Abstract. Aluminium (Al) is a phytotoxic element affecting the growth and yield of many crop plants, especially in the tropics. Yet, some plants are able to accumulate high levels of Al. The monogeneric family Symplocaceae represents an Al accumulating family including many tropical and evergreen species with high Al levels in their above ground plant tissues. It is unclear, however, whether Al accumulation also characterises temperate species of Symplocos, and whether or not the uptake has a beneficial growth effect. Here, we investigate if the temperate, deciduous species Symplocos paniculata is able to accumulate Al by growing seedlings and saplings in a hydroponic setup at pH 4 with and without Al. Pyrocatechol-violet (PCV) and aluminon staining was performed to visualize Al accumulation in various plant tissues. Both seedlings and saplings accumulate Al in their tissues if available. Mean Al levels in leaves were 4107 (±1474 mg kg⁻¹) and 4290 (±4025 mg kg⁻¹) for the seedlings and saplings, respectively. The saplings treated without Al showed a high mortality rate unlike the Al accumulating ones. The seedlings, however, showed no difference in growth and vitality between the two treatments. The saplings treated with Al showed new twig, leaf and root development, resulting in a considerable biomass increase. PCV and aluminon staining indicated the presence of Al in leaf, wood and bark tissue of the plants. S. paniculata shares the capacity to accumulate Al with its tropical sister species and is suggested to be a facultative accumulator. Whether or not Al has a beneficial effect remains unclear, due to developmental differences between seedlings and saplings. Al is suggested to be transported via the xylem transport system into the leaves, which show the highest Al levels. Radial transport via ray parenchyma to bark tissue is also likely given the high Al concentrations in the bark tissue.

Keywords: Accumulation; aluminium; aluminon; hydroponics; pyrocatechol-violet; Symplocos paniculata; Symplocaceae.

Introduction

Although abundantly present in all terrestrial biomes, aluminium (Al) is typically absent as nutrient and as trace element within biochemical pathways of the living biosphere (Pogue and Lukiw, 2014). It is considered to be phytotoxic to the majority of plants if the soil pH decreases below 5.5 (Delhaize and Ryan, 1995; von Uexküll and Mutert, 1995), which causes Al to become soluble while changing its hydroxide form Al(OH)₃ to toxic forms such as Al(OH)²⁺, Al(OH)³⁺ and Al⁵⁺ (Kinraide,
Aluminium accumulation in a temperate accumulating plant

Immediate responses of plants sensitive to Al exposure include cease of root growth, lesions of the root tissue and subsequent nutrient deficiencies due to impaired uptake (Ryan et al., 1994; Kochian et al., 2004; Yang et al., 2011; Kopittke et al., 2015). The majority of plants growing in acidic soils have developed various strategies to avoid the uptake of Al into their plant body. Known strategies are the increase of the soil pH to prevent the solubility of toxic Al\(^{3+}\) (Taylor, 1991), or the exudation and complexation of Al through various organic compounds (e.g. oxalate, malate, catechins or phenolics) at the root level (Taylor, 1991; Barceló and Poschenrieder, 2002; Watanabe et al., 2008). Some plants are known to accumulate more than 1000 mg kg\(^{-1}\) drymass of Al in their stems and leaves and are often referred to as Al-accumulators (Jansen et al., 2002; Haridasan, 2008). Al accumulating species are mainly found in tropical regions and are, with few exceptions, woody plants most of which also possess blue flowers and/or fruits (Hutchinson and Wollack, 1943; Chenery, 1948a, b, 1949). Physiological mechanisms underlying the uptake of Al into above-ground plant tissues have been investigated in Fagopyrum esculentum (Buckwheat), Melastoma malabathricum, Hydrangea macrophylla and Camellia sinensis (tea), but remain unclear in many plant groups (Ma et al., 1997a, b; Shen et al. 2002; Watanabe et al., 2005b; Fung et al., 2008; Hajiboland et al., 2013; Wang et al., 2015).

The uptake of Al from the soil into the plant body does not lead to an equal distribution within the above ground plant organs. Al is suggested to follow the transpiration stream through the xylem into the leaf mesophyll of the plant, where the highest concentrations can be found (Haridasan et al., 1986; Tolrà et al., 2011; Maejima et al., 2014; Schmitt et al., 2016). High concentrations of Al in the bark also indicate a potential transport pathway via the phloem tissue (Zeng et al., 2013; Schmitt et al., 2016). However, the mechanisms underlying the chemical detoxification within Al accumulating species remain poorly understood. Possible mechanisms observed in Fagopyrum and Hydrangea include secretion and chelation of Al through organic acids such as oxalic acid or citrate (Ma et al., 1997a, b; Nguyen et al., 2003). While Al accumulating angiosperms are rather rare, representing about 5 % of all angiosperm species, this feature is characteristic of various monophyletic plant groups (Jansen et al., 2002).

In some species such as Camellia sinensis and Melastoma malabathricum, the availability of Al has a beneficial effect on plant growth by improving the growth of roots and the nutrient and water uptake capacity (Watanabe et al., 2005a; Hajiboland et al., 2013). Generally, two different types of accumulators can be defined, namely, obligate and facultative. Obligate accumulators can only grow on metalliferous soils and are unable to survive if a particular element is unavailable. Facultative accumulators, however, are growing well regardless of whether the soil contains a given metal or not (Pollard et al., 2014). Applying these two terms to Al accumulators, however, is not straightforward because growth experiments with Al have been conducted for a few species only. No difference in plant growth was observed for Fagopyrum, Camellia and Hydrangea, indicating that these species might be facultative accumulators (Ma et al., 1997a; Ma and Hiradate, 2000; Tolrà et al., 2011). Obligate Al accumulation was reported for Miconia albicans: seedlings of this Melastomataceae species growing in the Brazilian cerrado show a pronounced chlorosis and die-back when growing in a calcareous soil with alkaline pH and no available Al (Haridasan, 1988).

Symplocos includes ca. 300 species of woody trees and shrubs (Fritsch et al., 2008). The monogenic family Symplocaceae is suggested to originate in Eurasia and has from there dispersed to the American continent (Fritsch et al., 2015). Symplocos is most commonly found in montane tropical and subtropical rainforests, but also present in low altitude, temperate regions in Eurasia and the Americas, where is it also cultivated as ornamental shrub with dark blue berries (Doney, 1945; Nooteboom, 1977; Wang et al., 2004; Fritsch et al., 2008). The natural distribution of Symplocos paniculata is on slopes in mixed forests above 800 m across Taiwan, China, Japan, Korea and India (Rong-fen and Nooteboom, 1996). Within the genus, S. paniculata, S. chinensis and S. tinctoria are the only Symplocos species with deciduous leaves (Nooteboom, 1977; Rong-fen and Nooteboom, 1996; Fritsch et al., 2008). Furthermore, according to recent phylogenetic analyses, two of these species (S. paniculata and S. chinensis) represent a sister clade to all other Symplocos species (Wang et al., 2004; Fritsch et al., 2008).

Symplocos is known to include many strong Al accumulators, as reported for three Symplocos species (S. ophirensis, S. odoratissima, S. ambangensis) from Sulawesi, with an average Al concentration of 24180 ± 7236 mg kg\(^{-1}\) in old leaves (Schmitt et al., 2016). Maejima et al. (2014) reported Al accumulation in S. chinensis, growing in an Al containing hydroponic setup (8309 ± 7236 mg kg\(^{-1}\)) The highest leaf level of Al in Symplocos species was recorded in leaves of S. spicata with 72300 mg kg\(^{-1}\) (von Faber, 1925). Some Symplocos species are of economic interest to traditional weaving communities across the Indonesian archipelago, where Al containing leaves are used as a mordant in the natural dying process (Jansen et al., 2002; Cunningham et al., 2011; Schmitt et al., 2016).

In this study, we investigated the Al uptake in the temperate, deciduous species Symplocos paniculata (Symplocaceae) using a hydroponic setup. We
hypothesize that *S. paniculata* has a similar Al accumulation capacity as the evergreen, (sub-)tropical species. Earlier, semi-quantitative tests performed on *Symplocos*, show that 141 specimens out of 142 tested ones were Al accumulating as summarized in Jansen et al. (2002). Alternatively, it is possible that Al accumulation has been lost or remains poorly developed in *S. paniculata* because this species takes an isolated phylogenetic position within the genus, together with its temperate and deciduous habitus. Furthermore, we investigated whether this species shows similar distribution patterns of Al in its plant organs as values recently reported in tropical evergreen species (Schmitt et al., 2016). If this would be correct, the highest Al levels would occur in its leaves, followed by bark tissue and wood. Staining of anatomical sections was applied to visualize the distribution of Al in aboveground organs and tissue (Chenery, 1948b; Watanabe et al., 1998; González-Santana et al., 2012). By performing a growth experiment with and without Al we also tested if Al has a beneficial effect on its growth (Hajiboland et al., 2013), which can be expected based on the close phylogenetic relationship between Symplocaceae and Theaceae.

**Methods**

**Plant material**

We obtained 16 potted plants of *Symplocos paniculata* from a nursery (Rein&Mark Bulk, Boskoop, The Netherlands), which will be referred to hereafter as ‘saplings’. The nursery grew the plants from seeds obtained from the UK. The saplings were ca. 1–3 years old and 15–50 cm tall. The pH of the soil was ca. 5.5.

In addition, 20 seedlings of *S. paniculata* were grown from seeds obtained from the Charles Keith Arboretum in North Carolina, USA in autumn 2014. The pulp of the fruit was removed and seeds were sown in sand in September 2014. The seeds were stored at 4 °C for 3 months and then placed outside at ambient temperature in January 2015. When the first leaves were visible in May 2015, seedlings were put into soil (bog soil with pumice, pH ca. 5.5) prior to a hydroponic experiment. A total of 20 seedlings (5–10 cm tall) were grown outside at the Botanical Garden of Ulm University in half-shaded conditions until the hydroponic experiment started in August 2015. These plants are hereafter referred to as ‘seedlings’.

**Hydroponic experiments**

The saplings were divided into two groups (i.e. *n* = 8 individuals per group), each group having the same number of specimens with a similar height and age. First, the soil was carefully removed from the plants with tap water. After removal of the substrate the roots were thoroughly rinsed with demineralized water. Plants were weighted and transferred into 35 l vessels, with four plants per vessel, respectively. The vessels contained a standard nutrient solution following Watanabe and Osaki (2001) and were constantly aerated with an aquarium pump (EHEIM air pump 200, EHEIM GmbH & Co KG, Deizisau, Germany). The solution consisted of 2.14 mM N (NH4NO3), 32 μM P (NaH2PO4·2H2O), 0.77 mM K (K2SO4; KCl = 1:1), 1.25 mM Ca (CaCl2·2H2O), 0.82 mM Mg (MgSO4·7H2O), 35.8 μM Fe (FeSO4·7H2O), 9.1 μM Mn (MnSO4·4H2O), 46.3 μM B (H3BO3), 3.1 μM Zn (ZnSO4·7H2O), 0.16 μM Cu (CuSO4·5H2O) and 0.05 μM Mo ((NH4)6Mo7O24·4H2O). The pH was adjusted to pH 4 with HCl and NaOH using a pH test stick (pH PAL Plus, ETI Ltd, UK). The nutrient solutions were changed on a weekly basis. The plants were kept in this reference solution for 2 weeks to adapt to the hydroponic environment. Then, two vessels including a total of eight plants were treated with the standard nutrient solution with added 1 mM L-1 AlCl3 until the end of the experiment (hereafter named +Al), whereas the other two vessels were not given any Al (−Al) [see Supporting Information]. Furthermore, the plants were sprayed with demineralized water on a daily basis to avoid dehydration and to reduce the heat stress by sun exposure at midday. During the third week all vessels were covered with a grey lid to prevent the growth of algae in the nutrient solution [see Supporting Information]. At the end of the experiment, all plants (including the fallen leaves) were removed from the vessels, weighed again, rinsed with demineralized water, put into plastic bags and kept in the freezer at −25 ºC for further analysis.

The experimental conditions of the seedling experiment in 2015 were analogous to the hydroponic setup with saplings in 2014 except for minor differences. Due to the smaller plant size of the seedlings, we used a vessel size of 4.5 l. Each vessel contained five plants, with four vessels in total (*n* _plants_ = 20 individuals). Two vessels were treated with the standard nutrient solution with 1 mM AlCl3 after 2 weeks of acclimatisation in the standard nutrient solution. At the end of the experimental phase, both the roots and leaves of the 20 saplings were scanned (EPSON Expression 10 000 XL, SEIKO EPSON Corporation, Nagano, Japan). Root morphological parameters were analysed with WinRhizo 2012 and leaf area was measured using ImageJ (Version 1.49, U. S. National Institutes of Health, Bethesda, Maryland, USA, http://imagej.nih.gov/ij/ (26 September 2016), 1997–2015). Both experiments were conducted inside the greenhouses of the Botanical Garden Ulm. Climate conditions were controlled to avoid considerable variation. Temperatures were kept above 12 ºC at night and below 33 ºC during the day, with a relative humidity between
total of 18 bark, 20 leaf, 20 root and 18 wood samples. The seedling experiment (root samples and 6 new root samples were analysed. wood samples, 16 leaf samples, 9 bark samples, 16 old roots, and roots, which were analysed separately. Root material was defined as the sum of tissue that was immersed in the nutrient solution. Although no differentiation between bark and wood tissue of the roots was made, the new roots of the +Al treatments were separated from the old roots and could easily be distinguished by their white colour (Fig. 1). The tissues were put into paper bags and oven dried at 75 °C for 24 h. Samples were then ground with a centrifuge mill (ZM1, Retsch, Germany); 50 mg of the sample was put into 10 ml PTFE vessels, and digested with HNO3 in a microwave oven (Mars 5 plus, CEM, Germany) at 200 °C for 30 min. Digestions were diluted to 25 ml with a 0.25 % (w/v) CsCl-solution as ionic buffer. The concentrations of Al, K, Ca, Mg and Fe were measured based on microwave-plasm atomic emission spectrometry (MP-AES, Agilent 4100, Agilent Technologies, Australia). Before starting the hydroponic experiment, Al concentrations in leaves of the saplings were measured to determine the pre-treatment concentration of Al in the plants, which was 390 mg kg⁻¹ in the leaf tissue. Furthermore, the Al concentration of 25 seeds and their fleshy pericarp was 3450 mg kg⁻¹ and 6150 mg kg⁻¹, respectively. In the sapling experiment (n = 16 specimens), 12 wood samples, 16 leaf samples, 9 bark samples, 16 old root samples and 6 new root samples were analysed. The seedling experiment (n = 20 specimens) included a total of 18 bark, 20 leaf, 20 root and 18 wood samples. Hence, all plants, including those that died, were used for our chemical analyses. We were unable to separate the bark and wood tissue for four sapling specimens and two seedling specimens, which were not included in our analyses.

**Biomass and root–shoot ratio**

The plants were weighed before they were put into the vessels at the beginning of the experiment. At the end of the experiment, the total weight of each plant was measured again. All leaves that were shed during the experimental phase were collected and included in the biomass calculation. As a relative measure, gain or loss of fresh biomass (Δm) was defined as:

\[ Δm = m_{after} - m_{before} \]

with \( m_{after} \) representing the plant weight in g of the whole plant at the end of the experiment, and \( m_{before} \) the plant mass before the experiment. The root–shoot ratio was calculated by dividing the dry mass of root tissue by the sum of dried above-ground tissues (i.e. leaves, bark and wood) for each plant. Both root–shoot ratio and \( Δm \) were calculated for all specimens in both experiments (saplings: \( n = 16 \); seedlings: \( n = 20 \)).

**Visualisation of Al accumulating tissue**

For the visualisation of Al in plant tissue, a total of eight saplings, four from each treatment, were sampled. Wood sections were cut with a sliding microtome. Leaf material was cut into 0.5 × 0.5 cm sections and fixed in FPA (10 % formalin [37 %], 5 % propionic acid, 50 % ethanol [95 %] and 35 % distilled water) over night. Samples were then dehydrated using a graded series of ethanol (30, 50, 70 and 96 %) and tert-butanol, and embedded in paraffin. The embedded leaf sections were cut with a rotary microtome, transferred on adhesive-coated objective slides (Menzel-Gläser, Thermo Scientific, Gerhard Menzel GmbH, Braunschweig, Germany) and a graded ethanol series (95, 70, 50 and 30 %). Two staining methods were applied to locate high levels of Al in the tissue sections. Staining was conducted using a 0.052 % (w/v) pyrocatechol-violet (PCV) solution in 2.5 % hexamine-NH4OH buffer (pH 6.2), which was applied for 15 min and then washed with the buffer solution prior to mounting (Watanabe et al., 1998). The second staining included ammonium aurin tricarboxylate (aluminon) (Chenery, 1948b; Clark and Krueger, 1985). After staining, the sections were washed in an additional ethanol series and mounted in Neo-Mount (Merck KGaA, Darmstadt, Germany).
Statistical analysis
A Shapiro–Wilk test was performed to check for normal distribution and to determine the statistical tests to be applied. Differences between group means were analysed using the non-parametric Wilcoxon rank-sum test (Abiomass, [Al], [Ca], [Fe]roots) or Welch’s t-test for parametric data (root–shoot ratio). Correlations between Al and calcium (Ca) were computed using the psych package of R with Spearman’s rho for non-parametric data. All statistical analyses were carried out with the freeware ‘R’ (RStudio, Version 0.99.902, RStudio Inc., Boston, USA).

Results
Hydroponic experiments on saplings and seedlings
After putting the saplings in the hydroponic solution for two weeks during the pre-treatment period, one sapling in the –Al treatment was considered dead according to visual properties (i.e. leaf desiccation and strong chlorosis) and showed no signs of recovery afterwards. A similar sapling was found in the +Al treatment, but was found to recover two weeks later. Regardless of the size of the plant and the +Al or –Al treatment, some leaves showed yellowish-brown leaf margins after 3 weeks (i.e. 2 weeks of pre-treatment and 1 week of +Al or –Al treatment). In the –Al treatment, basal leaves started to dry out without changing their colour. Two specimens from the +Al treatment showed a similar condition after the second week (i.e. before the +Al treatment started). In week 4, all saplings of the –Al treatment were seriously stressed and showed clear signs of desiccation. After week 5 (i.e. 3 weeks after starting with the Al treatment), new white roots were visible in the +Al saplings, which continued growing until the end of the experiment (Fig. 1). After 17 weeks of treatment, all plants in the –Al treatment were considered to be dead, with only the largest sapling remaining in an unhealthy, stressed condition. Except for two plants, all specimens in the +Al treatment developed new side branches and leaves and were healthy [see Supporting Information]. Changes in the fresh biomass during the hydroponic experiment showed a clear difference between the two treatments (Fig. 2). All –Al saplings had negative values with an average loss in fresh biomass of −3.45 g ± 3.46 g, whereas the +Al treatment showed a gain of 12.24 g ± 11.24 g. The highest loss was −12.14 g, and the largest increase was 28.62 g. A significant (Welch’s t-test, t = −2.3231, df = 11.718, P = 0.03903) difference was found between the root–shoot ratio of the –Al treatment (0.20 ± 0.08) and the +Al treatment (0.35 ± 0.14, Fig. 2).

In contrast to the saplings, the seedlings did not show visible differences in vitality (e.g. leaf chlorosis, desiccation) between the +Al and –Al conditions. No dead seedlings were observed in both conditions, although several plants dropped a few leaves. Furthermore, no difference in biomass (Wilcoxon rank-sum test, W = 62, P = 0.393) and root–shoot ratio (Welch’s t-test, t = −0.8788, df = 17.78, P = 0.3912) was found between the +Al and –Al seedlings (Fig. 2). The root morphology between the two conditions did not differ significantly from each other (Wilcoxon rank-sum test, root length: W = 76, P = 0.05243; root surface area: W = 75, P = 0.06301; root tips: W = 71, P = 0.123; root forks: W = 71, P = 0.123; Fig. 3), although the mean values of the root traits measured were higher in the +Al treatment than in the –Al seedlings.

Al accumulation in the plant tissues
There was a significant difference (Wilcoxon rank-sum test, W = 845, P < 0.01, Fig. 2) in Al concentration between the +Al saplings (5728 mg kg⁻¹ ± 4792 mg kg⁻¹) and –Al saplings (145 mg kg⁻¹ ± 117 mg kg⁻¹) at the entire plant level, summarizing all above and below-ground organs. The highest Al concentrations in the +Al treatment were found in the new roots (12936 mg kg⁻¹ ± 5294 mg kg⁻¹, see Supporting Information), whereas the stem wood tissue had the lowest concentration (1618 mg kg⁻¹ ± 842 mg kg⁻¹). Highest Al concentrations in the –Al treatment were found in the root tissue (196 mg kg⁻¹ ± 168 mg kg⁻¹) and the lowest in the stem wood (46 mg kg⁻¹ ± 21 mg kg⁻¹), respectively.

In the seedling experiment, the mean Al concentration of all tissues in the +Al treatment (3242 ± 1584 mg kg⁻¹) was significantly higher (Wilcoxon rank-sum test, W = 2235.5, P < 0.01) than the concentration in the –Al treatment (381 ± 226 mg kg⁻¹, Fig. 2). In the +Al treatment, the highest Al levels were found in the leaves with a mean concentration of 4107 (± 1474) mg kg⁻¹ dry mass (Fig. 4, see Supporting Information), followed by the root (3749 ± 1304 mg kg⁻¹), bark (3054 ± 419 mg kg⁻¹) and the wood tissue (1038 ± 357 mg kg⁻¹), respectively. The Al values were overall lower in the –Al treatment and had the highest levels in the bark tissues (481 ± 197 mg kg⁻¹). The leaf Al concentration in the –Al seedlings had a mean value of 456 (± 243) mg kg⁻¹ and was slightly lower than the bark concentration. The lowest concentration was found in the wood tissue (136 ± 32 mg kg⁻¹).

Concentration of additional elements
The Ca concentration at the entire plant level in the saplings was found to be significantly lower (Wilcoxon rank-sum test, W = 292.5, P = 0.04288) in the +Al treatment (4205 ± 4007 mg kg⁻¹) than in the –Al treatment
No significant correlation between Al and Ca was found (Spearman correlation, Al: \( r_s = 0.23, P = 0.27 \); +Al: \( r_s = -0.18, P = 0.31 \)). The K concentration in the sapling roots differed significantly (Wilcoxon rank-sum test, \( W = 41, P = 0.0293 \)) between the two treatments when comparing the new roots of the +Al treatment with the roots of the –Al treatment. However, no significant difference in K was found when comparing the old roots of the +Al treatment with the roots of the –Al treatment (Wilcoxon rank-sum test, \( W = 48, P = 0.1049 \)).

The concentrations of Ca and Mg were similar in both treatments for each plant organ of the seedlings. For both treatments, a significant, positive correlation was found between Al and Ca (Spearman correlation, –Al: \( r_s = 0.63, P < 0.01 \); +Al: \( r_s = 0.69, P < 0.01 \)). The Fe concentration in the roots was significantly (Wilcoxon rank-sum test, \( W = 10.5, P < 0.01 \)) lower in the +Al treatment (2644 ± 1437 mg kg\(^{-1}\)) than in the –Al condition (6668 ± 4242 mg kg\(^{-1}\), see Supporting Information).

Al visualisation based on light microscopy

Both staining solutions applied visualized Al in leaf tissues of the +Al plants, but not in the –Al plants (Fig. 5). PCV stained Al accumulating tissue in blue and was present in both the wood and the leaf tissue. In the wood tissue, tangential sections showed some blue stained ray parenchyma cells and axial parenchyma cells (Fig. 5). Cross-sections showed coloured cell walls in older growth rings and a weak blue colour in the pith parenchyma. In both tangential and cross sections, the bark tissue was blue. Colourless structures included most vessels, tracheids, and fibres in the younger wood. Leaf cross sections showed the presence of Al in the mesophyll, where the colour was stronger in the spongy parenchyma than in the palisade cells (Fig. 5). The vascular bundles of secondary and tertiary veins also contained Al, but no distinction between phloem and xylem could be made. The main vein, however, showed stained parenchyma cells in the xylem tissue and no colour in the vessels and tracheids within the vascular bundle.
Some cells within the phloem of the main vein were stained blue. Similar results as for PCV were observed for aluminon, which in the presence of Al was creating a red crimson pigment. Red cells were found across the pith, wood and bark tissue (Fig. 5), but unstained tissue could not be clearly distinguished from coloured tissue. Tangential sections visualised Al in both radial and axial parenchyma cells, with a more pronounced signal than the PCV stained sections. Not all parenchyma cells in the tangential sections showed the same colour intensity. The leaf sections stained with Aluminon showed a pale red colour in the cell walls of the leaf epidermis, spongy parenchyma and palisade cells. Furthermore, vascular bundles showed the presence of Al (Fig. 5), where the dye was more pronounced than in the rest of the leaf tissue. The main vein had an intense red colour in the parenchyma above the protoxylem, and similarly to the PCV staining, Al was found in the parenchyma cells of the xylem [see Supporting Information].

Discussion

Phylogenetic implications

Based on our hydroponic experiments, Symplocos was found to accumulate Al when soluble Al was available to the plants. For only a small proportion of Symplocos species exact Al concentrations have been measured, and all show high levels of Al in their leaves (Table 1). The values for S. chinensis and S. paniculata, however, do not reflect natural conditions, and may therefore over- or underestimate the Al levels of plants in their natural environment. Nevertheless, the overall accumulation in S. paniculata was lower than in S. chinensis, which accumulated on average 8309 ± 282 mg kg⁻¹ Al in its leaves after a 4-month hydroponic experiment with half the concentration used in our experiments (0.5 mM L⁻¹ AlCl₃) (Maejima et al., 2014). S. chinensis is closely related to S. paniculata and both species have been suggested to be taxonomically merged to a single species, or at least form a separate phylogenetic clade (Wang et al., 2004; Nooteboom, 2005).
Given the basal phylogenetic position of *S. paniculata* within the genus (Wang et al., 2004; Fritsch et al., 2008) and based on earlier reports on the phylogeny of Al accumulation within angiosperms (Chenery, 1948b, 1949; Jansen et al., 2002, 2003, 2004), this study suggests that Al accumulation characterizes all *Symplocos* species. Therefore, Al accumulation is not limited to tropical *Symplocos* species. In general, this trait is rare for temperate plants. An exception, however, is the Diapensiaceae family, which belongs to the same clade as Theaceae and Symplocaceae within the Ericales (Chenery, 1951; Jansen et al., 2004).

**Al distribution within the plants**

The variable Al concentrations in the tissues and organs most likely reflect the transport pathways of Al within the plant and show similar distribution patterns as naturally growing *Symplocos* trees in Indonesia (Schmitt et al., 2016). Although the chemical form in which Al is taken up by the plant is unknown, the primary uptake mechanism appears to be transport via the transpiration stream, as the highest concentrations were observed in the leaves of the plants. The most commonly reported forms of Al complexes detected in the xylem sap of Al accumulators are Al citrate, malate, oxalate and fluoride, as these molecules are known to have a strong binding affinity to Al (Watanabe and Osaki, 2001; Flaten, 2002; Morita et al., 2004; Satoh, 2006; Klug et al., 2015). The root represents the only organ that is directly in contact with soluble Al, explaining the high concentrations within the root tissue. High concentrations of Al have also been found in the bark tissue of *S. paniculata*, indicating a possible participation of the phloem tissue in Al uptake. A similar interaction has been reported for *Camellia oleifera* (Zeng et al., 2013) and is likely to occur as Al is known to have a high binding affinity to a variety of carbohydrates and ligands with carboxylate and pheno- late functional groups (Miltner and Zech, 1998; Flaten, 2002). It is unclear whether the distribution of Al into the phloem follows a transport pathway similar to photosynthetic products in leaves, or is transported axially from the roots and then spreads radially from the stem xylem into the bark. Radial transport from the xylem into vessel associated parenchyma cells and subsequent transport to the bark tissue via the ray parenchyma is indicated by...
the staining methods applied and supports a potential interaction between xylem and phloem (Fig. 5). A positive correlation between Al and Ca was found for the seedlings, which was also reported for the tropical Al accumulators in Sulawesi (Schmitt et al., 2016) and Sumatra (Masunaga et al., 1998), suggesting a possible link of both metals in their transport pathway.

Figure 5. Leaf and wood cross sections of *Symplocos paniculata* saplings stained with aluminon or pyrocatechol-violet (PCV) after treatment with 1 mM AlCl₃ (+Al treatment). (A) Stem cross section stained with aluminon. (B) Leaf cross section stained with aluminon. (C) Stem cross section stained with aluminon showing the presence of Al in the ray parenchyma (white asterisk). (D) Leaf cross section stained with PCV. (E) Tangential stem wood section of the –Al treatment stained with PCV. (F) Tangential stem wood section of the +Al treatment stained with PCV. (G) Leaf cross section of the –Al treatment after staining with aluminon. UE = upper epidermis, LE = lower epidermis, PP = palisade parenchyma, SP = spongy parenchyma, VB = vascular bundle. Scale bar = 100 μm.

Beneficial effect of Al

Whether or not Al has a beneficial effect on the growth of *S. paniculata* is complicated by the differences observed between seedlings and saplings. The seedlings did not seem to indicate a beneficial effect of Al on the growth of *S. paniculata*. None of the features measured (e.g. leaf area, leaf number, biomass, root-
shoot ratio) showed a significant difference between the $+$Al and $-$Al seedlings. However, a tendency towards induced root growth could be found in the $+$Al seedlings, although no statistical significance was found for this trend.

The saplings, however, showed a pronounced difference between the $-$Al and $+$Al treatment. During the first two weeks in hydroponics, all saplings showed a stress reaction, indicated by leaf discission and partial leaf chlorosis at the leaf margins. Eventually all but one specimen of the $-$Al treatment died, whereas the $+$Al treatment saplings recovered, flushed new leaves and produced new side branches. Furthermore, the $+$Al saplings significantly increased their root biomass, which was indicated by an increase in the root–shoot ratio. A similar increase in fine roots and increased biomass due to Al application was reported by Watanabe et al. (2005a) in saplings of *Melastoma malabathricum*. Fung et al. (2008) also reported a beneficial, stimulatory effect on the roots and an enhanced nutrient uptake in *Camellia sinensis* seedlings.

The difference in growth and plant mortality between the seedlings and saplings remain unclear. One explanation could be the origin of the seeds and the saplings, which were taken from different populations. The genetic variability could affect the growth of the plants and their susceptibility when transferred from a soil substrate to hydroponic condition. The potential impact of genetic variability on plants growing in hydroponics has been reported for *Sorghum bicolor* and *Fagopyrum esculentum* (Jordan *et al.*, 1979; Klug *et al.*, 2015), but may not explain the subsequent death of the $-$Al plants. Furthermore, earlier growth conditions between our seedlings and saplings differed, as the saplings were grown in a greenhouse, experiencing mild winter temperatures in contrast to the seedlings, which were germinated at ambient winter temperatures. The harsh winter conditions for the seedlings might have been beneficial for the hydroponics experiment, while the saplings obtained from a nursery were probably more sensitive to environmental changes. It is also possible that developmental differences in nutrient requirements contributed to the differences in our hydroponic experiment between seedlings and saplings.

Before we conducted our experiments, *S. paniculata* plants were grown in peat soil with a relatively low pH (ca. 5.5). Despite the fact that the species investigated is a temperate shrub, the natural distribution of *Symlocos* is in tropical to subtropical environments, mostly reflecting biomes with a low soil pH. Therefore, soil acidity below 5.5 cannot be regarded as a potential harm for the *S. paniculata* plants in our experimental setup, because the plants are naturally adapted to a low soil pH status (Brunner and Sperisen, 2013). Furthermore, a low soil pH is needed to ensure the availability of $\text{Al}^{3+}$ to the plants.

The only difference between the $+$Al and $-$Al treatments was the added 1 mM AlCl$_3$. Thus, a possible explanation for the death of the $-$Al saplings could be the stress reaction induced by the hydroponic condition.

---

### Table 1. Mean foliar aluminium concentration of 13 *Symlocos* species compiled from literature (see References) and the present study.

| Species                  | Author                     | Al [mg kg$^{-1}$] | SD  | References                     | Distribution   |
|--------------------------|----------------------------|------------------|-----|--------------------------------|----------------|
| *S. ambangensis*          | Noot.                      | 16.719           | 1.12| Schmitt *et al.* 2016          | SE-Asia        |
| *S. chinensis* (asterisk)| Jacq.                      | 8.309            | 282 | Maejima *et al.* 2014          | Eastern Asia   |
| *S. coreana*              | Jacq.                      | 14.766           | n.a.| Watanabe *et al.*, 2007       | Eastern Asia   |
| *S. crassipes*            | Jacq.                      | 33.883           | n.a.| Metali, 2010                  | SE-Asia        |
| *S. crataegoides*         | Jacq.                      | 6.56             | n.a.| Chenery, 1948a, b             | Central Asia   |
| *S. lancifolia*           | Jacq.                      | 23.05            | n.a.| Watanabe *et al.*, 2007       | Eastern Asia   |
| *S. myrtacea*             | Jacq.                      | 7.355            | n.a.| Chenery, 1948a, b             | Central Asia   |
| *S. odoratissima*         | Choisy ex Zoll.            | 23.383           | 9593| Schmitt *et al.*, 2016        | SE-Asia        |
| *S. ophirensis*           | C.B.Clarke                 | 21.352           | 2802| Schmitt *et al.*, 2016        | SE-Asia        |
| *S. paniculata* (asterisk)| Miq.                      | 4.125            | 3541| This study                    | Eastern Asia   |
| *S. prunifolia*           | Jacq.                      | 1.85             | n.a.| Watanabe *et al.*, 2007       | Eastern Asia   |
| *S. spicata*              | Jacq.                      | 72.3             | n.a.| von Faber, 1925               | Central Asia   |

Al concentration for *S. Chinensis* and *S. Paniculata* (asterisk) are based on hydroponic experiments and do not reflect natural conditions. Species distribution is based on the global biodiversity information facility (GBIF).
together with an impaired nutrient uptake. The limited availability of phosphorous (P) in the nutrient solution could also be a potential explanation for induced root growth in the +Al treatment, which is known to occur if plants experience a P-deficient substrate (Kochian et al., 2004; Kochian, 2012). Watanabe and Osaki (2001) reported a link between Al application and P nutrition in *M. malabathricum*, which did not indicate P as primary reason for growth enhancement. Konishi et al. (1985) observed a stimulatory effect of Al on the P-uptake in tea plants, which could also confirm our findings for the +Al treatment. A potential precipitation of Al with P, which reduces the P-availability, could be suggested (Watanabe et al., 2005a), and may account for the increased root growth observed. This scenario would explain the growth of the +Al treatment sapling as an indirect effect because of the added Al, which would subsequently precipitate the P and reduce its availability. However, the −Al treatment had the same nutrient solution with the same P concentration. Thus, the enhanced root growth due to P deficiency would also have been observed in the −Al treatment, which means that P availability cannot be regarded as the primary and only reason for the high mortality of the −Al saplings.

**Conclusions**

Our hydroponic experiments indicate that *S. paniculata* has the capacity to accumulate Al in its aboveground tissues. Al accumulation in this species appears to be facultative and the beneficial effect of Al on its growth is complicated by the different response to Al between seedlings and saplings. Variation in Al levels across different organs and tissues in *S. paniculata* is similar to tropical species within this genus, with the highest Al concentrations occurring in leaves and bark tissue.

**Sources of Funding**

This work was funded by the Juniorprofessuren-Programm of the Ministry for Science, Research and the Arts of Baden-Württemberg, Germany.

**Contributions by the Authors**

M.S. and S.J. planned the hydroponic experiments. M.S. conducted the hydroponic experiments, the elemental analysis and the anatomical observations. T.W. gave critical input to the hydroponic experiments and the setup. All authors contributed substantially to the writing of the manuscript.

**Conflict of Interest Statement**

None declared.

**Acknowledgements**

For technical assistance we thank Nadine Buchsteiner, Jakob Gerber, Ellen Salzer, Hans Malchus, Jutta Siegmund-Jonitz and Peter Zindl. We also thank Charles Keith from the Keith Arboretum in North Carolina for supplying us with *Symplocos paniculata* seeds.

**Supporting Information**

The following additional information is available in the online version of this article —

Figure S1. Hydroponic setup for the growth experiment with 16 specimens of *Symplocos paniculata* saplings in the greenhouses of the Botanical Garden at Ulm University. (A) Setup at the beginning of the experiment. (B) Setup at the end of the experimental phase. Except for the tallest plant (arrow), the −Al plants on the left side were all dead, whereas the +Al plants were resprouting and healthy. The grey cover plates prevented the growth of algae.

Figure S2. Magnified cross sections of the central leaf vein of *Symplocos paniculata* stained with two different techniques to visualize Al in the tissue in plants that were growing in a nutrient solution containing 1 mM AlCl$_3$ for 4 months. Black arrows show the presence of Al in the ray cells of the xylem tissue of the central vein. (A) Aluminon. (B) Pyrocatechol-violet. (C) Staining with aluminon in a plant growing without Al. Scale bar = 100 μm

Table S1. Elemental concentrations of various tissues of *Symplocos paniculata* growing in hydroponic solutions for 4 months ($n=8$ saplings per treatment) and 2 months ($n=10$ seedlings per treatment). Plants in the +Al-treatment were given 1 mM AlCl$_3$. All values are given in mean mg kg$^{-1}$ dry mass (± SD). New roots were only formed in the +Al seedling plants and not in the seedlings.

**Literature Cited**

Barceló J, Poschenrieder C. 2002. Fast root growth responses, root exudates, and internal detoxification as clues to the mechanisms of aluminium toxicity and resistance: a review. *Environmental and Experimental Botany* 48:75–92.

Brunner I, Sperisen C. 2013. Aluminium exclusion and aluminium tolerance in woody plants. *Frontiers in Plant Science* 4:1–12.

Chenery EM. 1948a. Aluminium in plants and its relation to pigments. *Annals of Botany* 12:121–136.

Chenery EM. 1948b. Aluminium in the plant world - Part I. General survey in dicotyledons. *Kew Bulletin* 2:173–183.
Chenery EM. 1949. Aluminium in the plant world - Part II. Monocotyledons and gymnosperms. Kew Bulletin 4:463–473.

Chenery EM. 1951. Some aspects of the aluminium cycle. Journal of Soil Science 2:97–109.

Clark RA, Krueger GL. 1985. Aluminon: Its limited application as a reagent for the detection of aluminium species. Journal of Histochemistry & Cytochemistry 33:729–732.

Cunningham AB, Maduarta IM, Howe J, Ingram W, Jansen S. 2011. Hating by a thread: natural metallic mordant processes in traditional Indonesian textiles. Economic Botany 65:241–259.

Delhaize E, Ryan PR. 1995. Aluminium toxicity and tolerance in plants. Plant Physiology 107:315–321.

Doney C. 1945. Shrubs for special uses. Plants and Gardens 1:18–51.

von Faber FC. 1925. Untersuchungen über die Physiologie der javanischen Solfataren-Pflanzen. Flora 118:89–110.

Flaten TP. 2002. Aluminium in tea—concentrations, speciation and bioavailability. Coordination Chemistry Reviews 228:385–395.

Fritsch P, Kelly LM, Wang Y, Almeda F, Kriebel R. 2008. Revised infrafamilial classification of Symplocaceae based on phylogenetic data from DNA sequences and morphology. Taxon 57:823–852.

Fritsch PW, Manchester SR, Stone RD, Cruz BC, Almeda F. 2015. Northern Hemisphere origins of the amphi-Pacific tropical plant family Symplocaceae. Journal of Biogeography 42:891–901.

Fung KF, Carr HP, Zhang J, Wong MH. 2008. Growth and nutrient uptake of tea under different aluminium concentrations. Journal of the Science of Food and Agriculture 88:1582–1591.

González-Santana IH, Márquez-Guzmán J, Cram-Heydrich S, Cruz-Ortega R. 2012. Conostegia xalapensis (Melastomataceae): an aluminium accumulator plant. Physiologia Plantarum 144:134–145.

Hajiboland R, Bohrami Rod S, Barceló J, Poschenrieder C. 2013. Mechanisms of aluminium-induced growth stimulation in tea (Camellia sinensis). Journal of Plant Nutrition and Soil Science 176:616–625.

Haridasan M. 1988. Performance of Miconia albicans (SW.) Triana, an aluminium-accumulating species, in acidic and calcareous soils. Communications in Soil Science and Plant Analysis 19:1091–1103.

Haridasan M. 2008. Nutritional adaptations of native plants of the cerrado biome in acid soils. Brazilian Journal of Plant Physiology 20:183–195.

Haridasan M, Paviani TJ, Schiavini I. 1986. Localisation of aluminium in the leaves of some aluminium-accumulating species. Plant and Soil 94:435–437.

Hutchinson GE, Vollick A. 1943. Biological accumulators of aluminium. Transactions of the Connecticut Academy of Arts and Sciences 35:73–128.

Jansen S, Broadley MR, Robbrecht E, Smets E. 2002. Aluminium hyperaccumulation in angiosperms: a review of its phylogenetic significance. The Botanical Review 68:235–269.

Jansen S, Watanabe T, Caris P, Geuten K, Lens F, Pyck N, Smets E. 2004. The distribution and phylogeny of aluminium accumulating plants in the Ericales. Plant Biology 6:498–505.

Jansen S, Watanabe T, Dessein S, Smets E, Robbrecht E. 2003. A comparative study of metal levels in leaves of some Al-accumulating Rubieaeae. Annals of Botany 91:657–663.

Jordan WR, Miller FR, Morris DE. 1979. Genetic variation in root and shoot growth of Sorghum in hydroponics. Crop Science 19:468–472.

Kinraide TB. 1991. Identity of the rhizotoxic aluminium species. Plant and Soil 134:167–176.

Kinraide TB. 1997. Reconsidering the rhizotoxicity of hydroxyl, sulphate, and fluoride complexes of aluminium. Journal of Experimental Botany 48:1115–1124.

Klug B, Kirchner TW, Horst WJ. 2015. Differences in aluminium accumulation and resistance between genotypes of the genus Fagopyrum. Agronomy 5:418–434.

Kochian LV. 2012. Rooting for more phosphorus. Nature 488:466–467.

Kochian LV, Hoekenga OA, Pineros MA. 2004. How do crops plants tolerate acid soils? Mechanisms of aluminum tolerance and phosphorous efficiency. Annual Review of Plant Biology 55:459–493.

Konishi S, Miyamoto S, Taki T. 1985. Stimulatory effects of aluminium on tea plants grown under low and high phosphorus supply. Soil Science and Plant Nutrition 31:361–368.

Kopittke PM, Moore KL, Lombi E, Gianoncelli A, Ferguson BJ, Blamey A, McNees NW, Nicholson TM, McKenna B, a, Wang P, Gresshoff PM, Kourousias G, Webb RI, Green K, Tollenaere A. 2015. Identification of the primary lesion of toxic aluminum in plant roots. Plant Physiology 167:1402–1411.

Ma JF, Hirade S. 2000. Form of aluminium for uptake and translocation in buckwheat (Fagopyrum esculenta Moench). Planta 211:355–360.

Ma JF, Hirade S, Nomoto K, Ishiwata T, Matsumoto H. 1997a. Internal detoxification mechanism of Al in Hydrangea. Plant Physiology 113:1033–1039.

Ma JF, Zheng SJ, Matsumoto H, Hirade S. 1997b. Detoxifying aluminium with buckwheat. Nature 390:569–570.

Maejima E, Hirade T, Sansen S, Osaki M, Watanabe T. 2014. Comparative analysis of aluminum accumulation in leaves of three angiosperm species. Botany 92:327–331.

Masunaga T, Kubota D, Hotta M, Wakatsuki T. 1998. Nutritional characteristics of mineral elements in leaves of tree species in tropical rain forest, West Sumatra, Indonesia. Soil Science and Plant Nutrition 44:315–329.

Metali, F. 2010. Factors controlling Al accumulation in plants: effects of phylogenoy, soil conditions and external nutrient supply. PhD Thesis, University of Aberdeen, United Kingdom.

Miltner A, Zech W. 1998. Carbohydrate decomposition in beech litter as influenced by aluminium, iron and manganese oxides. Soil Biology & Biochemistry 30:1–7.

Morita A, Horie H, Fujii Y, Takatsu S, Watanabe T, Yagi A, Yokota H. 2004. Chemical forms of aluminium in xylem sap of tea plants (Camellia sinensis L.). Phytochemistry 65:2775–2780.

Nguyen NT, Nakabayashi K, Thompson J, Fujita K. 2003. Role of exudation of organic acids and phosphate in aluminium tolerance of four tropical woody species. Tree Physiology 23:1041–1050.

Nooteboom HP. 1977. Symplocaceae. In: Van Steenis CGGJ, De Wilde WJJO, eds. Flora malesiana. Dordrecht: Kluwer Academic Publishers, 105–274.

Nooteboom HP. 2005. Additions to Symplocaceae of the old world including New Caledonia. Blumea 50:407–411.

Pogue A, Lukiw WJ. 2014. The mobilization of aluminium into the biosphere. Frontiers in Neurology 5:1–4.
Pollard AJ, Reeves RD, Baker AJM. 2014. Facultative hyperaccumulation of heavy metals and metalloids. *Plant Science* **217**:8–17.

Rong-fen W, Nooteboom HP. 1996. *Symplocaceae*. *Flora of China* **15**:235–252.

Ryan PR, Kinraide TB, Kochian LV. 1994. Al$^{3+}$–Ca$^{2+}$ interactions in aluminum rhizotoxicity. *Planta* **192**:98–103.

Satoh S. 2006. Organic substances in xylem sap delivered to above-ground organs by the roots. *Journal of Plant Research* **119**:179–187.

Schmitt M, Boras S, Tjoa A, Watanabe T, Jansen S. 2016. Aluminum accumulation and intra-tree distribution patterns in three *Arbor aluminosa* (*Symplocos*) species from central Sulawesi. *PIOS ONE* **11**:1–18.

Shen R, Ma JF, Kyo M, Iwashita T. 2002. Compartmentation of aluminium in leaves of an Al-accumulator, *Fagopyrum esculentum* Moench. *Planta* **215**:394–398.

Taylor GJ. 1991. Current views of the aluminum stress response: The physiological basis of tolerance. *Current Topics in Plant Biochemistry and Physiology* **10**:57–93.

Tolra R, Vogel-Mikuš K, Hajiboland R, Kump P, Pongrac P, Kaulich B, Gianoncelli A, Babin J, Regvar M, Poschenrieder C. 2011. Localization of aluminium in tea (*Camellia sinensis*) leaves using low energy X-ray fluorescence spectro-microscopy. *Journal of Plant Research* **124**:165–172.

von Uexküll HR, Mutert E. 1995. Global extent, development and economic impact of acid soils. *Plant and Soil* **171**:1–15.

Wang H, Chen RF, Iwashita T, Shen RF, Ma JF. 2015. Physiological characterization of aluminum tolerance and accumulation in tartary and wild buckwheat. *New Phytologist* **205**:273–279.

Wang Y, Fritsch PW, Shi S, Almeda F, Cruz BC, Kelly LM. 2004. Phylogeny and infrageneric classification of *Symplocos* (Symplocosaeae) inferred from DNA sequence data. *American Journal of Botany* **91**:1901–1914.

Watanabe T, Broadley MR, Jansen S, White PJ, Takada J, Satake K, Takamatsu T, et al. 2007. Evolutionary control of leaf element composition in plants. *New Phytologist* **174**:516–523.

Watanabe T, Jansen S, Osaki M. 2005a. The beneficial effect of aluminium and the role of citrate in Al accumulation in *Melastoma malabathricum*. *New Phytologist* **165**:773–780.

Watanabe T, Misawa S, Hiradate S, Osaki M. 2008. Characterization of root mucilage from *Melastoma malabathricum*, with emphasis on its roles in aluminum accumulation. *The New Phytologist* **178**:581–589.

Watanabe T, Misawa S, Osaki M. 2005b. Aluminum accumulation in the roots of *Melastoma malabathricum*, an aluminum-accumulating plant. *Canadian Journal of Botany* **83**:1518–1522.

Watanabe T, Osaki M. 2001. Influence of aluminum and phosphorus on growth and xylem sap composition in *Melastoma malabathricum* L. *Plant and Soil* **237**:63–70.

Yang JL, Zhu XF, Peng YX, Zheng C, Li GX, Liu Y, Shi YZ, Zheng SJ. 2011. Cell wall hemicellulose contributes significantly to aluminum adsorption and root growth in *Arabidopsis*. *Plant Physiology* **155**:1885–1892.

Zeng QL, Chen RF, Zhao XQ, Shen RF, Noguchi A, Shinmachi F, Hosegawa I. 2013. Aluminum could be transported via phloem in *Camellia oleifera* Abel. *Tree Physiology* **33**:96–105.