Lignin in Woody Plants under Water Stress: A Review

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ABSTRACT

Under tropical and sub-tropical conditions, perennial woody plants most often encounter drought stress due to erratic rainfall, which has become compounded as result of recent climatic changes. Control of the efficiency and safety of water transport from roots to shoots is fundamental for plant survival under water stressed conditions. Lignin plays an important role in the life of terrestrial plants. Depending upon the stress, lignin plays protective, sustaining and disruptive roles in addition to its involvement in plant growth, development and defense responses. This review aims to present and summarize reported evidence on the possible role of lignin of plant tissue (in special woody species) in response to water limiting condition which is expected to increase because of environment changes.

Keywords: tree survival, seedling production, tree hydraulics, wood anatomy.

Lignina em Plantas Lenhosas sob Estresse Hídrico: Uma Revisão

RESUMO

Sob condições tropicais e sub-tropicais, plantas lenhosas frequentemente deparam-se com condições de estresse hídrico devido à irregularidade das precipitações pluviométricas, agravada nos dias de hoje pelas mudanças climáticas. O controle da eficiência e segurança no transporte de água das raízes para tecidos aéreos é fundamental para a sobrevivência em condições estressantes. A lignina desempenha um papel importante na vida das plantas terrestres. Dependendo do estresse, a lignina desempenha função protetora, de sustentação, e disruptiva no crescimento, desenvolvimento e respostas de defesa das plantas terrestres. Esta revisão objetiva apresentar e resumir evidências relatadas sobre o possível papel da lignina no tecido vegetal (especialmente em espécies lenhosas) em resposta a condições limitantes de água aumentadas, resultantes de mudanças ambientais.

Palavras-chave: sobrevivência de árvores, produção de mudas, hidráulica de árvores, anatomia do lenho.
1. INTRODUCTION

Damage to plants in response to abiotic and biotic stresses is a worldwide ecologic and economic concern. Recent and predicted climate changes exacerbate this concern (Lobell & Gourdji, 2012). Environmental stresses not only have adverse effects on plant growth and productivity but also are expected to become more variable, severe, and widespread in decades to come. Prolonged and repeated severe environmental stresses affecting plant growth and development would bring down long-lasting effects in woody plants as a result of its long-term growth period. Plant tolerance to stress englobes a variation in the detail network and cascade of events or reactions leading to alleviation of potential stress-induced cellular injuries depending on the plant species that have evolved through environmental changes.

The development of lignin biosynthesis has been considered to be one key factor that allowed plants to flourish in terrestrial ecosystems. Therefore, to understand and improve water stress tolerance in plants by manipulating lignin in wood plant tissues becomes a necessity if tree survival and biomass production are to be increased and profitable in face of future climate changes. Plasticity in lignin biosynthesis may be an important feature in understanding future species distribution as impacted by changing water stress patterns.

2. TERRESTRIAL PLANTS AND WATER STRESS

Terrestrial plants are continuously exposed to changing environmental conditions that could potentially threaten survival. Water availability, which can be constrained by drought, salinity, or freezing is a major factor limiting plant growth and development in both agricultural and forestry settings. The effects of water limitation will likely worsen in the coming decades because of expected climate changes and the growing scarcity of fresh water available for irrigation caused by urbanization and the depletion of aquifers (Pennisi, 2008).

Access to water at the time of tissue development is a major environmental factor limiting plant productivity. The responses of plants to water stress and their relative importance for crop productivity vary with species, soil type, nutrients and climate. One-third of potential arable land suffers from inadequate water supply, and the productivity of the remainder is periodically reduced by drought (Arnell, 2004). In order to avoid limiting availability of water in the soil solution plants have evolved mechanisms to withstand water stress. Those mechanisms may include change in leaf anatomy and ultrastructure, reduction of leaf size, thickening of leaf cell wall, increase in number of large vessels, and reduction of stomata (Lisar et al., 2012).

Plants experience water stress either when the roots face water deficit or because of the high transpiration rate. Controlling the efficiency and safety of water transport from roots to shoots is fundamental for the survival of terrestrial plants under water stressed conditions. With changes predicted in seasonal drought patterns, especially under climate change, traditional/evolved mechanisms of drought tolerance or avoidance may no longer be uniquely able to guide plants to survival (De Micco & Aronne, 2012).

Water stress resulting from the withholding of water not only changes the physical environment for plant growth but also its physiology. Widely different plant responses have been reported depending on how the water stress is imposed to plants. Short-versus rapid-term and long-versus gradual-term essays results in a variety of responses. Prolonged and repeated severe stresses affecting growth and development would bring down long-lasting effects in woody plants as a result of its long-term growth period (Osakabe et al., 2012).

Terrestrial plant species possess distinctive indicators of stress tolerance at whole plant, tissue or cellular levels (Munns, 2002). Most adult woody plants can withstand periods of soil water deficits. However, seedlings of those very same species often cannot because of shallow root systems relative to available soil water (Pinto et al., 2012).

3. PLANT XYLEM ANATOMY

Water is transported from the roots to the leaves through the xylem of woody plants under negative pressure (Tyree, 1997). When water levels decrease, soil water potentials drop, pressure gradient in the xylem increases, and vessels are more likely to cavitate and lose their hydraulic conductivity due to water stress. Uninterrupted transport of water through the xylem
is essential for plant growth and survival because it replaces water lost by transpiration and allows stomata to remain open for photosynthesis (Ambrose et al., 2015).

Xylem cavitation can affect plant productivity and survival (McDowell et al., 2008; Brodribb & Cochard, 2009) while xylem vulnerability to cavitation is correlated with drought tolerance (Maherali et al., 2004; Pockman & Sperry, 2000; Tyree et al., 2003). However, lower vulnerability to cavitation is correlated with lower hydraulic conductivity (Cochard et al., 1992; Hacke et al., 2006).

The trade-off between safety and efficiency of plant water transport will depend on the pit area and the pit membrane structure. Inter-vessel pits, composed of small and thin porous membrane with overarching secondary cell walls are the main pathway for the water flow that prevent the passage of air bubbles from an embolized vessel to a conducting vessel (Jansen et al., 2009).

The extent in which wood plant species can conduct water and resist xylem cavitation in the stem is determined by vessel adaptation. Failure of the conductive tissue to resist high negative pressures can result in collapse of the conduit walls resulting in cavitation. Formation of gas bubbles, or embolism that cause blockage to the flow of water within conduits can result from cavitation. Embolism can be reversed (Shen et al., 2007) and may also play a developmental role in the formation of heartwood (Sperry et al., 1991).

Wide vessels have the advantage of a greater hydraulic efficiency (i.e., water transport capacity) because hydraulic efficiency of a xylem vessel increases with the fourth power of its diameter (Tyree & Zimmermann, 2002).

Terrestrial plants may also be able to reduce the risk of conduits imploding in response to the tensile forces acting on the walls by reducing the lumen diameter relative to the wall thickness of vessels (Hacke et al., 2001) or fibers (Jacobsen et al., 2005). This decrease in lumen diameter could reduce wood porosity. As a result, acclimation to water stress in trees is often associated with an increase of wood density (Al-Khalifah et al., 2006).

Denser wood shows thicker cell walls and stronger mechanical properties. Aref et al. (2013) studying anatomical adaptive changes of two Acacia woody species concluded that vessel-wall thickness in conjunction with inter-vessel pit membrane thickness showed a positive correlation with water stress in both species. Conversely, Awad et al. (2012) testing ten lines of Populus trema × Populus alba with modified gene expression of lignin metabolism reported no evidence of the mechanical versus hydraulic trade-off. Instead, they concluded that angiosperm wood species may have numerous ways to accommodate their mechanical properties without interfering with plant hydraulic characteristics.

Testing the relationship between water stress acclimation and mechanical properties in seedlings of three woody species, Christensen-Dalsgaard & Ennos (2012) emphasized the need for further studies relating physio-chemical parameters and lignin to terrestrial plant hydraulics.

4. THE ROLE OF LIGNIN IN TERRESTRIAL PLANTS

Lignins are found widely distributed in both angiosperms and gymnosperms (Figure 1). While in gymnosperms lignins present guaiacyl and syringyl, angiosperms also have all three types of lignins i.e., syringyl, hydroxy-phenyl and guaiacyl. Published estimates suggest that lignin represent about 25% of total plant terrestrial biomass (Boudet et al., 1995). Lignin is a 3-dimensional polymer which is the principal structural component of plant cell wall.

Physiologically, lignin plays an important aspect throughout the life of plants. Lignin affects seed coat which in turn modulates seed dispersal and dehiscence in many plant species (Liljegren et al., 2000). Lignins are also involved in fixation of atmospheric carbon dioxide by forming secondary thickenings of mature terrestrial plants via carbon cycle and by decomposing at an extremely slow rate that replenishes soil fertility and texture. Furthermore, lignification helps in binding plant fibers providing mechanical support to terrestrial plants (Li & Chapple, 2010).

The synthesis of lignin represents one of the most energy demanding biosynthetic pathway in plants requiring large amounts of carbon skeletons (Amthor, 2003). Lignin is composed of p-hydroxyphenyl (H), guaiacyl (G), and or syringyl (S) units, the proportions of which vary according to botanical and cytological aspects (Lewis & Yamamoto, 1990). The glucose requirements for biosynthesis of the three monolignol subunits (i.e., H, G and/or S) are thought to vary, with
H lignin subunit requiring the least amount of glucose 2.473 g of glucose per gram of lignin, followed by G lignin subunit 2.547 g of glucose per gram of lignin, and lastly S lignin subunit requiring 2.600 g of glucose per gram of lignin (Amthor, 2003).

5. THE LIGNIFICATION PROCESS

Lignification is a dynamic physiological process subject to modulation at different levels during normal plant development in response to several environment stresses (Boudet, 2000). The incorporation of lignin into cell walls results in structural rigidity and durability of plant tissues (Lee et al., 2007). Lignin production has been the target of genetic engineering in order to facilitate the production of wood-pulp and biofuel (Vanholme et al., 2008). Several transcription factors controlling lignin biosynthesis genes in response to developmental processes were identified (Osakabe et al., 1999; Boudet et al., 2003; Boerjan et al., 2003; Raes et al., 2003).

Lignin is found in the cell walls of specific tissue types such as traqueids and vessel elements of xylem (Esau, 1967). These cells are characterized by the presence of a specialized cell wall known as the lignified secondary cell wall. During water stress, the cell wall flexibility is affected by the levels of lignin and phenolic compounds (Moore et al., 2008). Within terrestrial plants, lignin content can vary greatly in different tissues. Lignin provides a hydrophobic surface allowing plants to transport water to heights of several meters (Carder, 1995; Koch et al., 2004) as well as contributes to the organism mechanical strength that make trees to support large amounts of biomass (Jagels et al., 2003).

On the other hand, lignin is a limiting factor in a number of agro-industrial processes such as chemical pulping and the processing of lignocellulosic plant biomass to produce bioethanol, as well as modulates forage digestibility. Lignin is very low in young shoots and high in wood. In different tree species, lignin content of wood can vary from 15% to 40% (Sarkanen & Ludwig, 1971) while within a species the average lignin content of wood is much less variable. The manipulation of lignin genes in mutants or transgenic plants suggests the existence of a high degree of plasticity (Lu et al., 2010) despite the fact that modifications are sometimes associated with a strong perturbation in the plant developmental program. In an individual plant, lignin content and composition can vary greatly between different tissues but are much less variable in the same cell wall.
tissue when plants are cultivated in standard conditions (Novaes et al., 2010).

6. LIGNIN AND PLANT FITNESS

In woody plants, secondary stem growth determines plant size, and responds to physiological and environmental signals. The large amount of assimilated carbon is both accumulated and stored in the secondary cell wall of woody plants, mostly as cellulose and lignin (Osakabe et al., 2012). Stress affects growth and development because plants need to re-allocate energy to adapt to the limiting condition (Skirycz & Inze, 2010).

According to Pedersen et al. (2005), it appears that reducing lignin content of crop plants can impact their agricultural fitness by reducing lodging resistance, disease resistance and water stress tolerance. However, the relationship among drought tolerance, post-drought recovery and lignin metabolism in plants have not been well revealed (Li et al., 2013).

While lignin biosynthesis is well characterized, the mechanism regulating lignin deposition in response to environmental stresses lacks agreement. The expression levels of the genes encoding cinnamoyl-CoA reductase, which is involved in lignin biosynthesis, increased after drought and localized changes in lignification were involved in the acclimation to drought stress (Fan et al., 2006) and is directly responsible for reduced root colonization by ectomycorrhizae fungi under fertilized conditions (Kleckewski et al., 2010).

In angiosperms, the water-conductive xylem vessel elements of secondary xylem and the primary xylem cells are rich in or contain only guaiacyl (G) lignin subunit, like the tracheids of the more ancient gymnosperms whereas the nonconductive xylem fiber cells are rich in syringyl (S) lignin subunit (Li et al., 2001). The fact that water-conducting cells in both gymnosperms and angiosperms are principally comprised of G lignin subunit suggests a strong selective pressure to conserve the pathway for and the regulation of G lignin subunit biosynthesis in the water conducting cells of xylem during land plant evolution (Peter and Neale, 2004). Moreover, transgenic poplars with severe reductions in xylem lignin contents were shown to have a reduced efficiency in water transport (Coleman et al., 2008; Kitin et al., 2010).

Evolution of xylem structure and architecture has been extensively examined from both functional and ecological perspectives (Bhaskar et al., 2007; Meinzer et al., 2010; Sperry et al., 2008). However, little advance has been achieved on the role of cell wall chemistry and in special lignin content and composition in how xylem traits or how xylem evolution affected the deposition and composition of lignin (Boyce et al., 2004).

7. LIGNIN AND PLANT HYDRAULICS

Support for the hypothesis that there exist a trade-off between wood strength or stiffness and xylem hydraulic function have been based on essays consisting of surveys across species and environmental gradients (Gartner, 1991a, b; Christensen-Dalsgaard et al., 2007; Jacobsen et al., 2007a, b; Pratt et al., 2007). Rarely addressed is how differences in lignin content may affect wood anatomical or hydraulic properties (Koehler et al., 2006; Horvath et al., 2010).

Results of Terzi et al. (2013) suggested that an increase of water stress could induce lignification in rolled leaves of Ctenanthe setosa. In addition, rubbing the internodes of plants have reduced elongation with alteration of cell wall mechanical properties and accelerated lignification (De Jaegher et al., 1985).

The induction of lignin in plants has been correlated stresses such as cold, drought or light as well as mechanical injuries in a number of plant species such as poplar, rice, pine, Arabidopsis and soybean (Moura et al., 2010). Moura-Sobczak et al. (2011) reported that Eucalyptus urograndis subjected to drought decreased the amount of lignin in the stem apical regions and increased lignin in the basal region while E. globulus showed an opposite behavior in apical regions and showed no significant changes in the basal regions. A hybrid between both species (E. urograndis) showed a pattern similar to E. urograndis in apical regions and similar to E. globulus in the basal regions.

Evidence is accumulating that a cell wall integrity sensing and maintenance mechanism exists in plants that coordinates the response to cell wall damage (Seifert & Blaauw, 2010). Mechanical stress can have a strong impact on the cell wall composition and architecture. For example, mechanical bending of stems or branches in some angiosperms induces the formation of tension wood containing lignin with an increased S/G ratio (Aoyama et al., 2001).
Additionally, there is evidence that expression of lignin biosynthetic genes can be correlated with lignin production in response to infection by pathogens such as in Arabidopsis exposed to Xanthomonas campestris, and in *Linum usitatissimum* cell cultures inoculated with *Botrytis cinerea* (Lauvergeat et al., 2001; Hano et al., 2006).

**8. FINAL CONSIDERATIONS**

Recent publications have reported that reduction in total lignin content can have a drastic effect on anatomical and physiological traits including reduced growth (Voelker et al., 2010) increased frequency of vessel collapse (Coleman et al., 2008; Kitin et al., 2010) and decreased hydraulic conductivity (Coleman et al., 2008).

Plants present a variety of lignin coordination techniques for monitoring synthesis, accumulation and deposition in response to various intrinsic and extrinsic signals. Lignin plays protective, sustaining and disruptive roles depending upon the situation or stress. Lignin is involved in fixing atmospheric carbon dioxide into secondary thickenings of adult terrestrial plants via the carbon cycle, and decomposes at a low rate compared to most components of dead plant tissues, contributing as a major constituent of humus which maintains soil fertility and texture.

During the lignification process, plant growth is inhibited because the initial enzyme phenylalanine ammonia lyase that is involved in monolignol biosynthesis pathway directly influences accumulation of lignin. Apart from providing mechanical strength, lignin content alteration has been cited during stressed conditions, thereby suggesting that various abiotic stresses have the ability to express the genes responsible for their synthesis.

The potential of altering plant lignin content by scientists and tree managers must be explored in the context to changing environment and anti-stress response triggered by environment stresses.

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