Elevated phosphorus impedes manganese acquisition by barley plants

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The occurrence of manganese (Mn) deficiency in cereal crops has increased in recent years. This coincides with increasing phosphorus (P) status of many soils due to application of high levels of animal manure and P-fertilizers. In order to test the hypothesis that elevated P may lead to Mn deficiency we have here conducted a series of hydroponics and soil experiments examining how the P supply affects the Mn nutrition of barley. Evidence for a direct negative interaction between P and Mn during root uptake was obtained by on-line inductively coupled plasma mass spectrometry (ICP-MS). Addition of a pulse of KH₂PO₄ rapidly and significantly reduced root Mn uptake, while a similar concentration of KCl had no effect. Addition of a P pulse to the same nutrient solution without plants did not affect the concentration of Mn, revealing that no precipitation of Mn–P species was occurring. Barley plants growing at a high P supply in hydroponics with continuous replenishment of Mn²⁺ had up to 50% lower Mn concentration in the youngest leaves than P limited plants. This P-induced depression of foliar Mn accelerated the development of Mn deficiency as evidenced by a marked change in the fluorescence induction kinetics of chlorophyll a. Also plants growing in soil exhibited lower leaf Mn concentrations in response to elevated P. In contrast, leaf concentrations of Fe, Cu, and N increased with the P supply, supporting that the negative effect of P on Mn acquisition was specific rather than due to a general dilution effect. It is concluded that elevated P supply directly interferes with Mn uptake in barley roots and that this negative interaction can induce Mn deficiency in the shoot. This finding has major implications in commercial plant production where many soils have high P levels.

Keywords: manganese deficiency, phosphorus, barley, chlorophyll fluorescence, ICP-MS, zinc

INTRODUCTION

Phosphorus (P) and manganese (Mn) are essential plant nutrients. Phosphorus is a constituent of nucleic acids, phospholipids and ATP, and is also involved in the regulation of key metabolic pathways (Yang and Finnegan, 2010; Byrne et al., 2011; Yao et al., 2011). Manganese (Mn) primarily acts as an activator of many enzymes and may in many cases be substituted by other nutrients, especially magnesium. However, in at least three cases Mn is irreplaceable (Burnell, 1988; Hansch and Mendel, 2009). These include (i) oxalate oxidase, catalyzing the conversion of oxalate and oxygen to hydrogen peroxide and carbon dioxide (Requena and Bornemann, 1999); (ii) superoxide dismutase (MnSOD), converting the oxygen free radical O₂⁻ to H₂O₂ and O₂ (Alscher et al., 2002; Poage et al., 2011); and (iii) the oxygen evolving complex (OEC) of photosystem II, placed in the thylakoid membranes of the chloroplasts and containing Mn bound in a 4-atom cluster (Ono et al., 1992).

Manganese deficiency is regularly observed in plants growing on soils with low Mn availability, but the frequency of the problem has increased during recent years (Hebbern et al., 2005). During the same period, soil-P levels have increased due to high application of animal manure and P-fertilizers (Kyllingsbæk and Hansen, 2007; Rejnevedt et al., 2010; MacDonald et al., 2011). It can therefore be hypothesized that the increased frequency of Mn deficiency may be influenced by the increased soil-P levels. Both P and Mn ions are rather immobile in soils implying that factors such as root length and root architecture, together with rhizosphere processes, have a major impact on their plant availability (Gahoonia and Nielsen, 1997; Gherardi and Rengel, 2004; Raghothama and Karthikeyan, 2005). The presence of Mn substantially inhibited root hair elongation in Arabidopsis, whereas Mn-deficient seedlings displayed stimulated root hair development (Yang et al., 2008). Also P starvation increases the number of lateral roots and root hair abundance (Gahoonia et al., 1997; Zhu et al., 2010; Miura et al., 2011). Simultaneously, nitrate uptake is inhibited by P starvation leading to excess uptake of cations relative to anions and acidification of the rhizosphere (Hedley et al., 1982; Schjoerring, 1986; Végh et al., 2009). The absence of these adaptive root responses in soils with high P status will reduce the ability of roots to acquire Mn.

Phosphorus interacts with other micronutrients than Mn, e.g., zinc (Zn), iron (Fe), and copper (Cu; Murphy et al., 1981; Broadley et al., 2010; Perez-Novoa et al., 2011). Phosphate fertilization of plants with low Zn status may lead to visible Zn deficiency symptoms, a phenomenon referred to as P-induced Zn deficiency (Cakmak and Marschner, 1987). It is not clear whether decreased Zn uptake or dilution following increased biomass production are the primary reason for P-induced Zn deficiency (Singh et al., 1988; Gianquinto et al., 2000; Zhu et al., 2001; Li et al., 2003). Genes encoding high-affinity orthophosphate transporters are up-regulated under Zn deficiency (Huang et al., 2000) resulting
in higher P uptake, potentially accompanied by P toxicity which can be alleviated by addition of Zn (Marschner and Cakmak, 1986; Silber et al., 2002).

A few cases of negative influence of high soil-P availability on Mn uptake have been reported, tentatively assigned to the reduced formation of cluster roots, root-induced changes in the rhizosphere or negative interactions with P during translocation from roots to shoots (Neilsen et al., 1992; Zhu et al., 2002; Shane and Lambers, 2005). However, the exact mechanisms behind the Mn–P interaction are still unknown.

The objective of the present study was to examine interactions between P and Mn in barley plants. A series of experiments were conducted with plants exposed to different Mn and P treatments in soil or hydroponics. The results show a negative effect of high P levels on Mn accumulation and it is suggested that P interferes directly with Mn at the uptake and/or translocation level.

**MATERIALS AND METHODS**

**PLANT CULTIVATION**

**Hydroponic experiments**

Seeds of the Mn-efficient genotype Vanessa and the inefficient genotype Antonia were germinated at 21°C in vermiculite. After 4 days, uniform seedlings were selected and transferred to opaque 4 l cultivation units, each holding four plants. The units were filled with aerated nutrient solution having the following composition: 0.2 mM KH₂PO₄, 0.2 mM K₂SO₄, 0.3 mM MgSO₄·7H₂O, 0.1 mM NaCl, 0.3 mM Mg(NO₃)₂·6H₂O, 0.9 mM Ca(NO₃)₂·4H₂O, 0.6 mM KNO₃, 0.8 μM Na₂MoO₄·2H₂O, 0.7 μM ZnCl₂, 0.8 μM CuSO₄·5H₂O, 2 μM H₂B₂O₇, 1 μM NiSO₄·6H₂O, 50 μM Fe(NO₃)₃·9H₂O, and 10 mM EDTA. Before use, the macronutrient stock solutions were purified with Chelex-100 resin (Sigma-Aldrich) in order to remove interfering on the availability of the ICP-MS instrument. Starting at least 7 days before the experiments, the nutrient solution was exchanged when they had attained growth stage 32 (Zadoks et al., 1974). The roots were washed in double deionised water for 5 min, followed by drying of shoots and roots.

**Soil experiments**

Soil was collected in a field cropped with cereals for more than 40 years without application of any organic or inorganic P or K fertilizers (Schjoerring and Nielsen, 1987). The NaHCO₃-extractable P content of the soil was 10.8 mg P kg⁻¹ dry soil and the pH 5.1 (0.01 M CaCl₂). The soil was mixed with quartz sand and sphagnum (2:3:1 v/v) and 0.3 g N kg⁻¹ mixed soil added in the form of NK-fertilizer with S, Mg, and B (Kemira 19-0-15; 50% nitrate-N, 50% ammonium-N). The pH (0.01 M CaCl₂) of the mixed soil was adjusted to 6.0 by mixing the soil with 1.25 g CaCO₃ kg⁻¹ soil. Four P levels were produced by addition of 0, 13.5, 41.4, or 83.2 mg KH₂PO₄ kg⁻¹ soil. In order to achieve similar K levels for all treatments, the K levels on Mn accumulation and it is suggested that P interferes directly with Mn at the uptake and/or translocation level.

| Table 1 | Daily addition rates of phosphorus and manganese per cultivation unit with four plants in the hydroponic experiment. |
|---------|-----------------------------------------------------------------------------------------------------|
| Nutrient | Treatment | Daily addition (μg) | Concentration after addition (μM) |
| Mn      | Low       | 0.33*              | 0.0015                           |
|        | Medium    | 3.30               | 0.015                            |
|        | High      | 9.90*              | 0.045                            |
| P       | Low       | 372                | 3                                |
|        | Medium    | 3720               | 30                               |
|        | High      | 18600              | 150                              |

*After 20 days of growth the daily Mn addition was abolished for the low treatment whereas the daily Mn addition for the high treatment was increased to 19.80 μg corresponding to a Mn concentration after addition of 0.090 μM.

**PLANT ANALYSIS**

**Chlorophyll a fluorescence**

The effect of Mn deficiency on PSII was recorded by measurement of the fluorescence induction kinetics of chlorophyll a using a Handy PEA (Plant Efficiency Analyzer, Hansatech Instruments, King’s Lynn, UK). A saturating light flash of 3000 μmol photons m⁻² s⁻¹ was applied for 2 s to leaves which had been dark adapted for 25 min. The fluorescence results were parameterized to Fv/Fm using the Handy PEA software (version 1.30).

**On-line measurements of nutrient uptake by inductively coupled plasma mass spectrometry**

Plants were pre-grown as described above for 25–53 days depending on the availability of the ICP-MS instrument. Starting at least 7 days before the experiments, the nutrient solution was exchanged daily. In addition, 24 h before the experiments, plants were pre-equilibrated to the uptake solution containing a reduced micronutrient supply composed of 0.01 μM Na₂MoO₄·2H₂O, 0.01 μM ZnCl₂, 0.01 μM CuSO₄·5H₂O, 0.01 μM H₂B₂O₇, 0.01 μM NiSO₄·6H₂O, 0.01 μM Fe(NO₃)₃·9H₂O, and 100 mM MnCl₂. This procedure was necessary in order to reduce the effects of ion-exchange processes between root and nutrient solution.

The experiments were conducted with plants growing in 300 ml vials (Bie & Berntsen, Denmark) and exposed to a 250 to 280-μmol m⁻² s⁻¹ photon flux density and 20°C (16/8 h) day/night.
temperature regime. Each series of measurements lasted up to 8 h and embraced one vial with one plant and one without plant (control). Each of the two vials was analyzed every 12 min during the experimental period. Pulses of 150 or 500 μM KH$_2$PO$_4$, or 200 μM KCl, were added at specified time points and the rate of P and Mn depletion from the nutrient solution were recorded by on-line ICP-MS (Agilent 7500c, Agilent Technologies, Manchester, UK). A multi-channel peristaltic pump continuously recirculated nutrient solution between an auto sampler vial (50 ml vial, Product no. 02-6100N, CM-LAB, Denmark) and the cultivation unit with plants. The tube inlet and outlet were vertically displaced in the auto sampler tube providing a steady-state volume of approx. 7.5 ml due to a flow rate difference between inlet and outlet. During sampling, a mixture of 1.75% HNO$_3$, 0.2% HF, and 10 mg l$^{-1}$ $^{45}$Sc was continuously added to the nebulizer (split-ratio 1:20) in order to reduce memory effects and to correct for drift and argon plasma instability. In separate experiments, continuous analysis of control nutrient solution without plants documented that no significant interferences caused by, e.g., precipitation and adsorption of Mn to surfaces of the experimental system were occurring during the experiments (Pedas et al., 2005).

**Nutrient composition of plant tissue**

Following freeze drying (Christ Alpha 2–4; Martin Christ GmbH), milling, and digestion, leaf and root samples were analyzed by ICP-MS as previously described (Hansen et al., 2009). For every 40 samples, four blanks without plant material and four with certified reference material (apple leaf, standard reference material 1515; US Department of Commerce, National Institute of Standards and Technology, Gaithersburg, MD, USA) were included. For total-N, determination 4 mg of dried plant material was weighed into tin capsules and analyzed in a system consisting of an ANCA-SL Elemental Analyzer coupled to a 20–20 Tracermass Mass Spectrometer (Europa Scientific Ltd., Crewe, UK).

**Root hair density**

Representative root samples in terms of size, root geometry, and thickness were analyzed with respect to root hair density using a scanner (ScanJet I1cx) and DT-Scan Software (Delta-T Devices, Cambridge, England) as described by Gahoonia et al. (2000).

**RESULTS**

**INTERACTIONS BETWEEN Mn AND P DURING UPTAKE BY ROOTS**

The influence of P on Mn acquisition was assessed in a series of on-line uptake experiments using ICP-MS. The uptake was carried out with the Mn-efficient genotype Vanessa growing at Mn concentrations <70 nM, i.e., mimicking the low Mn concentration in the rhizosphere soil solution of well-tilerated agricultural soils. All plants were at the time of the experiments in a healthy nutritional state with $F_{v}/F_{m}$ values of approximately 0.8, close to the theoretical optimum of 0.83 (Kriedemann et al., 1985). Prior to the addition of Mn/P pulses, plant roots had been exposed to the uptake solution for 24 h which resulted in Mn and P concentrations below 10 nM and 1 μM, respectively.

Plant roots receiving a Mn pulse in the absence of a simultaneous P pulse steadily depleted the nutrient solution for Mn (Figure 1). However, when the Mn pulse was accompanied by addition of 150 μM KH$_2$PO$_4$, the Mn uptake rate over the first h after the pulse decreased substantially from 7.8 to 1.7 nmol g$^{-1}$ DW h$^{-1}$ (Figure 1). Addition of a similar P pulse to the same nutrient solution without plants did not affect the concentration of Mn, revealing that no precipitation of Mn–P species was occurring (data not shown). Calculations using Geochem-EZ (version 1.0) showed that no uncharged Mn–P complexes, e.g., MnHPO$_4^{2-}$, were likely to form under the condition given and that >99.5% of the Mn was expected to be present as the free Mn$^{2+}$ ion (data not shown).

Tripling the P pulse to approximately 500 μM KH$_2$PO$_4$ completely inhibited the Mn uptake (Figure 2). Application of a similar P pulse, i.e., 500 μM KH$_2$PO$_4$ to only half of the roots in a split-root system instantaneously stopped the Mn uptake in roots exposed to extra P, while that in the other part of the root system was not affected (Figure 3). Approximately 2 h after addition of the P pulse, the Mn uptake resumed (Figure 3). When the KH$_2$PO$_4$ pulse was replaced with 200 μM KCl, no detectable decline in the Mn uptake rate occurred showing that the recorded effect was specific for P (Figure 4).

**PLANT GROWTH AND ELEMENTAL COMPOSITION AS AFFECTED BY Mn–P INTERACTIONS**

No visual deficiency symptoms were observed in any of the treatments (data not shown). The Mn treatments did not result in any clear differences in biomass production in the hydroponic experiments (Table 2). This was the case even though plants growing at the low Mn level became clearly Mn deficient as evidenced by a decline in the $F_{v}/F_{m}$ value to below 0.6 (Figure 5). Biomass production in the medium P treatment was significantly higher than that in the low P treatment and the root/shoot ratio decreased about 50% with increasing P supply showing preferential shoot growth in response to extra P (Table 2). In both the Mn-inefficient genotype Antonia
The elemental composition of the fully expanded youngest leaf was analyzed in order to reveal the influence of the interactions between P and Mn on the nutrient supply to growing tissues. Plants growing in hydroponics with frequent replenishment of Mn were used to avoid indirect dilution effects associated with formation of depletion zones in the rhizosphere soil when biomass production increases in response to elevated P. The P concentration in the youngest leaf increased significantly \((P < 0.05)\) with the P level in the root medium (Figures 6C,F). In the low Mn treatment, the Mn concentration in the youngest leaf decreased below 15 \(\mu\text{g g}^{-1}\) at low P supply and further down below 10 \(\mu\text{g g}^{-1}\) at medium and high P supply (Figures 6A,D). Overall, the Mn concentration in the youngest leaf was in both genotypes negatively correlated with the P supply \((P < 0.05)\). Conversely, the Fe and Cu concentrations in the youngest leaf were only marginally affected by increasing P supply (Figure A1 in Appendix), while the corresponding N concentration increased (Figure A2 in Appendix), supporting that the negative interaction between Mn and P concentrations was specific rather than due to a general dilution effect. The Zn concentration in the youngest leaf decreased with increasing P supply as was the case for Mn (Figures 6B,E). At low P supply, the Zn concentration in the youngest leaf was up to 70% higher than at medium and high P supply. The negative effect of P supply on the Zn concentration was most pronounced at low Mn supply and almost disappeared at ample Mn supply (Figures 6B,E).

Increasing P supply to soil-grown plants of the genotype Antonia resulted in more than two-fold increase in shoot weights (Figure 7A) and three-fold increase in foliar P concentrations (Figure 8B). Conversely, root hair density decreased 35% (Figure 7B). Leaf Zn and Mn concentrations were negatively affected by increasing P supply, decreasing 37 and 29%, respectively, while the Fe and Cu concentrations increased 42 and 71%, respectively (Figure 8A).
the P concentration, ranging from 75% inhibition at 150 μM P (Figure 1) to complete abolishment following addition of 500 μM P (Figure 2). Addition of a similar P pulse to the same nutrient solution without plants did not affect the concentration of Mn, revealing that precipitation of Mn–P species were not occurring (data not shown). It has been hypothesized that Mn $^{2+}$ may form an ion-par with HPO$_4^{2-}$ resulting in the uncharged soluble complex $\text{MnHPO}_4^0$ (Lindsay, 1979). Using Geochem-EZ (version 1.0) we therefore examined to which extent inorganic soluble Mn–P complexes were likely to form after addition of P. According to the Geochem calculations, no uncharged Mn–P complexes were likely to form and $>99.5\%$ of the Mn was present as the free Mn $^{2+}$ ion (data not shown). The observed decline in Mn uptake appeared to increase with the P concentration, ranging from 75% inhibition at 150 μM P (Figure 1) to complete abolishment following addition of 500 μM P (Figure 2). Addition of a similar P pulse to the same nutrient solution without plants did not affect the concentration of Mn, revealing that precipitation of Mn–P species were not occurring (data not shown). It has been hypothesized that Mn $^{2+}$ may form an ion-par with HPO$_4^{2-}$ resulting in the uncharged soluble complex $\text{MnHPO}_4^0$ (Lindsay, 1979). Using Geochem-EZ (version 1.0) we therefore examined to which extent inorganic soluble Mn–P complexes were likely to form after addition of P. According to the Geochem calculations, no uncharged Mn–P complexes were likely to form and $>99.5\%$ of the Mn was present as the free Mn $^{2+}$ ion (data not shown). The observed decline in Mn uptake appeared to increase with

**DISCUSSION**

Previous studies of interactions between Mn and P have been carried out with plants growing in soil (Neilsen et al., 1992; Shane and Lambers, 2005). In the present study we primarily used a hydroponic growth system to examine how interactions between Mn and P affect growth, Mn uptake, and nutrient concentrations in root and leaf tissues. One of the advantages associated with nutrient solutions instead of soil is that effects of P addition on Mn uptake by the roots can be measured dynamically. Using a very sensitive on-line ICP-MS system, enabling analysis in the nM range, we have here demonstrated that Mn uptake respond instantaneously and specifically to changes in the external P concentration (Figures 1–4). The decline in Mn uptake appeared to increase with

**Table 2 | Total biomass (shoot + root) and root/shoot ratios of barley genotypes Antonia and Vanessa grown in a hydroponic system at different Mn and P levels.**

| P level | Biomass production (g DW) | Root/shoot ratio |
|---------|---------------------------|----------------|
|         | Low Mn | Medium Mn | High Mn | Low Mn | Medium Mn | High Mn |
| Antonia |        |          |         |        |          |         |
| Low     | 5.5    | 4.8      | 5.1     | 0.41   | 0.38     | 0.38    |
| Medium  | 7.4    | 9.0      | 7.3     | 0.21   | 0.23     | 0.19    |
| High    | 7.6    | 8.2      | 7.0     | 0.20   | 0.21     | 0.21    |
| Vanessa |        |          |         |        |          |         |
| Low     | 6.6    | 6.9      | 7.5     | 0.40   | 0.40     | 0.43    |
| Medium  | 10.3   | 11.8     | 12.0    | 0.21   | 0.23     | 0.21    |
| High    | 9.4    | 12.5     | 10.9    | 0.19   | 0.22     | 0.19    |

Values are means ($n=3$). Values followed by different letters are significantly different ($P < 0.05$).

**Figure 5** Chlorophyll $a$ fluorescence measurements of hydroponically grown plants receiving different Mn and P levels. $F_v/F_m$ values are shown for the Mn-inefficient genotype Antonia (A,B) and the efficient genotype Vanessa (C,D). Inserts (A) and (C) show $F_v/F_m$ values depending on the Mn level summarized across the P levels. Inserts (B) and (D) show $F_v/F_m$ values for plants grown at low Mn supply and increasing P additions. Each data point represents the mean ± SE ($n=6–18$).
application of a P pulse is therefore unlikely to reflect changes in Mn speciation but rather occurred due to a direct interaction between Mn$^{2+}$ and H$_2$PO$_4^-$ during Mn uptake.

Plants acquire Pi by active uptake into the epidermal and cortical cells of the root via proton–Pi symporters of the PHT1 family (Raghothama, 2000). The high-affinity barley Pi transporter HvPHT1;1 is highly expressed in root segments with dense root hairs, is localized to the plasma membrane and shows preferential selectivity for Pi relative to SO$_4^{2-}$ and NO$_3^-$ (Preuss et al., 2011). Uptake of Pi via this transporter leads to a depolarization of the membrane potential (Preuss et al., 2011) which hypothetically may affect Mn$^{2+}$ uptake. In addition, the accompanying proton uptake
may lead to a transient alkalinisation of the apoplastic solution around the root cells which potentially can promote Mn$^{2+}$ adsorption to negative charges in the cell walls and limit the concentration of Mn$^{2+}$ at the uptake sites.

The external P concentration may affect different ion transporters via phosphorylation/dephosphorylation reactions (Bayle et al., 2011). In yeast, protein phosphatase 2B has been shown to be essential for prevention of Mn entry into the cells (Farcasanu et al., 1995). Transporter gene families that have been implicated in Mn uptake by plant roots include natural resistance-associated macrophage protein (Nramp) transporters and zinc-regulated transporter/iron-regulated transporter (ZRT/IRT1)-related protein (ZIP) transporters (Pittman, 2005; Pedas et al., 2008; Cailliatte et al., 2010). None of these are presently known to interact with P. However, an alternative pathway for Mn uptake was identified in yeast where the high-affinity H$^+$-P$^{-}$ co-transporter Pho84 was shown also to mediate low-affinity Mn transport (Jensen et al., 2003). Barley and *Arabidopsis* have various P transporters with a significant similarity to Pho84 (Leggewie et al., 1997). The *Arabidopsis* Pho1 transporter shares extensive amino acid homology with the yeast Pho84 transporter and is involved in xylem loading and long-distance transfer of Pi from the root to the shoot (Hamburger et al., 2002). Overexpression of PHO1 in *Arabidopsis* leaves has recently been shown to lead to a dramatic efflux of Pi out of cells and into the xylem vessel, revealing a crucial role for PHO1 in Pi efflux (Stefanovic et al., 2011). A potential link between phosphate efflux and Mn acquisition remains to be investigated.

A split-root experiment showed that the initial rapid decline in Mn uptake in response to P addition only occurred in roots exposed to the P pulse, while the other root part continued to absorb Mn with the same rate as before (Figure 3). Thus, systemic feed-back signals in the root system, or between roots and shoots, were not responsible for the Mn–P interaction. However, after approximately 2 h, the Mn uptake started to recover again in the roots exposed to the P pulse (Figure 3). This gradual acclimation may involve long-distance signaling affecting P acquisition and translocation between roots and shoots (Schachtman and Shin, 2007; Liu et al., 2009).
MicroRNAs play an important role in the regulation of stress responses in plants and some miRNAs respond both to changes in P as well as Mn availability (Valdés-López et al., 2010).

**PHYSIOLOGICAL EFFECTS OF Mn-P INTERACTIONS**

Measurements of fluorescence induction kinetics of chlorophyll a demonstrated that the low Mn treatment induced a decline in Fv/Fm values (Figure 5), showing that the plants were suffering from Mn deficiency although no visual symptoms were present (Hebborn et al., 2009; Husted et al., 2009). The decline in Fv/Fm values for plants growing at low Mn levels was dependent on the supply of P, occurring faster for plants growing at medium or high P levels compared to low P (Figures 5B,D). Ample P supply thus reduced Mn acquisition, translocation and/or the amount of Mn functionally available for photosynthesis. The ability to perform state transitions and the amount of the photosynthetic key protein PsbA (D1 protein) have previously been shown to be influenced by the plant Mn status (Husted et al., 2009). The Mn concentration in the youngest leaf of both genotypes growing in low Mn treatment was approximately 12 μg Mn g DW⁻¹ (Figures 6A,D) which is below the critical level for Mn (Reuter et al., 1997). The elemental analysis of the plant tissue thus confirmed the Chl a fluorescence measurements and both approaches documented the risk of inducing Mn deficiency in the youngest leaf by increasing P levels.

The Mn concentration in plants results from the integration of two dynamic processes, dry matter accumulation and Mn absorption. If dry matter accumulation increases at a faster rate than Mn absorption it will result in a decrease in the Mn concentration. This inverse relationship between growth and mineral concentration is termed the dilution effect (Jarrell and Beverly, 1981). The Mn concentration in the youngest leaf was clearly negatively affected by increasing P supplies (Figures 6A,D). However, for the following reasons we consider it unlikely that this decrease was due to a general dilution effect: Firstly, the plants were grown in hydroponics with continuous replenishment of Mn, ensuring that the availability of Mn to the roots was the same throughout the experiment. This is in contrast to soil, where increased biomass production in response to P application during early growth stages may enhance Mn depletion in the rhizosphere and reduce the amount of Mn available for uptake and translocation to the youngest leaf, thereby leading to dilution of its Mn concentration. Secondly, the fact that the weight of the youngest leaf was not affected by the P treatments (data not shown) supports that the decline in Mn concentration reflected a negative effect of P on the actual Mn import rather than increased growth. Thirdly, the Fe and Cu concentrations did not decrease, but were only marginally affected by increasing P supply (Figure A1 in Appendix), as was also the case for N (Figure A2 in Appendix), thus contradicting a general dilution effect. Also barley plants growing in soil showed decreased Mn and Zn concentrations in the leaves in response to P supply, while Cu and Fe increased (Figure 8). Reduced foliar Mn concentrations under high soil-P availability are in accordance with previous experiments (Nielsen et al., 1992; Shane and Lambers, 2005) who similarly attributed this to physiological interactions. In a sand-culture experiment with eight barley cultivars, Zhu et al. (2002) showed that increased P availability reduced Zn shoot concentration in all cultivars, but only Mn concentrations in some of the cultivars. In the present work, there were no clear differences between the Mn-efficient genotype Antonia and the Mn-efficient genotype Vanessa in their response to elevated P, neither in terms of fluorescence induction kinetics of chlorophyll a (Figure 5) nor in the Mn concentration in the youngest fully emerged leaf (Figure 6). This was the case although Vanessa at all combinations of P and Mn produced up to 50% more biomass than Antonia (Table 2).

**CONCLUSION**

We have here shown that elevated P supply directly interferes with Mn uptake in barley roots and that this negative interaction can induce Mn deficiency in the shoot. The practical implication of this is that increased soil-P levels may be expected to lead to increased prevalence of Mn deficiency in barley crops.

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FIGURE A1 | The concentration Cu (A,B) and Fe (C,D) in the youngest fully emerged leaf of the Mn-inefficient genotype Antonia (A,C) and Mn-efficient genotype Vanessa (B,D) grown in hydroponics at different Mn and P levels. Different letters above columns mark significant differences between treatments ($P > 0.05$). Data are means ± SE ($n = 3$).
Figure A2 | The concentration of nitrogen (N) in the youngest fully emerged leaf of the Mn-inefficient genotype Antonia (A) and Mn-efficient genotype Vanessa (B) grown in hydroponics at different Mn and P levels. Different letters above columns mark significant differences between treatments (\(P > 0.05\)). Data are means ± SE (\(n = 3\)).