change the degree of mycorrhizal colonization; no morphological differences were found between transformed and non-transformed flax lines and arbuscules, which are indicative of a functional symbiosis, were well developed in all cases except line M50. The lines B14 and W92.40 showed even higher arbuscule abundance in comparison with the parent cultivars. These observations are in agreement with most of previously published data showing no negative effect on fungal growth in plants overexpressing defense-related genes (Liu 2010). The transformations in the present flax lines increase resistance to pathogenic fungi via increased phenylpropanoid levels (W92.40) (Lorenc-Kukula et al. 2005), antioxidant capacity (W92.40, GT) and overproduction of glycosyltransferase (GT) (Lorenc-Kukula et al. 2004) and β-1,3-glucanase ectopic expression of a potato β-1,3-glucanase gene in flax (B14). Since there is some similarity between the soil-borne pathogens and AMF in their resources such as space and photosynthate within the root, such defense "pathways" could be potentially deleterious to both groups of fungi (Whipps 2004). However, as reviewed by Dumas-Gaudot et al. (2000) and Khan et al. (2010), AMF are capable themselves of inducing hydrolytic enzymes and pathogenesis-related proteins without being affected. For example, AMF are not inhibited by Class I chitinases (Arlorio et al. 1991; Salzer et al. 1997; Vierheilig et al. 1993) that are deleterious to pathogenic fungi such as Rhizoctonia solani. Similarly, AMF are not affected by an antifungal protein that targets Verticillium albo-atrum or Botrytis cinerea in transformed aubergine plants (Turrini et al. 2004a, b). This differential targeting of pathogenic and mycorrhizal fungi could be due to mechanisms of recognition specificity (Vierheilig et al. 2008; Khan et al. 2010; Stefani and Hamelin 2010) but this issue requires further in-depth research.

The effect of genetically modified crops on mycorrhizal fungi has so far been mostly addressed by analysis of mycorrhizal colonization levels which, as mentioned above, are largely unaffected. However, the present data indicate that, although growth of the mycorrhizal transformed flax plants was not different from the parent lines, transformation may alter plant responses to AMF which could be relevant for biomass production and crop quality under agricultural conditions. While no significant differences were observed in Md concerning P and Zn, G. intraradices was less...
effective in stimulating growth or photosynthetic performance and, in one case P uptake (M50), by transformed as compared to parent flax plants. The magnitude of mycorrhizal responses can be influenced not only by the plant but also environmental factors (Smith and Gianinazzi-Pearson 1988, Smith and Read 2008, Jakobsen et al. 2002). In the GM flax lines, the lower mycorrhizal dependency parameters in reference to plant growth and photosynthesis efficiency resemble the situation between modern varieties or landraces and their ancestors (Hetrick et al. 1992, 1993; Boyetchko 1996), and both might be less adapted to an agriculture turned towards low fertilizer inputs, especially in phosphorus and nitrogen, where AMF can be the key players in establishment of ecosystem function and diversity (Thompson 1994, van der Heijden et al. 1998, Klironomos et al. 2000).

In conclusion, it was shown that the genes introduced into flax lines to increase resistance against pathogens did not have a negative effect on the colonization of flax by AMF but may affect some plant responses to the symbiosis. Plant growth and photosynthetic activity were not significantly improved by G. intraradices in the transformed as compared to parent lines, but P nutrition was still enhanced in all but one which may be promising for better survival of seedlings due to better nutrition of seeds. The fact that the transformed flax lines, which meet the demands for better quality of fiber and higher resistance to pathogens, do not appear to be negatively affected in their interactions with AMF should make them suitable for use in agriculture.

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