A Study of the Molecular Pathways of Wood Formation using Peptide Mass Spectrometry

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Cover: From transcription to cell wall synthesis
(photo: R. Nilsson & K. Olofsson)
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Abstract
Wood consists essentially of secondary plant cell walls, composed of a cellulose and hemicellulose network impregnated with lignin. It is formed via the development of derivatives of a thin layer of cells (the vascular cambium) into xylem cells through cell division, expansion, secondary wall formation, lignification and, finally, programmed cell death. Throughout the formation of the resulting, complex matrix (wood) diverse proteins are involved. Hence, biosynthesis of wood is tightly regulated at the molecular level, primarily by transcriptional regulation, together with various post-transcriptional processes.

In the studies this thesis is based upon, a key technique for studying proteins, liquid chromatography coupled to mass spectrometry, was used to elucidate pathways of wood formation. First, proteins involved in the central transcriptional unit Mediator, regulating most essential processes in plants, were identified in arabidopsis. Mediator subunits that most likely provide secondary cell wall activity are differentiated paralogous subunits located in the tail part of the complex. Point mutations in the Mediator tail subunit, Med5#2 affect secondary cell wall lignification. Moreover, control of xylem expansion may be indirectly regulated by Med25. A strategy to integrate transcript-, protein- and metabolite-data was then developed, using data acquired from analyses of poplar mutants with perturbed wood development. The results showed that lignin biosynthesis is heavily affected at both transcript and protein levels in the mutant with the strongest phenotypic deviations. Interestingly, when transcript levels are decreased, the protein levels are increased. Finally, the front of cell wall synthesis at poplar plasma membranes was dissected. A high coverage of proteins known to be involved in the cell wall synthesizing machinery and associated components was captured, as well as several potential new ones.

Keywords: proteomics, transcription, xylem, mediator, regulation, plasma membrane, development, arabidopsis, poplar, *Populus*, mass spectrometry

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E-mail: Robert.Nilsson@genfys.slu.se
Varför ska man göra lätt för sig, när det är så lätt att göra svårt för sig.

Ronny Eriksson
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List of Publications

This thesis is based on the work contained in the following papers, referred to by Roman numerals in the text:


* To be considered joint first authors

Papers are reproduced with the permission of the publishers.
The contribution of Robert Nilsson to the papers included in this thesis was as follows:

I Planned identification analysis jointly with co-authors, extracted and analysed peptides using mass spectrometry, evaluated the data, and assisted in the writing of the paper.

II Planned design of experiments, extracted proteins and metabolites, analysed peptides using mass spectrometry, evaluated the data, and wrote the paper jointly with co-authors.

III Planned design of protein identification experiments, extracted and analysed peptides using mass spectrometry, evaluated the data, and wrote the paper jointly with co-authors.
## Abbreviations

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<th>Description</th>
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<tr>
<td>4CL</td>
<td>4-coumarate:CoA ligase</td>
</tr>
<tr>
<td>AMBIC</td>
<td>ammonium bicarbonate</td>
</tr>
<tr>
<td>Ara</td>
<td>arabinose</td>
</tr>
<tr>
<td>At</td>
<td><em>Arabidopsis thaliana</em></td>
</tr>
<tr>
<td>C3H</td>
<td>4-coumarate 3-hydroxylase</td>
</tr>
<tr>
<td>C4H</td>
<td>cinnamate-4-hydroxylase</td>
</tr>
<tr>
<td>CAD</td>
<td>cinnamyl alcohol dehydrogenase</td>
</tr>
<tr>
<td>CAld5H</td>
<td>coniferyl aldehyde 5-hydroxylase</td>
</tr>
<tr>
<td>CAZyme</td>
<td>carbohydrate-active enzyme</td>
</tr>
<tr>
<td>CCoAOMT</td>
<td>caffeoyl-CoA O-methyltransferase</td>
</tr>
<tr>
<td>CCR</td>
<td>cinnamoyl-CoA reductase</td>
</tr>
<tr>
<td>CE</td>
<td>carbohydrate esterase</td>
</tr>
<tr>
<td>CESA</td>
<td>cellulose synthase</td>
</tr>
<tr>
<td>CID</td>
<td>collision-induced dissociation</td>
</tr>
<tr>
<td>COMT</td>
<td>caffeic acid/5-hydroxyconiferaldehyde O-methyltransferase</td>
</tr>
<tr>
<td>CSC</td>
<td>cellulose synthase complex</td>
</tr>
<tr>
<td>DNA</td>
<td>deoxyribonucleic acid</td>
</tr>
<tr>
<td>DoE</td>
<td>Design of Experiments</td>
</tr>
<tr>
<td>EMS</td>
<td>ethyl methanesulfonate</td>
</tr>
<tr>
<td>EST</td>
<td>expressed sequence tag</td>
</tr>
<tr>
<td>Fuc</td>
<td>fucose</td>
</tr>
<tr>
<td>Gal</td>
<td>galactose</td>
</tr>
<tr>
<td>GalUA</td>
<td>galacturonic acid</td>
</tr>
<tr>
<td>GC</td>
<td>gas chromatography</td>
</tr>
<tr>
<td>GH</td>
<td>glycoside hydrolase</td>
</tr>
<tr>
<td>Glc</td>
<td>glucose</td>
</tr>
<tr>
<td>GlcUA</td>
<td>glucuronic acid</td>
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<tr>
<td>GME</td>
<td>GDP-D-mannose 3,5-epimerase</td>
</tr>
<tr>
<td>Acronym</td>
<td>Full Form</td>
</tr>
<tr>
<td>---------</td>
<td>-----------</td>
</tr>
<tr>
<td>GPI</td>
<td>glycosylphosphatidyl inositol</td>
</tr>
<tr>
<td>GT</td>
<td>glycosyl transferase</td>
</tr>
<tr>
<td>GTF</td>
<td>general transcription factor</td>
</tr>
<tr>
<td>HCT</td>
<td>p-hydroxycinnamoyl-CoA:quinate shikimate p-hydroxycinnamoyltransferase</td>
</tr>
<tr>
<td>HTM</td>
<td>heat and trapping module</td>
</tr>
<tr>
<td>IEF</td>
<td>isoelectric focusing</td>
</tr>
<tr>
<td>LC</td>
<td>liquid chromatography</td>
</tr>
<tr>
<td>Man</td>
<td>mannose</td>
</tr>
<tr>
<td>miRNA</td>
<td>microRNA</td>
</tr>
<tr>
<td>mRNA</td>
<td>messenger RNA</td>
</tr>
<tr>
<td>MS</td>
<td>mass spectrometry</td>
</tr>
<tr>
<td>MS/MS</td>
<td>tandem mass spectrometry</td>
</tr>
<tr>
<td>NAD</td>
<td>nicotinamide adenine dinucleotide</td>
</tr>
<tr>
<td>O2PLS</td>
<td>bidirectional OPLS</td>
</tr>
<tr>
<td>OPLS</td>
<td>orthogonal projections to latent structures</td>
</tr>
<tr>
<td>PAL</td>
<td>phenylalanine ammonia-lyase</td>
</tr>
<tr>
<td>PCA</td>
<td>principal component analysis</td>
</tr>
<tr>
<td>PCR</td>
<td>polymerase chain reaction</td>
</tr>
<tr>
<td>PIC</td>
<td>preinitiation complex</td>
</tr>
<tr>
<td>pre-mRNA</td>
<td>precursor mRNA</td>
</tr>
<tr>
<td>Ptr</td>
<td><em>Populus trichocarpa</em></td>
</tr>
<tr>
<td>Ptt</td>
<td><em>Populus tremula x tremuloides</em></td>
</tr>
<tr>
<td>Q-ToF</td>
<td>quadrupole time-of-flight</td>
</tr>
<tr>
<td>RG</td>
<td>rhamnogalacturonan</td>
</tr>
<tr>
<td>Rha</td>
<td>rhamnose</td>
</tr>
<tr>
<td>RHM</td>
<td>UDP-L-rhamnose synthase</td>
</tr>
<tr>
<td>RNA</td>
<td>ribonucleic acid</td>
</tr>
<tr>
<td>RNAi</td>
<td>RNA interference</td>
</tr>
<tr>
<td>RNAP</td>
<td>RNA polymerase</td>
</tr>
<tr>
<td>rRNA</td>
<td>ribosomal RNA</td>
</tr>
<tr>
<td>SAD</td>
<td>sinapyl alcohol dehydrogenase</td>
</tr>
<tr>
<td>SDS-PAGE</td>
<td>sodium dodecyl sulfate polyacrylamide gel electrophoresis</td>
</tr>
<tr>
<td>siRNA</td>
<td>short interfering RNA</td>
</tr>
<tr>
<td>UGE</td>
<td>UDP-D-glucose 4-epimerase</td>
</tr>
<tr>
<td>UPLC</td>
<td>ultra performance liquid chromatography</td>
</tr>
<tr>
<td>UXS</td>
<td>UDP-D-apiose/xylose synthase</td>
</tr>
<tr>
<td>WT</td>
<td>wild-type</td>
</tr>
<tr>
<td>Xyl</td>
<td>xylose</td>
</tr>
</tbody>
</table>
1 Introduction

55% of Sweden's surface area is covered by productive forest (Anon., 2008). Although the growth of this dense forest is much less rapid than that of its tropical counterparts, its wood can be profitably refined by using advanced management, harvesting, and transport techniques. When refining wood, the quality and value of the final products are determined by the composition of the major cell-wall polymer (a blend of cellulose, hemicellulose and lignin). Differences in the precise constitution of the hemicellulose mixture aside, the normal composition of hardwoods (such as poplar, birch, or oak) is comparable to that of softwoods such as pine and spruce (Pauly & Keegstra, 2008). Traditional breeding and genetic engineering can be used to alter the constitution and composition of the wood and thereby facilitate its refinement. However, such alterations may compromise the biological functionality of the cell wall by reducing its pathogen resistance, its ability to withstand mechanical stress, or its capacity for intracellular transport or cell signaling; such potential problems must be addressed. Field trials (or mixed field/greenhouse studies) are necessary to characterize the new phenotypes and expose pleiotropic effects. Potentially undesirable side-effects can be minimized in various ways. First, even relatively minor changes in wood chemistry can result in dramatic improvements to the refining process; one may not need to significantly disrupt the existing functionality of the system to achieve desirable results. Second, benefits can be sometimes obtained even if the altered trees constitute only a small fraction of the total feedstock, and they can be propagated in greenhouses or restricted fields. An alternative method to avoid disrupting existing traits that are important in survival and fitness would be to allow the tree to grow normally and only induce the desired changes shortly before it is to be harvested. An inducible gene expression system could be used to turn on wood modifying processes at the desired
time in a specific cell type or tissue. At present, there are only a few such systems that are both suitable for field trials and commercially viable (Corrado & Karali, 2009). Basic knowledge of how wood formation is regulated can also be applied in the development of improved methods for the pretreatment of feedstocks for the pulp and paper industries, for biofuel production and for other renewable materials. Currently, most research on trees is performed on the angiosperm poplar (Populus spp.) because of its relatively small genome compared to other trees. In addition, poplar is an attractive model system because its genome has been sequenced, various tools for studying it have been developed, and its growth and vegetative propagation are both rapid (Jansson & Douglas, 2007; Taylor, 2002). Despite its small genome, poplar has a more complex wood structure than do softwoods, in that it has additional vessel elements. The relatively close phylogenetic relationship between poplar and the primary plant model system, arabidopsis (Arabidopsis thaliana), makes comparative studies possible.

Paper I describes a study of a gene regulation system that is common to both poplar and arabidopsis, and to plants in general, using arabidopsis as the model system. Many of arabidopsis' genes have been well characterized; because of its rapid generation time, it has also lost some redundant genes that are essential in other plants (Jansson & Douglas, 2007). The remainder of the work described in this thesis (papers II and III) focused on studying wood formation, using poplar as the model system. In this context, it should be noted that by using a combination of short- and long-day conditions, in conjunction with regular removal of the inflorescence stems, arabidopsis can be forced to produce extensive secondary xylem (Demura & Fukuda, 2007).

1.1 Wood formation

The formation of wood (xylogenesis) involves cell division, cell expansion, cell wall thickening and programmed cell death (PCD). Additionally, the deposition of an aromatic polymer (lignification) in the secondary cell wall makes it rigid and impermeable. After this extensive synthesis and deposition, the mature wall can represent over 95% of the cell's dry weight, with the cell wall proteins accounting for only 5-10% of this (Brett & Waldron, 1996; Cassab & Varner, 1988).

1.1.1 Cell differentiation and physiology

Wood formation in the trunk starts at the shoot apical meristem (SAM) of the young plant. The shoot apical meristems develop a cylindrical lateral meristem called the vascular cambium that is able to generate all of the cell
types necessary for wood formation. Wood formation occurs simultaneously
and in parallel in the roots and the branches. The vascular cambium contains
stem cells known as the initials, as well as partially differentiated phloem and
xylem mother cells. There are two types of initials. Radial initials give rise
to ray cells, which are essential for the translocation of nutrients between
phloem and xylem and the storage of nutrients. Fusiform initials, by
contrast, form xylem and phloem cells. The xylem cells are further
subdivided into tracheary elements, vessel elements (only found in
angiosperms), xylem fibres and axial parenchyma.

The earliest stages of wall formation (cell plate formation) can be
monitored during cell division. The cell plate grows from the centre of the
cell out towards the pre-existing cell walls, and fuses with them. The zone
in which plate formation occurs is called the phragmoplast; it possesses an
extensive network of microtubules that are involved in the transport of
vesicles from the endomembrane system. The vesicles carry the proteins and
other materials that are used to build the cell plate, and lipids to generate its
plasma membrane. At the same time, channels (plasmodesmata) form around
some of the microtubules linking the two nascent daughter cells. These
microtubules may also form a desmotubule, a tube of membrane that spans
the plasmodesma and is continuous with the endoplasmic reticulum. As
cytokinesis proceeds, the phragmoplast loses definition and the cell plate
joins with the parental cell wall to form the middle lamellae. The separated
daughter cells deposit further material to generate a primary wall. The
structurally-important cellulose microfibrils are usually randomly or
longitudinally oriented in the primary wall.

The cells expand during the primary wall stage. Fibers and axial
parenchyma cells expand primarily in radial direction while vessel elements
also exhibit substantial tangential growth. Fiber elongation is achieved by
intrusive tip growth that requires wall biogenesis and dissolution of the
middle lamellae between neighboring cells. Cell wall enlargement is
regulated by the cell's turgor pressure, growth substances that alter the pH of
the wall, and by the turnover and cross-linking of wall components,
mediated by enzymes and various ligands.

As the xylem cells reach their final size, secondary wall formation takes
place. Three distinct layers, denoted S1, S2 and S3, form inside the primary
wall. The S1 layer is formed first. Initially, its microfibril angle is
approximately perpendicular to the cell's long axis; as the layer grows, this
angle increases in a clockwise direction (as seen from the cell lumen),
eventually reaching the longitudinal arrangement that characterizes the
thickest S2 layer. In poplar the S2 layer is proportionally thinner in the
vessel elements than in the fibres. Finally, the microfibril angle undergoes a second clockwise increase, restoring the transverse arrangement in the S3 layer; overall, the fibrils describe a transverse helix spanning the three layers. The changes in microfibril orientation during the formation of successive cell wall layers are always paralleled by the reorientation of cortical microtubules. An additional tertiary layer called the protective layer is formed within the contact ray cells after autolysis of the neighboring vessel element (Déjardin et al., 2010; Mellerowicz et al., 2001; Brett & Waldron, 1996; Larson, 1994).

Distinctive structures are formed in reaction wood, which is generally formed in xylem tissue in response to mechanical stress of the stem. In poplar, the reaction wood takes the form of tension wood; its formation is characterized by an increased rate of cell division on the tension wood side, and reduced wood formation on the opposite side of the stem. On the tension wood side, the vessels are reduced in size and density and the fibers form a gelatinous layer (G layer) over the inner side of the secondary cell wall (Mellerowicz et al., 2001).

The last stage of wall formation is programmed cell death. At this stage the tonoplast (vacuole membrane) ruptures and releases hydrolytic enzymes into the cytoplasm to degrade the cell's contents. The cell walls are left intact, and constitute the wood.

The process of cell wall lignification occurs in parallel with the final stages of xylem cell differentiation but is not directly connected to programmed cell death. Lignified parenchyma cells remain alive for several years (Déjardin et al., 2010). The extent of lignification can vary significantly among cells within a tissue; the most extensive deposition is normally observed in vessels (Donaldson et al., 2001). Lignification proceeds in different phases. Initially, following the initiation of S1, the corners of the cell walls become lignified, along with the middle lamella. After the S2 layer is completed, lignification proceeds further, and massive deposition occurs when the S3 layer is finished. Ultimately, the concentration of lignin within the wall is highest in the middle lamella and cell corners, and is relatively low in the S2 layer. However, the overall lignin content of the secondary wall is greater than that of other regions due to its larger volume (Boerjan et al., 2003).

1.1.2 Biochemical composition of wood

Wood is a composite material with two phases, a microfibrillar phase and a matrix phase. The microfibrillar phase has a high degree of crystallinity and a homogenous chemical composition: it consists of cellulose microfibrils. The
The chemical composition of the non-crystalline matrix phase is more complex. It consists of a variety of polysaccharides (hemicelluloses), proteins and phenolic compounds (lignin) (Brett & Waldron, 1996).

The most abundant component of the secondary wall of poplar is cellulose (43–48%), while pectin is the most abundant component of the primary wall (~47%). The cell plate is probably made of callose when it is first formed, but is re-made from cellulose at later developmental stages (as observed in tobacco) (Mellerowicz et al., 2001). The biopolymer compositions of the primary and secondary cell walls of poplar are shown in Figure 1.

![Biopolymer composition](image)

**Figure 1.** Biopolymer composition (% dry weight) of the primary and secondary cell walls of poplar. Adapted from Mellerowicz et al. (2001).

The detailed compositions of the major biopolymers (in terms of their constituent monomers) in the primary and secondary cell walls of poplar are compiled in Table 1.
Table 1. Monomer composition of the main components of the primary and secondary cell walls of poplar, from Takahashi Schmidt thesis (Schmidt, 2008).

<table>
<thead>
<tr>
<th>Polysaccharides or lignins</th>
<th>Occurrence</th>
<th>Major monomers</th>
<th>Backbone</th>
<th>Substituents</th>
</tr>
</thead>
<tbody>
<tr>
<td>Cellulose</td>
<td>Primary and secondary cell wall</td>
<td>β-Glc</td>
<td>β-1,4-Glc</td>
<td>Unbranched.</td>
</tr>
<tr>
<td>Xyloglucan</td>
<td>Mainly primary cell wall</td>
<td>β-Glc, α-Xyl, β-Gal, α-Fuc</td>
<td>β-1,4-Glc</td>
<td>α-Xyl-1,6-β-Glc, β-Gal-1,2-α-Xyl-1,6-β-Glc, α-Fuc-1,2-β-Gal-1,2-α-Xyl-1,6-β-Glc. β-Gal is partially acetylated.</td>
</tr>
<tr>
<td>Glucuronoxylan</td>
<td>Mainly secondary cell wall</td>
<td>β-Xyl, α-GlcUA</td>
<td>β-1,4-Xyl</td>
<td>α-GlcUA-1,2-β-Xyl. Some α-GlcUA are as 4-O-methyl ether. Some β-Xyl are acetylated.</td>
</tr>
<tr>
<td>Glucomannan</td>
<td>Mainly secondary cell wall</td>
<td>β-Glc, β-Man</td>
<td>β-1,4-Glc/β-1,4-Man (ratio 1:2)</td>
<td>Unbranched.</td>
</tr>
<tr>
<td>Pectin Homogalacturonan</td>
<td>Primary cell wall</td>
<td>α-GalUA</td>
<td>α-1,4-GalUA</td>
<td>Unbranched. Some α-GalUA are as methyl ester.</td>
</tr>
<tr>
<td>Pectin RG-I</td>
<td>Primary cell wall</td>
<td>α-GalUA, α-Rha, β-Gal, α-Ara, α-Fuc</td>
<td>α-1,4-GalUA-α-1,2-Rha</td>
<td>Rich in α-Ara and/or β-Gal, attached to O-4 of α-Rha.</td>
</tr>
<tr>
<td>Pectin RG-II</td>
<td>Primary cell wall</td>
<td>Eleven different glycosyl residues. Usual sugars are α-GalUA, β-Rha, α-Gal, α-Fuc, α-Rha, β-GalUA, α-Ara, β-GlcUA.</td>
<td>At least eight α-1,4-GalUA</td>
<td>Two structurally distinct disaccharides attached to C3 and two oligosaccharides attached to C2 of the backbone.</td>
</tr>
<tr>
<td>Lignins</td>
<td>Middle lamella, primary cell wall (higher conc.) and secondary cell wall (lower conc.) during secondary cell wall stage.</td>
<td>Guaiacyl (G) and syringyl (S) units and traces of p-hydroxyphenyl (H) phenylpropanoid units.</td>
<td></td>
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</tr>
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</table>
1.2 The central dogma and beyond

The central dogma of molecular biology states that sequential information in DNA is transferred by transcription to RNA and then by translation to protein in a unidirectional process (Crick, 1970). These processes have been studied extensively since the dogma was first articulated by Francis Crick in 1958. Recent work has shown that these processes are much more complex than the dogma implies, and that many more actors are involved than was initially thought, including microRNAs (miRNAs) and short interfering RNAs (siRNAs) (Carthew & Sontheimer, 2009; Vaucheret, 2006). In addition, DNA carries numerous regulatory elements, both within and outside genes and promoter regions; these regulatory elements are the focus of much current research. The proteins involved in these processes are often incorporated into large and dynamic multi-protein complexes (Sikorski & Buratowski, 2009). The most important enzyme complexes in the transcription of genomic DNA to RNA are the RNA polymerases (RNAPs).

1.2.1 Transcriptional regulation

In eukaryotes, there are three nuclear polymerases: RNAP I, which transcribes the ribosomal RNA (rRNA); RNAP II, which handles the bulk of the transcription of genes to precursor messenger RNA (pre-mRNA) and also that of most small nuclear RNA (snRNA) and miRNA; and RNAP III, which transcribes genes that encode short structural RNAs, including transport RNAs (tRNAs) and 5S rRNA. Plants have two additional polymerases, RNAP IV and V, which mediate siRNA-directed DNA methylation and gene silencing (Ream et al., 2009; Wierzbicki et al., 2009; Onodera et al., 2005).

RNAP II transcribes all protein-coding genes. It consists of 12 proteins in humans, yeast and plants (Ream et al., 2009; Myer & Young, 1998). To initiate transcription, RNAP II needs to bind to a number of other proteins to form the preinitiation complex (PIC). Minimally, transcription in vitro requires the presence of the basal initiation factors TFIIA, TFIIB, TFIID, TFIIE, TFIIF, and TFIH. These components are called general transcription factors (GTFs). The Mediator protein complex transmits information from regulatory DNA elements to core promoters and the basal machinery (Kim et al., 1994; Koleske & Young, 1994) (Figure 2). There is some debate over whether Mediator should also be classified as a GTF (Sikorski & Buratowski, 2009).
Mediator was first identified as being necessary for transcriptional activation \textit{in vitro} in yeast (\textit{S. cerevisiae}) (Flanagan et al., 1991; Kelleher et al., 1990). The yeast Mediator complex consists of 25 proteins, comprising four subdomains: head, middle, tail and kinase (Asturias et al., 1999). The kinase domain is only associated with free Mediator (i.e. Mediator that is not bound to RNAP II) (Samuelsen et al., 2003). The head and middle domains bind to the C-terminal domain of RNAP II while the tail interacts primarily with DNA-bound transcription factors. When so bound, the Mediator can act as a molecular switchboard and mediate signals from hormones, receptors and transcription factors (van den Berg et al., 2010; Khidekel & Hsieh-Wilson, 2004). A key feature of Mediator is its ability to regulate both positive and negative signalling pathways. Different subunits have been shown to act antagonistically on the same genes in yeast. Several Mediator subunits in yeast are essential, primarily those located in the head domain (van de Peppel et al., 2005). Metazoan Mediators have been identified in \textit{D. melanogaster}, \textit{C. elegans}, mouse, and human cells. They share 22 subunits with the yeast Mediator (Myers & Kornberg, 2000) but also contain eight subunits (Med 23–Med 30) that are not found in \textit{S. cerevisiae}. Finally, Mediator has also been purified from the fission yeast \textit{S. pombe}, which contains a somewhat smaller Mediator. However, 19 of the shared metazoan and \textit{S. cerevisiae} Mediator subunits and one of the metazoan–specific Mediator subunits are conserved in the \textit{S. pombe} Mediator. These results led to a unified nomenclature for Mediator subunits in all species (Bourbon et
Analysis of the genomic sequence of the model plant *arabidopsis* revealed that only 8%–23% of the proteins involved in transcription are related to genes found in other eukaryotic genomes, indicating that many plant transcription proteins evolved independently. In contrast, 48%–60% of the *arabidopsis* genes that encode translation proteins have counterparts in other eukaryotic genomes, indicating that the functions of the corresponding proteins are highly conserved. It is notable that >5% of the *arabidopsis* genome codes for ~1700 transcription factors (TFs), about 41% of which are from families specific to plants. It is therefore likely that coregulators that function at the interface between the rapidly evolving regulatory TFs (activators and repressors) and the more conserved general transcription machinery should contain both conserved and species-specific domains.

*Arabidopsis* also contains several tandem gene duplications and large scale duplications on different chromosomes, accounting for >60% of the genome (Blanc et al., 2000; Riechmann & Ratcliffe, 2000; Lin et al., 1999; Mayer et al., 1999). Some duplicated sequences subsequently underwent rearrangement and divergent evolution, but still, up to 40% of the *arabidopsis* genome seems to consist of pairs of closely related sequences.

Regulatory TFs are nuclear proteins that can regulate transcription in particular tissues and in response to cellular signals. The general view of regulatory TFs is that they bind to short DNA sequences (consensus motifs) at specific sites, typically in the promoter region of the relevant gene, and also to enhancers, which may be situated upstream or downstream from the transcriptional unit, or even within the unit itself. The transcription-affecting DNA motifs at the binding site are called regulatory elements (Latchman, 1998); it should be noted that the absence of a consensus motif does not preclude TF binding. Various mechanisms for TF activation in the absence of a proximal consensus site have been suggested: binding at a distal site that contains a consensus motif and then activating the site in question through protein–protein interactions; ‘piggyback’ binding that is mediated by protein–protein interactions with a second factor and that does not involve the DNA-binding domain of the first factor; and assisted binding to a site that is similar to the consensus site, which is enhanced by protein–protein interactions with another site specific DNA-binding factor or with a specifically modified histone. Factors that can be recruited to the genome in multiple ways are able to participate in multiple signaling pathways. The binding of a single regulatory TF can be essential, but not sufficient, for regulation of a specific gene. Overall, gene regulation in eukaryotes seems to
be heavily dependent on the cooperative and communal action of regulatory TFs (Farnham, 2009).

1.2.2 Alternative splicing of primary transcripts

Primary transcripts (pre-mRNA) generated by RNAP II can be spliced in various ways and therefore generate multiple templates (mRNA) from a single gene. In the nucleus the pre-mRNA are capped, spliced and polyadenylated before being exported to the cytosol for subsequent translation (Reddy, 2007). In plants, 20% of the genes undergo alternative splicing; the functionality of most of these has not yet been characterized (Barbazuk et al., 2008). The role of alternative splicing as a developmental and/or tissue-specific regulator is a topic of current interest (Srivastava et al., 2009).

1.2.3 Post translational regulation

Transcripts exiting the nucleus at the nuclear pores are translated, relocated or stored before ultimately being degraded (Bailey-Serres et al., 2009). Post-translational modifications (PTMs) of proteins are of great importance in transcription and gene regulation. The covalent binding of various molecules to the amino acids of proteins makes the proteome two to three orders of magnitude more complex than would be expected solely on the basis of the number of genes in the genome and the various alternative splicings they can undergo (Walsh et al., 2005). Around 300 different PTMs (Witze et al., 2007) have been identified; they affect the target protein's localization, stability, and interactions, and thus, its activity. In general, only a subset of the translated proteins are modified; this complicates the search for regulatory proteins and complexes (Jensen, 2006). Furthermore, some PTMs are reversible, rapid, and highly dynamic. For instance, phosphorylation is a reversible PTM that is commonly involved in signal transduction. Kinases and phosphatases regulate the phosphorylation of their targets with temporal resolutions measured in minutes (Olsen et al., 2006). Transcription of the genomic chromatin requires a cascade of PTMs of both the histones and the transcriptional machinery (Weake & Workman, 2008; Zhou et al., 2008; Pfluger & Wagner, 2007).
1.3 Genes in the pathway of wood formation

Wood formation is subject to highly regulated genetic control, primarily at the transcriptional level. Transcription factors (TFs) are some of the first active genes in the pathways leading to wood formation. The following section provides an overview of the current models of the regulation of wood formation, from the cellular to the tissue level.

1.3.1 Transcription factors and hormones

The first set of TFs that control early vascular development are regulated by hormones and miRNA (Demura & Fukuda, 2007) (Figure 3). In this model vascular formation is initiated by the binding of the TF ARF5 to the DNA following the recruitment of the inhibitory protein IAA12 by the auxin-binding protein TIR-1. This causes the degradation of IAA12 via the ubiquitin proteasome pathway. Such regulation via proteolysis is rather slow, with a lag time of 10-15 min; other hormone (auxin) responses are typically detected within a few minutes (Demura & Fukuda, 2007). Downstream of ARF5, there are two classes of TFs that regulate tissue patterning and polarity. The first class comprises four genes encoding class III homeodomainleucine zipper (HD-ZIPIII) transcription factors: IFL1, ATHB8, ATHB9 and ATHB14. The second comprises three members of the GARP-type transcription factor family: KAN1, KAN2 and KAN3. The HD-ZIPIII and KAN families have a complex pattern of overlapping and antagonistic functions involving control of the stability of the HD-ZIPIII transcripts by miRNA. Furthermore, it is possible that brassinosteroids (BRs) may be involved in up-regulating the HD-ZIPIII genes to promote xylem formation.
It was recently shown that a group of NAC domain TFs (PtrWNDs) may be key regulators of the cellulose, xylan and lignin pathways in poplar. The PtrWNDs are most likely orthologs of the arabidopsis AtSND1, a master transcriptional switch which activates a network of other TFs (Figure 4) involved in the control of the pathways that form the three main components of wood (Zhong et al., 2009).

Figure 3. Control of the early stages of tissue formation by transcriptional regulators and hormones. Arrows indicate activation; T-bars indicate inhibition. Adapted from Demura and Fukuda (2007).

Figure 4. Transcriptional regulation of secondary wall formation in arabidopsis. Arrows indicate activation. Adapted from Zhong et al. (2008).
The TFs downstream of AtSND1 are required for normal secondary wall biosynthesis in Arabidopsis. In situ hybridization localizes expression of all PtrWND genes to developing vessels and fibers. Additionally, PtrWND transcripts are observed in ray parenchyma cells, and a subset of the PtrWNDs are expressed strongly in phloem fibers. Overexpression of PtrWND2B and PtrWND6B (by the CaMV 35S promoter) in Arabidopsis induces the downstream TFs associated with secondary wall formation as well as the biosynthetic genes for cellulose (AtCesA4, AtCesA7 and AtCesA8), xylan (AtIRX8, AtIRX9 and AtFRA8) and lignin (At4CL1 and AtCCaAOMT1). Moreover, ectopic deposition of cellulose, xylan and lignin is observed in PtrWND2B and PtrWND6B overexpressors. However, only about 20% of these transgenic plants exhibit the ‘curly leaves’ phenotype. In Arabidopsis 35S-AtSND1, around 50% of the overexpressors have a characteristic visual phenotype in which the rosettes are small and the leaves are stunted and have severely upward-curling blades (Zhong et al., 2006). It is thought that the ‘curly leaves’ phenotype is only observed in a few of the transgenic plants because of differences in the cells’ competence to form secondary walls (Zhong et al., 2009).

The R2R3-MYB family is a large group of TFs that are heavily involved in the transcriptional activation of secondary wall formation. There are 192 R2R3-MYB family members in poplar (P. trichocarpa). An analysis of transcript abundance in poplar indicated that 23 out of 180 R2R3-MYB encoding genes are involved in xylem differentiation. It has been suggested that R2R3-MYB proteins may function as regulators of processes that are limited to discrete cells, organs, or conditions (Wilkins et al., 2009). Paper II describes a study investigating the role in wood formation of an R2R3-MYB TF which is upregulated in the latter stages of xylem formation (Karpinska et al., 2004). According to the previously-suggested nomenclature (Wilkins et al., 2009), this gene should be denoted PttMYB090 (in the article, it is referred to as PttMYB21a). Phylogenetic analyses suggest that this gene is one of two MYBs in poplar (PtrMYB090 and PtrMYB161), both of which are orthologs of AtMYB52 from Arabidopsis. PttMYB090 is rapidly (0.5–3h) induced following bending (tension) of the stem, suggesting that it may be involved in the early stages of tension wood regulation. Furthermore, antisense constructs of PttMYB090 increases the expression of CCoAOMT (caffeoyl-CoA O-methyltransferase), an enzyme involved in lignin biosynthesis (Karpinska et al., 2004). It was recently shown that expression of AtMyb52 activates all three of the major secondary wall biosynthetic pathways. However, this
alone was not sufficient to induce ectopic deposition of secondary walls (Zhong et al., 2008).

Prior to the identification of Mediator in Arabidopsis as described in paper I, the mediator subunit AtMed33b (AtMed5 using the updated nomenclature (Bourbon, 2008); accession number At2g48110) was identified during a screen for altered phenylpropanoid compounds in EMS-mutagenized plants. By means of a combination of mapping and sequencing, abnormally low levels of phenylpropanoid species were linked to amino acid substitutions at conserved positions in At2g48110. Plants bearing the mutant allele, named *ref4*, have reduced levels of all of the major phenylpropanoids, including lignin, flavonoids, and sinapate esters. The unusual semidominant behaviour associated with the mutant phenotype is characteristic of a mutation in a TF, transporter, or a component of a signalling cascade. It has been shown that dominant mutant traits can arise from point mutations that increase or stabilize mRNA transcripts or the proteins they encode, preventing normal turnover. For example, the atr1D mutant exhibits upregulated transcription of tryptophan pathway genes due to stabilization of the ATR1 MYB transcription factor mRNA. Moreover, the *ref4* mutant could be rescued by a second point mutation exchanging a proline for a leucine located in a WW protein-interaction domain (DWPSPA). The WW domain is conserved in all At2g48110 plant homologs and is a submotif within a proline-directed serine kinase phosphorylation site (DWPSPAA) (Stout et al., 2008).

1.3.2 Wall building enzymes and supporting scaffolds

Carbohydrate-active enzymes (CAZymes) are responsible for the synthesis, modification, and degradation of the carbohydrate polymers that comprise the cell wall, both in terms of the bonds that comprise the polymers' backbones and those at branch points. These enzymes are also responsible for the glycosylation of proteins, lipids and nucleic acids. In poplar, CAZyme transcripts are most abundant in woody tissues (Geisler-Lee et al., 2006). CAZymes are classified into different groups on the basis of their amino acid sequences. This is facilitated by their modular composition, featuring one or more catalytic domains and a variable number of independent substrate binding sites. Enzymes are assigned to a particular class on the basis of conserved active-site residues, molecular mechanisms, and protein folds (Davies & Henrissat, 2002). There are four classes of CAZymes; glycosyl transferases (GTs), glycoside hydrolases (GHs), polysaccharide lyases (PLs) and carbohydrate esterases (CEs). The CAZyme system provides a framework that is useful in the study of wood formation.
in poplar, for which gene annotation is limited and is largely based on functional characterizations of analogous genes from Arabidopsis.

Xylem-specific CAZymes in poplar have been identified using microarray analysis of narrow tissue sections taken from developing xylem. In total 25 GTs and 9 GHs were found to have secondary wall-specific expression patterns, but no PL or CE genes were significantly expressed (Aspeborg et al., 2005).

The key cellulose-synthesizing enzyme complexes are located in the plasma membrane and contain at least three types of cellulose synthase (CesA) proteins (Somerville, 2006) belonging to the GT2 family. PttCesA4, PttCesA8-A, PttCesA8-B and PttCesA7-x (classification according to Kumar et al. (2009)) are particularly highly expressed during xylogenesis (Aspeborg et al., 2005) and are therefore plausible candidate members of these complexes. The corresponding CesA homologs in Arabidopsis are associated with secondary cell wall formation (Persson et al., 2005). The transcript of the cellulose synthase-like gene, PttGT2A is very abundant at the onset of secondary cell wall formation, indicating that the protein it encodes may be a mannan or xylan synthase.

Several genes in the GT8 family of enzymes that catalyze the formation of \( \alpha \)-glycosidic linkages are expressed at high levels during secondary wall formation. PttGT8A, PttGT8B, and PttGT8C may be self-glucosylating proteins that serve as primers for polysaccharide synthesis. PttGT8D, PttGT8E, PttGT8F, and PttGT8G may be specifically involved in pectin biosynthesis in the contact ray cells. Two members of the GT14 family, PttGT14A and PttGT14B are expressed in xylem and have plant-specific features; they await functional characterization. In animals, GT14 enzymes catalyze the transfer of \( \beta(1-6) \)-linked N-acetylglucosaminyl and \( \beta \)-linked xylosyl residues to proteins. PttGT31A and PttGT31B from the GT31 family are also expressed, but no plant GT31 has been functionally characterized (in animals, these enzymes form \( \beta \)-glycosidic linkages between \( \alpha \)-linked nucleotide sugars). Two genes from the GT43 family (in animals, enzymes in this family are \( \beta \)-glucuronyltransferases), PttGT43A and PttGT43B, may be involved in fiber development. Four genes from the GT47 family are highly expressed; the first two, PttGT47A and PttGT47D, may be involved in the transfer of glucuronyl side chains to 4-O-methylglucuronoxylan in the secondary wall. Alternatively, the enzymes may be involved in the synthesis of rhamnogalacturonan II in ray cells. The second two, PttGT47B and PttGT47C, have not been characterized. One gene from the GT61 family, PttGT61A, is highly expressed during xylogenesis. The enzymes in subfamily A catalyze the transfer of Xyl from
UDP-Xyl to the core β-linked Man of N-linked oligosaccharides in glycoproteins.

The proteins encoded by genes from the GH9 family are typically endoglucanases which act on β(1-4)-glucan polymers (cellulose). The expression of PttCel9A is up-regulated during secondary cell wall formation. PttCel9A is probably a homolog of KOR1 in arabidopsis, which has been proposed to release cellulose chains from a putative sitosterol-glucoside primer or to be involved in the assembly or editing of cellulose microfibrils. PttXyn10 (a member of the GH10 family) is expressed in the xylem and may be involved in xylan hydrolysis or remodeling during xylogenesis. PttXET16L is a highly-expressed member of the GH16 family; it may have a role in the endohydrolytic degradation of xyloglucan. PttGH17 (a member of the GH17 family, which code for enzymes that degrade β(1-3)- or β(1-3)(1-4)-glucans) may have a role in callose degradation. PttGH19A from the GH19 family is expressed in a xylem-specific fashion, but its function is unknown. The GH28 family contains enzymes associated with pectin degradation and cell wall modification. Of these enzymes, PttGH28A is highly expressed and may be involved in the modification of the primary wall and the region between it and the developing secondary walls. The members of the GH35 family that have been characterized to date are β-galactosidases that act on various substrates, including arabinogalactans, galactolipids, and pectin. PttBGal35A and PttBGal35B are active during xylogenesis and may play a role in pit membrane adaptation and the hydrolysis of primary walls in the perforations, by degrading the galactan side chains of rhamnogalacturanan I or xyloglucan. Alternatively, their function may have to do with modifying arabinogalactan proteins. Members of the GH51 family have been shown to have α-L-arabinofuranosidase activity, releasing Ara from substrates such as arabinan, arabinoxylan, gum arabic, and arabinogalactan. PttGH51 is expressed during xylem formation and might act on homogalacturonan and the side chains of rhamnogalacturonan I.

Although the pathways involved in the biosynthesis of carbohydrate polymers in poplar have not yet been fully characterised, those involved in the biosynthesis of lignin are known in much greater detail. Lignin is a complex aromatic heteropolymer derived mainly from three monomers with differing degrees of methoxylation. These monolignols generate p-hydroxyphenyl (H), guaiacyl (G), and syringyl (S) phenylpropanoid units when incorporated into the lignin polymer. Angiosperm (poplar) lignin consists mainly of G and S units, with traces of H units. The monomers are synthesized from phenylalanine (derived from the shikimate biosynthesis
pathway) via phenylpropanoid and monolignol-specific pathways (Boerjan et al., 2003). Recently, 95 phenylpropanoid gene models extracted from poplar (P. trichocarpa) were investigated. 18 genes that were considered likely to encode monolignol biosynthesis enzymes were found to be highly transcribed in xylem undergoing differentiation. The current model of monolignol biosynthesis involves 10 enzyme families (Shi et al., 2010), as shown in Figure 5.
Figure 5. The biosynthetic pathway leading to monolignol (Shi et al., 2010). PAL (phenylalanine ammonia-lyase), C4H (cinnamate-4-hydroxylase), 4CL (4-coumarate-CoA ligase), HCT (p-hydroxycinnamoyl-CoA:quinate shikimate p-hydroxycinnamoyltransferase), C3H (4-coumarate 3-hydroxylase), CCoAOMT (caffeoyl-CoA O-methyltransferase), CCR (cinnamoyl-CoA reductase), CAld5H (coniferyl aldehyde 5-hydroxylase), COMT (caffeic
Finally, several cytoskeleton genes are essential for the deposition and remodeling of the secondary cell wall. It is well established that microtubules have an important role in wall formation. The large family of tubulin genes in poplar contains some distinct isoforms that are expressed particularly strongly in tissues undergoing secondary cell wall thickening (Oakley et al., 2007). The role of actin is less well understood. One model suggests that actin transports organelles containing cellulose synthase complex (CSC) around the cell, pausing at sites marked by transverse actin. The CSCs are incorporated into the plasma membrane at these sites, and are maintained beneath sites where wall synthesis is occurring by microtubules (Wightman & Turner, 2008).

1.4 Functional genomics

In the post-genomics era, functional genomics studies have largely focused on applying profiling techniques for the parallel monitoring of various aspects of cellular activity, such as transcription, protein synthesis, and the production of metabolites (Carrari et al., 2006; Kolbe et al., 2006; Rischer et al., 2006; Hirai et al., 2005; Tohge et al., 2005; Clish et al., 2004; Hirai et al., 2004; Kleno et al., 2004; Oresic et al., 2004; Gygi et al., 1999). This approach has become possible mainly due to the increasing availability of the instrumentation required for high-throughput characterization of biological samples, such as the microarray technology used in transcript profiling (Schena et al., 1995) or the use of high performance chromatography in conjunction with mass spectrometry for peptide or metabolite profiling (de Hoffmann & Stroobant, 2001). The objective of such studies is to examine organisms as integrated systems of genetic, protein, metabolic, pathways and cellular events in order to achieve a higher level of understanding of the interplay between molecular and cellular components.
2 Objectives

The objective of the work described in this thesis was to understand the process of wood (cell wall) formation in terms of proteomics. Wood formation is a complex process involving a large set of genes. In an attempt to obtain a holistic view of the pathway, we have studied a range of processes, from the transcription of genes to the synthesis of the wall itself. Recent developments and improvements in nanoflow liquid chromatography and electrospray ionization mass spectrometry have made it possible to study proteins (via peptides) in unprecedented detail. It has become possible to validate hypothetical model pathways on the basis of expression analysis (transcripts) and to both identify individual gene products and quantify their levels of expression. Paper I describes the determination of the composition of the co-regulatory transcription complex called the “Mediator”. The Mediator is thought to act as a molecular switchboard for upstream transcriptional activators of various different pathways. The correlation between transcription and protein expression (including a subset of metabolites) is discussed in paper II. Finally, paper III describes the isolation and characterization of plasma membranes from various tissues of poplar, which contain the major components involved in cell wall synthesis. A detailed understanding of the molecular pathway of wood formation would make it possible to manipulate the composition of the wood and to fine tune the regulation of the synthetic flow for sustainable growth. This knowledge would be of use in academic research and in the industrial production of pulp, paper, biofuel, and other renewable materials from forestry.
3 Material and methods

3.1 Liquid chromatography-mass spectrometry

The first techniques for the analysis of proteins and peptides using mass spectrometry (MS) were only suitable for use with volatile samples that could be vaporized without undergoing thermal destruction. In 1988, John Fenn and co-workers made a major breakthrough when they invented the concept of electrospray ionization. The electrospray ion source made it possible to perform MS analysis on liquid samples, especially those which had been purified in advance by liquid chromatography (LC) (Whitelegge, 2008). In order to analyze complex protein extracts and their digests, different separation methods are required. It is mainly the huge dynamic range in protein abundance that makes sample processing necessary. To analyze proteins of low abundance, it is advisable to start by extracting only differentiated tissues, individual cell types, or specific subcellular compartments in which the target protein is relatively abundant. Alternatively, where applicable, affinity purification of individual proteins and their associated complexes often gives superior results. When purifying individual proteins, extracts are commonly separated on the basis of protein size by gel electrophoresis after complete denaturation, and on the basis of the proteins' isoelectric points using isoelectric focusing (IEF). Peptides from digested proteins are separated by liquid chromatography and analyzed using mass spectrometry.

3.2 Plant material and Protein extraction

Two plant model systems and several protein extraction methods were used in the work described in this thesis. First, arabidopsis was grown in cell
suspension cultures (paper I) and extracted in order to isolate a protein complex by means of ion exchange chromatography, immunoprecipitation and gel electrophoresis (Figure 7). Second, we studied hybrid aspen (*Populus tremula × Populus tremuloides*) clones grown in greenhouses (paper II) and in the open field (paper III). The studies described in paper II used wild-type (WT) T89 clones and two different transgenic lines bearing antisense constructs of PttMYB090 (PttMYB21a). The antisense plants were named G3 and G5, corresponding to the “21III” and “21V” lines described by Karpinska *et al.* (2004). The experiment was designed so as to enable the separate analysis of developmental and genotype effects (Figure 6); Design of Experiments (DoE) was also used to reduce the sample size (see the Bioinformatics section). Tissue containing differentiated xylem cells was ground in liquid nitrogen, and samples of this powder were extracted to isolate three specific types of biomolecules: proteins (from 20 mg of powder), total RNA (from 50 mg) and metabolites (from 10 mg). Highly water-soluble proteins were extracted using a variation on the method of Giavalisco *et al.* (2003), adapted for use with LC-MS rather than a gel system (Figure 8).

![Figure 6. Overview of the study design. The different sample categories are shown, including three genotypes (G3, G3 and WT) and three internodes (A-C) on a 3 × 3 grid. The transcript, metabolite, and protein contents of samples from all 9 categories were measured. The smaller white circles explain the contents of the larger filled circles.](image)
In the work described in paper III, the plasma membranes of fresh leaves and of cambium/phloem and xylem tissues were isolated by a combination of aqueous polymer two-phase partitioning and ultra centrifugation, after which their proteins were separated by gel electrophoresis (Figure 9).
3.2.1 Isolation of protein complex from cell culture

Figure 7. Overview of the procedure used when isolating protein complexes from cell cultures.
3.2.2 Isolation of water-soluble proteins from xylem tissue

Figure 8. Overview of the procedure used when extracting water-soluble proteins from differentiated xylem tissue.
3.2.3 Isolation of plasma membrane proteins

Figure 9. Overview of the procedure used when extracting plasma membranes from differentiated xylem tissue.
3.3 Protein digestion

Protein sequencing by MS/MS relies on the prior digestion of the proteins to peptides. We have performed digestion on proteins separated using gel systems and on some separated using non-gel based systems. In order to maximise the yields from gel systems, high gel surface areas are desirable, as this improves both rehydration and the recovery of peptides. However, there is a lower limit on the size of the gel pieces that can be used (and thus an upper limit on the possible gel surface area) because gel pieces smaller than 0.5 mm\(^3\) clog the pipet tips and cause losses of gel material (Havlíš et al., 2003). The duration of the incubation time with the enzyme is also critical: the penetration of trypsin into the gel matrix is controlled by diffusion, which is much slower than the swelling of the dehydrated gel. Consequently, proteases that are larger than trypsin (e.g., thermolysin) are much less efficient at in-gel digestion than in-solution digestion (Havlíš et al., 2003). The handling of proteins separated by SDS-polyacrylamide gel electrophoresis requires laborious techniques that can easily introduce contamination. Potential contaminants include dust (keratin) that adheres to plastic tubes by static electricity, detergents from sample buffers, and residual reagents used in sample treatment. Reduction/alkylation is a general method for sample treatment that usually improves separation and identification but can also reduce sensitivity (Borchers et al., 2000). We omitted the reduction/alkylation step in the work described in paper III; the omission increased the sensitivity of the analysis and also considerably reduced the sample treatment time.

3.4 Nanoflow liquid chromatography

The complex peptide mixtures generated in these studies were separated using liquid chromatography (LC). The stationary phase of the column was functionalised with linear C\(_{18}\) (n-octadecyl) chains, which give the column a high capacity and make it very hydrophobic. Peptides are retained on the column primarily due to the combined hydrophobicity of their individual amino acids (Krokhin et al., 2004). Separation is carried out using a nanoscale flow of the liquid phase (water, acetonitrile and 0.1% formic acid) to achieve efficient electrospray ionization and thereby increase sensitivity. To facilitate the use of sample injection volumes in the 1-10 µl range, a pre-column was used to concentrate and wash the samples (at flow rates measured in µl/min). Thus, the sample is injected and washed on the
trapping column (Figure 10) with the HTM (heat and trapping module) valve open, allowing the excess eluent to flow into the waste container. The HTM valve is then closed, redirecting the flow to the analytical column to separate the sample at a flow rate of 200-600 nl/min.

Figure 10. Setup of the nanoscale UPLC system. The injection valve allows for loading of the sample into the sample loop (10 µl) and, in conjunction with the HTM (heat and trapping module) valve, delivers it to the analytical column. The binary pumps provide a steady flow of the liquid phase (water, acetonitrile and 0.1% formic acid), mixed according to a user-specified gradient. The analytical column is housed in a heated sleeve for thermal control and enhanced reproducibility.

3.5 Electrospray ionization

In the electrospray ionization (ESI) of peptides, an acidic, aqueous peptide solution is sprayed through a small-diameter needle. A high positive voltage is applied to this needle to produce a Taylor cone from which droplets of the solution are sputtered. The protons from the acid give the droplets a positive charge, causing them to move away from the needle towards the negatively charged instrument. During the course of this movement, evaporation reduces the size of the droplets until the number and proximity of the positive charges split the droplet into a population of smaller, charged droplets. The evaporation process is often accelerated using a stream of heated nitrogen gas. The droplet-splitting is repeated until the small size and high charge of the droplets causes desorption of the protonated peptides into the gas-phase (Kinter & Sherman, 2000). The acidic conditions used in
electrospray ionization result in protonation of the peptides' basic functional groups, such as the N-terminal amine and the basic side chains of lysine (K), arginine (R) and histidine (H). As a result, multiply-charged peptide ions dominate the mass spectrum.

3.6 Time-of-flight mass spectrometry

A hybrid quadrupole time-of-flight (Q-ToF) mass spectrometer was used in all of the MS experiments discussed in this thesis. This instrument can also be classified as a tandem-in-space instrument, in contrast to instruments with only one mass analyzer, which perform tandem-in-time analysis. The operational principals of time-of-flight mass analysers are elegantly simple. Ions are accelerated by a fixed electric field to a kinetic energy (and thus a velocity) that is inversely proportional to their m/z (equation), and then travel at this velocity through a field-free area.

\[ v = \left( \frac{2 \cdot V \cdot z}{m} \right)^{1/2} \]

\( m = \text{mass in kg, } v = \text{velocity in m/s, } V = \text{accelerating voltage, } z = \text{charge} \)

Therefore, ions with low m/z travel more rapidly than ions with high m/z. The time required for ions to travel through the field-free region can be accurately measured and used to calculate their m/z. The acceleration voltage and the length of the flight tube are two critical factors that affect the resolution and sensitivity of the measurement (Kinter & Sherman, 2000). To avoid increasing the length of the flight tube, a reflector is used. Current instruments have two reflectors (not shown in Figure 12) to accommodate “W-mode” ion travel, which improves resolution. In our study, we sought to maximise the sensitivity of the instrument, and therefore operated it in “V-mode”. Despite this, isotopic resolution was still achieved.
Figure 11. Schematic diagram of a hybrid Q-Tof mass spectrometer (V-mode). In a peptide fragmentation tandem mass spectrometry (MS/MS) experiment, the first stage involves selection of the desired precursor (intact) peptide ion, which is accomplished using the quadrupole mass filter (MS1). The second stage involves fragmentation in the collision cell, a hexapole lens system in which ions are accelerated in argon gas and all fragments are transmitted further without regard to their m/z. The third stage employs an orthogonal time-of-flight mass analyzer (MS2) which performs the mass analysis. For molecular weight measurement (MS), the first quadrupole is used in rf-only mode and transmits all peptide ions; no fragmentation is induced in the collision cell, and mass analysis is performed using the time-of-flight mass analyzer.

3.7 Peptide fragmentation

In mass spectrometric sequencing, the information that describes the amino acid sequence of the peptide is contained in the product ion spectrum. The product ion spectrum is acquired from a tandem mass spectrometry experiment by using low-energy collision-induced dissociation (CID). By using the first mass filter to select intact peptides, individual peptides within a narrow mass range can be accelerated in a collision cell. In the collision cell, which is typically filled with argon gas, intermolecular collisions convert kinetic energy (10eV to 50eV) to vibrational energy in the peptide ions. The ions’ internal vibrational energy causes fragmentation according to
the mobile proton hypothesis. In the gas-phase, protons bind strongly to basic groups and remain bound even after collisions. In contrast, protons bound to less basic functional groups can move around 'within' the peptide to any one of its amide bonds by means of internal solvation. The migration of the mobile protons produces sub-populations of peptide ions and directs subsequent charge-site-directed fragmentation. Fragmentations occur primarily at the protonated amide bonds, producing the characteristic y and b-type ions, from the C- and N-termini respectively (Figure 12). Detection of the fragment ions in a second high resolution mass analyzer allows the fragments' masses to be measured with sufficient accuracy to determine the identities of their constituent amino acids. However, current instruments cannot distinguish between the isobaric amino acids isoleucine and leucine. Peptides produced by tryptic digestion are ideally suited for ESI and CID fragmentation. The charged C-terminal lysine and arginine residues of the fragments produced by tryptic cleavage give rise to predictable fragmentation spectra that can readily be related to specific amino acid sequences.

Figure 12. Peptide ion fragmentation nomenclature. Low-energy collision-induced dissociation primarily causes fragmentation at the peptide bonds (peptide or amide linkage) generating b- and y-ions. Internal fragmentations generate combinations of a-/x-ions or c-/z-ions as well as immonium ions characteristic of each amino acid.
3.8 Bioinformatics

3.8.1 Preprocessing of raw data
Continuous mass spectral data from the peptide ion (MS) and fragment ions (MS/MS) require processing before they can be used in database searches. The typical workflow for processing spectral data involves background reduction, smoothing, centroiding, offset calibration, deisotoping and charge state deconvolution. The optimal processing settings for a given spectrum depend on the sample quality and the acquisition parameters used on the instrument. Different database search algorithms require the use of specific settings for optimal performance. Processing ultimately generates a basic list (peak list) of singly charged, monoisotopic masses for the intact peptide and its fragments.

3.8.2 Databases and search engines
The tools used for comparing mass spectrometry results to data in sequence databases are known as database-dependent algorithms because they are heavily dependent on the sequence database. Peak lists from processed raw data can be submitted to a large number of different commercial and publicly available database search engines (Matthiesen & Jensen, 2008). No matter what search engine is used, no results will be generated if the corresponding protein sequence is not present in the database. With common search parameters, exact amino acid matches are required. Good sequence coverage is available for Arabidopsis because of the existence of extensive EST collections (from different ecotypes) and genomic sequencing data. Furthermore, there is a well-organized gene annotation system for Arabidopsis, which allows for quick access to and use of genes with alternative splicing. Poplar is a much more difficult species to study, in part because the ecotype whose genome was sequenced is P. trichocarpa (Tuskan et al., 2006), but research is often performed on hybrids or on natural populations of aspen (P. tremula) which exhibit substantial genetic variation (Luquez et al., 2008). Furthermore, the whole-genome duplication event that occurred in poplar after its lineage diverged from that of Arabidopsis greatly increased the complexity of its proteome. There are approximately 45,000 putative protein-coding genes in poplar, including nearly 8,000 pairs of paralogous genes (Tuskan et al., 2006) whose protein products are too similar to be readily distinguished by MS/MS sequencing. The presence of paralogous genes and genetic variation makes the process of merging MS/MS results with data from a different source like an EST database unfeasible. Search engine programs like PLGS from Waters make it possible
to conduct a secondary database search to find modified peptides derived from proteins found in the preliminary first search. This facilitates the detection of mutations, genetic variations that affect codon usage and splicing, and post-translational protein modifications. Alternatively, one can use the de novo sequencing method, which does not rely on sequence databases. The process of algorithm-based de novo sequencing starts with the identification of significant ions, followed by examining the internal immonium ions to determine the peptide's amino acid composition, creation of b- and y-ion-evidence lists, determination of peptide sequences from the spectrum, generation of complete sequences (possible candidate sequences), and finally, scoring and ranking of the various calculated sequences (Reinders et al., 2004). Manual or automated de novo sequencing is essential when evaluating peptides that have an ambiguous fragmentation pattern. While search engines will always identify the structure in the database that most closely matches the experimental data, the use of incomplete databases means that this method has the potential to generate false positive matches. Methods for estimating the false positive rate are therefore required.

Extracting peptide markers from LC-MS experiments and connecting these markers to individual proteins necessitates the use of rather elaborate data preparation techniques. The general workflow for the process is shown in Figure 13.
Careful experimental planning is required if one is to correctly interpret the results obtained from the analysed samples and their associated biological annotations. Where possible, we employed design of experiments (DoE) (Box et al., 1978) in order to avoid drawing inaccurate conclusions from our experimental results. In the study described in paper II, a large set of samples (90 samples, with 10 biological replicates) was subjected preliminary metabolic profiling using GC-MS. The resulting metabolite profiles were subsequently explored using Principal Component Analysis (PCA) (Jolliffe,
in order to identify similarities and differences between the biological replicates, using the first principal component (along which the greatest variance in the data lies). The four most diverse biological samples for each genotype and internode were selected as candidates for further profiling in terms of their transcript and protein contents. The essential principle of this selection strategy was to ensure that the natural variability in the biological samples was not underestimated.

### 3.8.3 Bidirectional multivariate regression

The study described in paper II employed a bidirectional multivariate regression method (Trygg & Wold, 2003; Trygg, 2002) called O2PLS that identifies joint covariation between two data sets as well as systematic variation that is unique to each data set (Figure 14). Both the jointly covarying and unique sources of variation are composed of smaller units which are referred to as latent variables that describe independent effects in the data.

![Figure 14. Overview of the components of the O2PLS model. The variation in each data set (matrix) is separated into predictive (joint) variation, data set specific variation and nonsystematic (residual) variation.](image)

To handle multiple datasets, further processing is required. In paper II we integrate data from three different analytical platforms (transcript, protein and metabolite). Stepwise processing was employed to identify joint covariance, extract platform specific variation and finally, to analyse the matrices (Figure 15).
Figure 15. The O2PLS-based integration framework. The different steps used to identify joint and platform-specific variation for three data sets by means of O2PLS are shown. In Step 1, the joint covariance structures from the transcript and protein data sets are identified. In Step 2, the joint covariance structures from the joint transcript-protein variation in Step 1 and the metabolite data are utilized to identify the joint covariance between all data sets. In Step 3, the joint covariance structures are removed from each data set and specific systematic variation is extracted.

3.8.4 Visualizing bioinformatics data

We have used various approaches to obtain overviews of complex datasets and to capture the underlying biological relationships. To visualize the relationship between the genes of interest in paper II, we employed a multilayer network algorithm by forming the union of multiple minimal spanning trees (MST), each of which was constructed by random resamplings of a large data set of 1024 microarrays (Grönlund et al., 2008). Such network visualization techniques can reveal interesting pathways by
application of the “guilt by association” principle. We integrated the network clusters with the correlation loading scores obtained from the O2PLS analysis to identify affected pathways.
4 Results and discussion

4.1 Composition of the plant transcriptional regulator

Mediator was originally identified as a central transcriptional coregulator in yeast (Kim et al., 1994; Flanagan et al., 1991; Kelleher et al., 1990), but it is now clear that it also plays the same important role in higher eukaryotes (Bourbon et al., 2004). A Mediator-like complex would be expected to play a major role in the regulation of wood formation at the transcriptional level during the appropriate stages in development. It was therefore puzzling that prior to our studies, no Mediator-like complex had been reported in plants. The nature of the Mediator complex we isolated from arabidopsis (paper I) suggests an explanation for this mystery: the subunits of the plant Mediator show very little sequence homology with the corresponding subunits from yeast and metazoans (Table 2).

Table 2. Homology between from arabidopsis and S. cerevisiae Mediator subunit 6, 7, 10, 14, 15 and 21 summarized

<table>
<thead>
<tr>
<th>Gene</th>
<th>identity</th>
<th>similarity</th>
<th>length (aa)</th>
</tr>
</thead>
<tbody>
<tr>
<td>AtMed6</td>
<td>24%</td>
<td>41%</td>
<td>191</td>
</tr>
<tr>
<td>AtMed7</td>
<td>33%</td>
<td>48%</td>
<td>188</td>
</tr>
<tr>
<td>AtMed10</td>
<td>26%</td>
<td>50%</td>
<td>139</td>
</tr>
<tr>
<td>AtMed14</td>
<td>21%</td>
<td>41%</td>
<td>171</td>
</tr>
<tr>
<td>AtMed15</td>
<td>26%</td>
<td>38%</td>
<td>132</td>
</tr>
<tr>
<td>AtMed21</td>
<td>24%</td>
<td>41%</td>
<td>139</td>
</tr>
</tbody>
</table>

This low homology is not so surprising considering that the arabidopsis genome encodes ~1 700 transcription factors, of which ~700 are specific to plants (Riechmann & Ratcliffe, 2000). Since these plant-specific
transcription factors will execute their function through Mediator, it was perhaps to be expected that the plant Mediator subunits would differ from those in yeast and humans.

In total, we identified 24 arabidopsis Mediator subunits by LC-ESI-MS/MS after purification by immunoprecipitation using antibodies raised against AtMed6, AtMed7, and ultimately, AtMed32 which would be named AtMed2 (Bourbon, 2008). We were unable to find arabidopsis homologs of the conserved Med1 subunit or the metazoan-specific Med26 subunit. The positions of gel spots (from 2D gels) and slices (from 1D gels) containing peptides that correspond to the identified Mediator subunits can be found in Figure 2 of paper I. Multiple spots corresponding to AtMed8 were observed. This is probably due to degradation, but may be indicative of post-translational modifications that affect its isoelectric focusing (Figure 16).

![Figure 16](image)

Figure 16. 2D gel electrophoresis of arabidopsis Mediator. Immunoprecipitated proteins were separated along the first dimension by isoelectric focusing followed by separation along the second dimension with SDS-PAGE (12%). Circles indicate spots where peptides belonging to AtMed8 were identified after in-gel tryptic digestion and LC-MS/MS analysis.

It is apparent that arabidopsis Mediator contains homologs of nearly all of the subunits from the head and middle domains in yeast, where, these subunits are situated in close proximity to RNAP II. However, it was not possible to identify homologs of many of the subunits that comprise the tail
domain in the yeast Mediator. After the publication of paper I, a comprehensive bioinformatics analysis of the evolutionary origin of the Mediator was undertaken; our recent results were examined along with the rest of the literature data (Bourbon, 2008). This detailed comparison of Mediator subunits from 70 eukaryotes with completed genome sequences (>90%) clarified the relationships between the divergent tail subunits. It was concluded that AtMed32 is a homolog of yMed2, AtMed33a/b is a homolog of yMed5a/b, and AtMed27 is a homolog of yMed3. Additionally, At5g63480 was found to be a homolog of yMed30; at the time of writing paper I, we had not been able to determine which Mediator domain it belonged to. The updated names of the Mediator subunits identified in our study are shown in Table 3. Of the four domains, the tail is the least conserved between different species. It is therefore likely that the plant-specific functions of the Arabidopsis Mediator are carried out by the tail subunits.

Table 3. Mediator subunits of Arabidopsis identified by LC-ESI-MS/MS

<table>
<thead>
<tr>
<th>Protein name</th>
<th>New def. Bourbon</th>
<th>Accession number</th>
<th>Mediator module</th>
</tr>
</thead>
<tbody>
<tr>
<td>Med32</td>
<td>Med2</td>
<td>At1g11760</td>
<td>Tail</td>
</tr>
<tr>
<td>Med27</td>
<td>Med3</td>
<td>At3g09180</td>
<td>Not identified</td>
</tr>
<tr>
<td>Med4</td>
<td>Med4</td>
<td>At5g02850</td>
<td>Middle</td>
</tr>
<tr>
<td>Med33a</td>
<td>Med5#1</td>
<td>At3g23590</td>
<td>Tail</td>
</tr>
<tr>
<td>Med33b</td>
<td>Med5#2</td>
<td>At2g18110</td>
<td>Tail</td>
</tr>
<tr>
<td>Med6</td>
<td>Med6</td>
<td>At3g21350</td>
<td>Head</td>
</tr>
<tr>
<td>Med7a</td>
<td>Med7#1</td>
<td>At5g03220</td>
<td>Middle</td>
</tr>
<tr>
<td>Med7b</td>
<td>Med7#2</td>
<td>At5g03500</td>
<td>Middle</td>
</tr>
<tr>
<td>Med8</td>
<td>Med8</td>
<td>At2g03070</td>
<td>Head</td>
</tr>
<tr>
<td>Med9</td>
<td>Med9</td>
<td>At1g55080</td>
<td>Middle</td>
</tr>
<tr>
<td>Med10a</td>
<td>Med10#1</td>
<td>At5g41910</td>
<td>Middle</td>
</tr>
<tr>
<td>Med10b</td>
<td>Med10#2</td>
<td>At1g26665</td>
<td>Middle</td>
</tr>
<tr>
<td>Med11</td>
<td>Med11</td>
<td>At3g01435</td>
<td>Head</td>
</tr>
<tr>
<td>Med14</td>
<td>Med14</td>
<td>At3g04740</td>
<td>Tail</td>
</tr>
<tr>
<td>Med15</td>
<td>Med15#1</td>
<td>At1g15780</td>
<td>Tail</td>
</tr>
<tr>
<td>Med16</td>
<td>Med16</td>
<td>At4g04920</td>
<td>Tail</td>
</tr>
<tr>
<td>Med17</td>
<td>Med17</td>
<td>At5g20170</td>
<td>Head</td>
</tr>
<tr>
<td>Med18</td>
<td>Med18</td>
<td>At2g22370</td>
<td>Head</td>
</tr>
<tr>
<td>Med19</td>
<td>Med19#1</td>
<td>At5g12230</td>
<td>Head</td>
</tr>
<tr>
<td>Med20</td>
<td>Med20#2</td>
<td>At2g28230</td>
<td>Head</td>
</tr>
<tr>
<td>Med21</td>
<td>Med21</td>
<td>At4g04780</td>
<td>Middle</td>
</tr>
<tr>
<td>Gene</td>
<td>Locus</td>
<td>Protein Name</td>
<td>Position</td>
</tr>
<tr>
<td>--------</td>
<td>---------</td>
<td>--------------</td>
<td>----------</td>
</tr>
<tr>
<td>Med22a</td>
<td>Med22#1</td>
<td>At1g16430</td>
<td>Head</td>
</tr>
<tr>
<td>Med22b</td>
<td>Med22#2</td>
<td>At1g07950</td>
<td>Head</td>
</tr>
<tr>
<td>Med23</td>
<td>Med23</td>
<td>At1g23230</td>
<td>Not identified</td>
</tr>
<tr>
<td>Med25</td>
<td>Med25</td>
<td>At1g25540</td>
<td>Not identified</td>
</tr>
<tr>
<td>Med28</td>
<td>Med28</td>
<td>At3g52860</td>
<td>Not identified</td>
</tr>
<tr>
<td>Med30</td>
<td>Med30</td>
<td>At5g63480</td>
<td>Not identified</td>
</tr>
<tr>
<td>Med31</td>
<td>Med31</td>
<td>At5g19910</td>
<td>Middle</td>
</tr>
</tbody>
</table>

The identification of At2g48110, which encodes AtMed5#2, (AtMed33b) as a subunit of Mediator is the first direct link between Mediator and the wood formation pathway. Point mutations at this locus have been shown to affect the phenylpropanoid pathway and lignification, (Stout et al., 2008) which strongly suggests that Mediator is an important target of this pathway. Regulation is most likely effected by interactions between Mediator and the various regulatory TFs that are involved in the transcriptional networks that coordinate wood formation (Zhong & Ye, 2007). A member of the MYB TF family has already been shown to interact with a Mediator tail subunit in yeast two-hybrid screens (submitted). Paper I focuses on the AtMed25 subunit (PHYTOCHROME AND FLOWERING TIME 1, PFT1) and its involvement in flowering. It has recently been shown that the induction of flowering triggers xylem expansion in the hypocotyl and secondary growth in the roots in arabidopsis (Sibout et al., 2008). The data support the hypothesis that flowering-related signals from the shoot control xylem expansion in the hypocotyl and roots.

The partitioning of the Mediator complex into core and peripheral elements is echoed in the structure of one of its subunits, AtMed25. We have shown that the N-terminal domain of AtMed25, which interacts with the Mediator core in human cells, is conserved in arabidopsis. However, human Med25 has a domain close to the C-terminus that interacts with the transcriptional activator VP16. This domain is conserved in metazoans but is not conserved in arabidopsis. By contrast, the corresponding C-terminal region is conserved between different plants (Figure 17). It seems that Mediator may have evolved to accommodate species-specific transcriptional regulators by incorporating subunits that possess one highly conserved domain that interacts with the other Mediator subunits together with a second, species-specific domain that interacts with the appropriate regulators of transcription.
Interestingly, we identified peptides that map to different paralogs of the AtMed7, AtMed10, AtMed19, AtMed20, and AtMed22 subunits in our purified arabidopsis Mediator. Gene duplications that generated paralogs have played important roles in evolution. Some excellent examples of subfunctionalization and neofunctionalization after gene duplication were identified in arabidopsis by analysing 280 phylogenetically identified paralogous pairs. The expression of these pairs was measured in six organs; 85% of them exhibited significant gene by organ interactions, indicating that they have undergone subfunctionalization and/or neofunctionalization (Duarte et al., 2006). Because we purified the arabidopsis Mediator from a cell suspension culture, we are unable to determine whether different paralogs are present in a single complex or even whether they are normally expressed in the same cell or cell type. It is possible that Mediator complexes containing different paralogs have different functions and that different paralogs are expressed in different tissues or in different developmental stages. Mediator has been reported to have tissue- and cell-specific functions. One way in which this could arise would be if the Mediator complexes of the different cell types or tissues contain different subunits. The corresponding paralogous genes in poplar may be involved in the regulation of tree specific traits like wood formation, long-term perennial growth, and seasonality.

Our studies on the isolation of Mediator also highlighted the importance of using an appropriate protein database in work of this kind. In this case, the NCBI database proved to have the greatest sequence coverage of At2g48110. Alternative gene models for the At2g48110 locus from the NCBI database were found to match the acetylated N-terminal peptide and a peptide having a slightly different C-terminus that was not present in the TAIR database. Moreover, we identified the full length mRNA arising from this locus; its size is consistent with a protein having a molecular weight of 16 kDa.
4.2 Correlation between transcripts, proteins and metabolites

Paper II describes a functional genomics strategy for the integrated analysis of transcript–(microarray), protein– and metabolite–data. The strategy relies on the use of designed experiments that are specifically tailored to the study at hand. However, the methodology is unsupervised in the sense that no prior knowledge of the sample is used in the modelling. We decided to demonstrate this strategy using poplar mutants that exhibit abnormal wood development. The mutants incorporate antisense constructs of PttMYB090 (PttMYB21a); this protein is involved in lignin biosynthesis (Karpinska et al., 2004) and is part of the regulatory TF network that controls wood formation (Zhong et al., 2009; Zhong et al., 2008). We collected xylem tissue from wild-type (WT) and two transgenic lines (G3 and G5) at three internode positions (denoted A–C). Our experimental design (Figure 6) allowed us to analyse the growth characteristics and genotype effects individually and to look at synergistic effects. Multiple O2PLS models were used to integrate and reduce the large datasets generated. Two main effects were revealed by analysis of the joint covariation in the datasets; the effects were visible in all three profiles (transcript, protein, and metabolite). The first joint effect is the internode gradient, which reflects the common developmental progress of the samples, regardless of their genotype (Figure 18, A). The second joint effect reflects differences between the G5 and G3 genotypes, and is not sensitive to the precise internode examined (Figure 18, B). We refer to the first phenomenon as the internode effect and to the second as the genotype effect.

Figure 18. The internode and genotype effects. (A) The internode gradient is seen along the second joint score vector. (B) The genotype effect is shown as a combination of the first (G5 versus G3 and WT) and third joint score vectors (G3 versus G5 and WT).
These effects reflect the experimental setup shown in Figure 6. This verifies the fitness of the integrative methodology, as information on the nature of the samples was not factored into the experimental design. The results indicate that a considerable part (40% on average) of the existing variation in the steady-state system can be linked to changes in transcript levels, which are in turn reflected in the quantities of specific proteins observed and in the levels of individual metabolites. Much of the variation in the data reflects variation within individual profiles (e.g. within transcript levels) rather than interaction effects, and seems to be related to various housekeeping-type events. For example, we observed transcript-specific variation that was associated with processes involving chromatin remodelling and the assembly of protein-DNA complexes (Fig 8, paper II). This may be due in part to the fact that while microarray technology permits near-exhaustive monitoring of the transcriptome, the techniques used for proteomics and metabolomics generally measure only the most abundant proteins and metabolites. Secondly, processes like chromatin remodelling are regulated by complex post-transcriptional modifications and/or post-translational events that are not readily captured by the latter methods without using advanced enrichment techniques. This limitation restricts the scope for the simultaneous analysis of different kinds of -omics datasets using a global profiling approach; further technological advances are required to facilitate more comprehensive analyses.

The difficulties associated with combining transcript and proteomics data were highlighted in our studies of monolignol biosynthesis. We were able to identify the key enzymes involved in the biosynthetic pathway (Figure 19) but were only able to relate a few of these to specific extracted peptide markers. Additionally, we sought to identify the genes corresponding to these proteins using the microarray. However, the inherent complexity of the poplar genome, with its large number of duplicated genes, made it difficult to perform comparisons between the datasets. Even if it were possible to uniquely relate a peptide to a specific gene model, it would not be possible to uniquely identify the corresponding spot on the array. Each gene model on the spotted array usually has multiple reporters (PU-numbers) and it is essential to treat these in a uniform manner. Extensive filtering of the dataset was necessary, which made it difficult to obtain deeper insights into the biology underpinning the data.
Despite the difficulties associated with using correlations between the transcript data and the proteomics data to identify unique gene models, the integrated analysis did reveal some interesting correlations. We were able to confirm previously-identified links between specific transcripts, proteins, and metabolites, and also discovered a number of potential new associations. The changes in the levels of photosynthesis-related transcripts, glucose metabolism proteins, and amino acids on going upwards from internode to internode (i.e. upwards along the developmental gradient) were as expected. Such changes are an unavoidable necessity of plant growth, and were accurately detected by all three profiling techniques. Specific effects due to the down-regulation of the PttMYB090 (PttMYB21a) gene, which is known to affect plant growth and lignin biosynthesis, were observed in the mutants (Karpinska et al., 2004). We found that the normal growth gradient and the differences between the G5 and G3 genotypes are independent of one another. This would imply that with some exceptions, the mutants share the essential growth characteristics of a normal plant, although their
growth is somewhat slowed. The levels of important factors in both cell growth (specifically, tubulin and protein translation elongation factors) and lignin biosynthesis (COMT2 and CCoAOMT2) are heavily affected in both the transcript and protein data sets for the stronger G5 mutant. Importantly, it was observed that the reduction in transcript levels due to the G5 genotype resulted in an increase in the levels of the proteins encoded by those transcripts. Similar phenomena have previously been observed in yeast (Gygi et al., 1999); they indicate that there are various active mechanisms of post-translational regulation and consequently, the levels of proteins will not be strongly correlated with those of the corresponding transcripts. Additionally, the changes in the levels of COMT2 and CCoAOMT1/2 in the G3 and G5 mutants correlated with changes in the levels of metabolites such as quinic acid that are involved in lignin biosynthesis.

4.3 Components of wood formation in the plasma membrane

In the work described in paper III, we identified more than 900 proteins in the plasma membranes isolated from leaves, xylem, and cambium/phloem obtained from young poplar trees (see Table 1, paper III and Supplemental Table 3, which is available online). One beneficial effect of using big trees in proteomics studies is that one can readily obtain rather large samples; we took advantage of this. A comprehensive protocol for the isolation and purification of plasma membranes was employed (Figure 9), involving the aqueous polymer two-phase partitioning technique previously described by Larsson et al. (1994). In the work described in paper I, we were able to successfully identify proteins digested after purification by 1D gel separation; 2D separation was not necessary. We therefore adopted the same approach when studying the plasma membrane proteins. 1D gel separation does not separate protein isoforms or other modified versions and provides higher protein to gel ratios than does 2D separation; this is favourable for gel digestion (Havlíš et al., 2003) and subsequent identification of the digested proteins. The protein bands were well focused, making it possible to use multiple injections, and individual bands was analysed over several mass ranges to increase the number of identified proteins. Because of the large number of proteins identified, we focused on proteins with transmembrane domains, as was done in previous studies. It should be noted that this restriction will exclude both true peripheral proteins that associate with the membrane by means of lipid anchors and also those that bind to the membrane as subunits of larger protein complexes. Up to 7% of the
identified proteins that lack a transmembrane domain are predicted to have lipid anchors.

The primary conclusion of this study was that more than 40% of the 956 proteins identified were found in the plasma membranes of all three tissues examined, and may thus be classified as “housekeeping” proteins. The most valuable results in terms of increasing our understanding of wood formation in poplar were obtained by comparing the differences in protein levels between the tissues. A graphical model of the plasma membrane and the complexes involved in cell wall synthesis was created, and cell wall related genes from arabidopsis were superimposed on this (Figure 20). Extra symbols were added to indicate proteins identified in a particular tissue. In several cases, we were able to identify unique peptides corresponding to specific poplar gene models, and thus to confirm their existence and location.

Figure 20. Proteins involved in wood formation. The figure shows a schematic model of the cellulose-synthesizing complex and other proteins associated with wood formation, such as
the enzymes involved in lignin biosynthesis. The numbers outside each symbol refer to the
gene model ID in Table I of paper III for integral proteins and supplemental Table 3 for
soluble proteins. Only proteins detected in the xylem plasma membranes are shown; they are
denoted by a red triangle. Proteins that are also found in the plasma membranes of leaves are
marked by a green triangle, while those that are found in cambium/phloem bear a blue
triangle. A star in a colored field indicates that the protein is highly ranked within that tissue.
Where available, the proteins’ names in poplar are shown below the symbol. Abbreviations
for the NAD-dependent epimerase dehydratase family are: UGE, UDP-D-glucose 4-
epimerase; GME, GDP-D-mannose 3,5-epimerase; RHM, UDP-L-rhamnose synthase;
UXS, UDP-D-apiose/xylose synthase. Abbreviations for the lignin enzymes are: CAD,
cinnamyl alcohol dehydrogenase; COMT, caffeic acid O-methyltransferase; CCoAOMT,
caffeoyl-CoA 3-O-methyltransferase; HCT, hydroxycinnamoyltransferase; PAL,
phenylalanine ammonia-lyase.

8% of the 213 integral membrane proteins examined are involved in cell
wall and carbohydrate metabolism. The other integral membrane proteins
are discussed in detail in paper III; transport proteins constitute the largest
class (41%), followed by receptors (14%), and proteins involved in
membrane trafficking (8%). “Others,” of which various stress-induced
proteins are the largest group, represent only 17%, and “unknowns” only
9%. Several of the ABC transporters show xylem-specific localization,
indicating that they may be involved in secondary cell wall formation. Space
constraints prohibited the inclusion of our phylogenetic analysis of the
receptors in paper III. This analysis is shown in Figure 21. Some xylem-
specific groups were identified from this important collection of signalling
proteins Of these proteins, only the LRR RLKs denoted by IDs 186 and
188 have (to the best of our knowledge) previously been associated with
wood formation; this association was suggested on the basis of coexpression
analyses (Persson et al., 2005).
Figure 21. Phylogenetic Analysis of Plasma Membrane Receptor-Like Kinases (RLKs) and their Tissue Localization. Both highly-ranked and subset RLKs detected by mass spectrometry are included (two truncated sequences, denoted by IDs 197, 198 were excluded). The numbers outside each symbol refer to the gene model ID in Table 1, paper III (which contains top-ranked integral proteins only, indicated by numbers in bold) and Supplemental Table 2 (which contains both top rank and subset integral proteins). According to the phylogenetic analysis, four subfamilies of RLKs are represented: CrRLK1L (Catharanthus roseus RLK1-Like), LRR (Leucine-Rich Repeat), PERK (Proline Extensin-like Receptor Kinase), and SD (S-locus glycoprotein-like Domain); the subfamilies are distinguished by their extracellular N-terminal ligand-binding domains (Shiu & Bleecker, 2001). Proteins found in leaves are denoted by a green triangle, those in xylem by a red triangle, and those in cambium/phloem by a blue triangle. A star in a colored field indicates that the protein is highly ranked within that tissue.

Proteins involved in cell wall and carbohydrate metabolism were most abundant in the xylem plasma membranes, in keeping with the role of the xylem in wood formation. Many peripheral proteins have essential roles in supporting the complex machinery that constructs the secondary cell wall.
We have identified proteins derived from a range of established genes and confirmed their expression and localization. Several novel candidates were also identified that have roles in less well-understood cellular processes that are nevertheless important in secondary cell wall formation.
5 Conclusions and future perspectives

The work described in this thesis has generated new insights into the molecular pathways leading to wood formation from a variety of perspectives, highlighting its universal importance in plant development.

We began by identifying the central transcriptional complex known as Mediator, which is involved in regulation of many essential processes in plants. We also identified subunits of the Mediator complex that are likely to be involved in regulating secondary cell wall formation. Subunits belonging to the tail of Mediator that are conserved within the plant kingdom are likely to perform plant specific functions. We demonstrated the existence of expressed and translated paralogous Mediator subunit genes that are unique to plants. A comparative analysis of Mediator genes from arabidopsis and poplar suggested that such paralogous genes may be involved in the regulation of tissue- and tree-specific traits such as wood formation. A previous study had shown that point mutations at the At2g48110 locus, which encodes Mediator subunit AtMed5#2 (AtMed33b), affect the phenylpropanoid pathway and lignification. This discovery may have practical applications: it has been stated that even modest decreases in the lignin content of biofuel feedstock brought about by such semidominant mutations could increase the efficiency of cellulosic biofuel production. Further, the reduced-lignin phenotypes caused by such transgenes may ultimately be more stable than those generated using RNAi-based strategies over successive generations or years in annual and perennial crops, respectively (Stout et al., 2008). The AtMed25 Mediator subunit is also known as PHYTOCHROME AND FLOWERING TIME 1 (PFT1); as its name suggests, it is involved in flowering. It turns out that the induction of flowering also triggers xylem expansion in the hypocotyl and secondary growth in the roots of arabidopsis (Sibout et al., 2008), suggesting that AtMed25 may have a role in wood formation. Unexpectedly we also found
a conserved domain in the plant protein AtMed25 that corresponds to the interaction domain of the human transcriptional activator Med25-VP16. The finding suggests that it may be possible to develop inducible chimeric systems for medical applications. Inducible systems derived from plant components can rely on chemicals that should have a negligible effect on mammals, or indeed, on animals in general (Corrado & Karali, 2009). From the perspective of plant research, considerable emphasis should be put on finding the activators that interact not only with Med25, but with all of the plant Mediator subunits. The identification of these interacting partners would make it possible to identify the events that trigger the transcriptional initiation of many known pathways back to their primary transcriptional initiation. The identification of such connections would in turn help in fleshing out the details of the mechanisms by which the pathways are regulated.

We suggest that Design of Experiments should be used in all future work involving the simultaneous analysis of levels of multiple different classes of biomolecules, as should our method for the integration of such data as described in paper II. Using these techniques, we were able to separate growth effect from genotype effects in mutants whose growth is severely affected. However, methods for combining data from transcriptomics, proteomics, metabolomics and other -omics are still in their infancy. In terms of proteomics in particular, many challenges remain to be overcome. While it is possible to achieve almost complete coverage of the transcriptome, proteomics is restricted to the analysis of relatively abundant species. Variations in the extent of transcription associated with nuclear processes performed by proteins of low abundance such as Mediator can be observed and isolated. However, although it is likely that such processes are also regulated at the protein level, regulation of this kind cannot be readily analysed with existing technology. In order to analyse more proteins, including sub populations of proteins that have undergone post-translational modification, it will be necessary to combine several different extraction procedures. In poplar and other trees, gene duplications make it difficult to distinguish certain proteins from one-another. Progress in discriminating between the isobaric amino acids isoleucine and leucine would be useful in this respect. Additionally, it would be desirable to automate the identification of markers (i.e. fragments having a specific mass that elute at a known time) that represent individual peptides. Automation could be achieved in a variety of ways. It is theoretically possible to use computational methods to predict the identity of all of the fragments that would be produced by digestion of all of the predicted proteins (of which poplar has
approximately 45 000) of a given species. One could then calculate those peptides' masses and their properties that influence MS detection; the physiochemical properties of peptides are more predictable than those of large proteins. Furthermore, liquid chromatography elution times can also be accurately modelled (Krokhin et al., 2004). In theory, then, it is possible to predict the expected MS ion current of the digested proteome down to the isotope patterns generated by individual peptides. The growing collection of peptides from arabidopsis and poplar that have been identified with a high degree of confidence can be used to identify "proteotypic peptides" that are represented in specific tissues or samples, reducing the complexity of the analysis. As predictions get more accurate, there will be less need to perform numerous identification experiments on actual samples, and therefore smaller quantities of the sampled material will be needed. Ultimately, new algorithms will make it possible to rapidly identify markers with a high degree of confidence. However, the management and analysis of complex samples will still require extensive separation. Methods that exploit the readily-predicted isoelectric points of peptides are attractive in this context. However, current methods based on the isoelectric point require the use of ampholytes. Unfortunately, most ampholytes are incompatible with MS analysis, and so some effort should be focused on the development of innovative ampholyte-free methods that exploit peptide autofocusing (Tomáscaron et al., 2007).

Finally, we examined cell wall synthesis in poplar plasma membranes. This work was initially intended to be a screening study to aid in the planning of future experiments based on the acquisition of quantitative time series to examine processes that occur in the plasma membranes and are important for plant development. However, the "screening" proved to be very successful and identified close to 1 000 proteins, along with their tissue distribution patterns. The sheer number of proteins identified made it possible to analyse the behaviour and activity of different pathways in specific tissues. The identities of many proteins that had been assigned on the basis of single peptide hits from known pathways were supported by their association with others involved the same pathway. We were also able to identify the components of the machinery of cell wall synthesis and the proteins that support it. In future, it may be possible to extend this work by using gel free methods like LCMSE (Silva et al., 2006) to better quantify the proteins. However, the gel based system are useful when analysing plasma membranes, as the gel effectively removes the detergents and lipids that are inevitably present as contaminants in separated plasma membranes.
Compared to other species that have previously been used in proteomics research, poplar presents a number of unusual challenges. We have been able to overcome some of these, and intend to apply the knowledge and experience we have gained by doing so to the analysis of even more challenging species such as spruce and pine.
6 Sammanfattning

Trädstammens vedceller (xylemet) består huvudsakligen av sekundär cellvägg uppbyggd av ett nätverk av cellulosa, hemicellulosa och lignin. Veden bildas av dotterceller från ett tunnt lager av celler (kambiet) till xylenceller som i sin tur genomgår cellexpansion, sekundär cellväggsyntes, lignifiering och apoptos. Under hela utvecklingen av den komplicerade matrisen (veden) deltar ett stort antal proteiner. Biosyntesen av xylemet kräver därför en strikt reglering på molekylär nivå främst genom transkriptionell reglering med olika posttranskriptionella processer.

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