Identification of additive, dominant, and epistatic variation conferred by key genes in cellulose biosynthesis pathway in *Populus tomentosa*†

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†Sequence data corresponding to all candidate genes have been deposited in the GenBank Data Library, and the accession numbers were shown in text or tables.

Edited by Dr Satoshi Tabata

Received 4 August 2014; Accepted 22 October 2014

Abstract

Economically important traits in many species generally show polygenic, quantitative inheritance. The components of genetic variation (additive, dominant and epistatic effects) of these traits conferred by multiple genes in shared biological pathways remain to be defined. Here, we investigated 11 full-length genes in cellulose biosynthesis, on 10 growth and wood-property traits, within a population of 460 unrelated *Populus tomentosa* individuals, via multi-gene association. To validate positive associations, we conducted single-marker analysis in a linkage population of 1,200 individuals. We identified 118, 121, and 43 associations ($P < 0.01$) corresponding to additive, dominant, and epistatic effects, respectively, with low to moderate proportions of phenotypic variance ($R^2$). Epistatic interaction models uncovered a combination of three non-synonymous sites from three unique genes, representing a significant epistasis for diameter at breast height and stem volume. Single-marker analysis validated 61 associations (false discovery rate, $Q \leq 0.10$), representing 38 SNPs from nine genes, and its average effect ($R^2 = 3.8\%$) nearly 2-fold higher than that identified with multi-gene association, suggesting that multi-gene association can capture smaller individual variants. Moreover, a structural gene–gene network based on tissue-specific transcript abundances provides a better understanding of the multi-gene pathway affecting tree growth and lignocellulose biosynthesis. Our study highlights the importance of pathway-based multiple gene associations to uncover the nature of genetic variance for quantitative traits and may drive novel progress in molecular breeding.

Key words: Chinese white poplar (*Populus tomentosa*), epistasis, pathway-based multiple gene association, transcript profiling, validation population
1. Introduction

Trees have unique features that distinguish them from most herbaceous species, including large sizes, long lifespans and woody, perennial growth. Many tree species occur in widely distributed populations that harbour a wealth of diversity. Large, diverse tree populations can be exploited for breeding to improve economically important properties. In addition, deciphering the mechanisms by which forest trees have adapted to their changing environments is an intriguing research problem. Tree growth and wood formation are complex dynamic processes that require coordinate regulation of diverse metabolic pathways in the whole plant. Improving economically important traits in trees by molecular marker-assisted selection (MAS) will require improving our understanding of the molecular mechanisms underlying phenotypic variation, particularly for traits with complex genetic architecture, such as wood properties or growth. Research has been hindered by trees’ large size, long generation times, and the lack of mutants for reverse genetic approaches.

As an alternative approach, recent work has examined complex traits in trees by forward genetic methods, such as quantitative trait loci (QTL) and association mapping of genes or alleles underlying quantitative traits. Single-marker association studies have identified candidate functional single-nucleotide polymorphisms (SNPs) that explain a small proportion of the additive variation (≤5%) on average in growth or wood properties, and many of these genes have not been cloned in full length yet. It is well known that genes do not work in isolation; instead, complex cellular pathways and molecular networks are often involved in phenotypic variation. Variation in quantitative traits includes additive, dominant, and epistatic effects that are conferred by multiple variants within interactive genes in different biological pathways. The complete effects of genetic variation for multiple full-length genes [including promoter, introns, untranslated regions (UTRs), and exons] in related biological pathways remain to be addressed in forest trees. Multi-gene association models take advantage of large numbers of SNPs in population-wide linkage disequilibrium (LD) with QTL regions harbouring functional loci affecting complex traits. Studies using multi-gene association may identify genetic variants that jointly have significant effects but individually make only a small contribution.

Our previous studies using single-marker association (i.e. mixed linear models) have identified sites of SNPs in several candidate genes that located in close proximity to the causal polymorphisms or even the functional variant itself for growth and wood-property traits. To extend these observations, this study investigates the nature of genetic variance (additive, dominant, and epistatic effects) for 11 full-length candidate genes related to cellulose biosynthesis in 10 quantitative traits, using multi-gene association, as well as haplotype-based association approaches, in a population of 460 unrelated individuals of *Populus tomentosa*. To verify the most significant associations, we conducted single-marker analysis in a family-based linkage population consisting of 1,200 individuals. Moreover, tissue-specific transcript abundances of these genes provide a better understanding of the multi-gene network, affecting tree growth and lignocellulose biosynthesis.

2. Materials and methods

2.1. Population materials and phenotypic data

The discovery population (association population) consisted of 460 24-yr-old unrelated individuals, representing almost the entire natural distribution range of *P. tomentosa*, i.e. Southern region, Northwestern region, and Northeastern region (30°–40°N, 105°–125°E). Plants were randomly sampled in 2009 from a collection of 1,047 natural *P. tomentosa* in Guan Xian County, Shandong Province, China (36°23’ N, 115°47’E) and used for the initial multi-gene association (Supplementary Table S1, Du et al.12). In addition, 40 unrelated individuals were randomly selected from this association population and used to identify SNPs.

To confirm the association results identified in the discovery population, a validation population (linkage population) consisting of 1,200 hybrid individuals were randomly selected from 5,000 F1 progeny established by controlled crossing between clone “XY01” (*Populus alba* × *Populus glandulosa*) as the female and clone ‘LM 50’ (*P. tomentosa*) as the male. The progeny were planted in 2008 in the Xiao Tangshan horticultural fields, Beijing, China (40°2’N, 115°50’E) using a randomized complete block design with three replications per genotype.

Ten growth and wood-property traits were scored for all individuals in the two populations, using at least three replications per genotype. The growth traits were tree height, diameter at breast height (DBH), and stem volume. The wood-property traits were holocellulose, α-cellulose, hemicellulose, and lignin content (chemical composition), and fibre length, fibre width, and microfibre angle (physical properties) (Supplementary data S1–S2). The detailed sampling and measurement methods, and phenotypic variance for these 10 traits in the two populations were conducted as previously described. Pearson’s correlations for these 10 phenotypic traits were shown in Supplementary data S3.

2.2. RNA extraction and cDNAs identification

Thirty ramets of 1-yr-old *P. tomentosa* clone ‘LM50’ were selected and planted at Xiao Tangshan horticultural fields, Beijing, China (40°2’N, 115°50’E). A piece of bark was peeled off from the main stem at ∼1.0 m height of 1-yr-old *P. tomentosa* clone ‘LM50’ using a chisel. The newly formed xylem (∼3 mm thick) was first removed from the exposed surface, and then mature xylem was collected by further scraping into the stem at ∼2 mm thick; finally, the collected samples were placed into 50-ml Falcon tubes filled with liquid nitrogen and stored in a freezer at −80°C until RNA extraction. Using the Plant Qiagen RNaseasy kit, total RNA from the stem xylem tissues was extracted and then reverse transcribed into cDNA with the SuperScript First-Strand Synthesis system (Life Technologies, Carlsbad, CA, USA). The stem xylem cDNA library was constructed using the SuperScript cDNA Library System (Life Technologies, Rockville, MD, USA). The detailed procedures for constructing the cDNA library were previously described by Li et al.21 In our study, the cDNA library consisted of 5.0 × 10⁶ pfu with an insert size range of 1.0–4.0 kb. By random end-sequencing of 30,000 cDNA clones and comparison with all sequences in NCBI (http://www.ncbi.nlm.nih.gov), we chose 11 cDNAs with high similarities to *Arabidopsis* or *Populus trichocarpa* sequences to examine the relationship with lignocellulosic biosynthesis (Supplementary Table S2). All cDNA sequences chosen were deposited in GenBank (accession numbers shown in Supplementary Table S2).

2.3. DNA extraction and full-length genomic DNA identification

Using the DNeasy Plant Mini kit (Qiagen China, Shanghai), total genomic DNA was extracted from fresh young leaves. For sequencing the full-length genomic DNA sequences, specific primers were designed based on each cDNA sequence. PCR amplification was performed...
according to Zhang et al.\textsuperscript{22} All the genomic DNA of candidate genes were obtained by direct sequencing of both strands, using conserved T7 and SP6 primers in the BigDye Terminator Cycle Sequencing Kit (version 3.1; Applied Biosystems, Foster City, CA, USA) and a 4300 DNA Analyzer (Li-Cor Biosciences, Lincoln, NE, USA). Full-length genomic DNA sequences of all 11 candidate genes were isolated by PCR amplification from the \textit{P. tomentosa} LM50 clone (Supplementary Table S2). Of these, five of these genes were previously reported in single-marker association studies (Supplementary Table S2). Also, these 11 full-length gene sequences were deposited in GenBank (accession numbers shown in Supplementary Table S2).

2.4. Tissue-specific expression analysis

Fresh tissues (root, stem phloem, stem cambium, developing xylem, mature xylem, expanding leaf, expanded leaf, and apex) were sampled from 1-yr-old plants of \textit{P. tomentosa} (clone LM50). Developing xylem tissues were collected by scraping the thin (∼1.0 mm) and partially lignified layer on the exposed xylem surface of main stems at the 1.0 m height of plant, and the liquid stem cambium was collected from the exposed xylem surface as described previously.\textsuperscript{21} The main root (5 cm in length), expanding leaf (the 2–3rd leaf from the stem tip), expanded leaf (the 5–6th leaf), and the final shoot apex (2–3 mm in length) were immediately collected from 1-yr-old \textit{P. tomentosa} clone ‘LM50’. The extraction of RNA from various fresh tissues and the reverse transcription into cDNA was used for testing of tissues-specific transcript abundance for all 11 candidate genes. Real-time quantitative PCR (RT-qPCR) was performed on a 7500 Fast Real-Time PCR System (ABI) using the LightCycler-FastStart DNA master SYBR Green I kit (Roche). The qPCR program and the real-time amplification reaction were performed as described by Du et al.\textsuperscript{16} All reactions were performed in triplicate technical and triplicate biological replications, respectively. The melting curve was used to check the specificity of the amplified fragments, and all data were analysed using the Opticon Monitor Analysis software 3.1 tool (Bio-Rad, USA), following the manual protocol. The results obtained for the different tissues were standardized to the levels of \textit{Actin} gene (the internal control). The specific primer pairs were individually designed to target the 3′ UTR of each gene, using Primer Express 3.0 software (Applied Biosystems) (Supplementary Table S3). Mean value and standard deviation (S.D.) for expression data of all repetitions in each set was determined using SPSS Statistics Version 19.0 (SPSS Inc., Chicago, IL, USA) (Details not shown).

To better understand the co-expression interactions among these candidate genes, Pearson’s correlations between each pair of genes were determined for all 11 genes, according to their tissue-specific transcript abundances in SPSS Statistics, version 19.0. A structural network representing significant gene–gene correlations was visualized using the NodeXL Excel Template, version 1.0.1.238 (http://nodexl.codeplex.com/).

2.5. SNP discovery and genotyping

To identify SNPs, full-length genomic DNA of all 11 candidate genes were sequenced and aligned among 40 unrelated individuals from the association population of \textit{P. tomentosa}, without considering insertions/deletions (INDELS). All 440 sequences were deposited in GenBank (Supplementary Table S4). Sequence alignments and manual editing were performed as described by Du et al.\textsuperscript{16} Next, all common SNPs (minor allele frequency ≥0.05) identified were genotyped using the Beckman Coulter (Franklin Lakes, NJ, USA) sequencing system.

2.6. Nucleotide diversity and LD

Diploid sequences were phased into haplotypes with the Phase v. 2.1, using 10,000 iterations of the Bayesian Markov Chain Monte Carlo (MCMC) Model.\textsuperscript{23} We then used alignments of these 40 phased haplotypes for each gene to estimate the number of segregating sites, nucleotide diversity, neutrality tests, and the pattern of codon usage with DnaSP version 5.10.\textsuperscript{24} Nucleotide diversity was estimated using the average number of pairwise differences per site (\(\pi\))\textsuperscript{25,26} and the number of segregating sites (\(θ\)).\textsuperscript{27} Neutrality test statistics, Tajima’s \(D\)\textsuperscript{28} and Fu and Li’s \(D^*\) and \(F^*\),\textsuperscript{29} were calculated using data for the whole population with 10,000 coalescent simulations. The pattern of codon usage was measured using relative synonymous codon usage (RSCU).\textsuperscript{30}

The squared correlation of allele frequencies \((r^2)\)\textsuperscript{31} between each pair of common SNPs (frequency ≥0.05) within candidate genes was calculated with 10\(^5\) permutations, using TASSEL Ver. 2.0.1 (http://www.maizegenetics.net/). To assess the extent of LD within the sequenced gene regions, the decay of LD with physical distance (base pairs) within each candidate locus and over all candidate genes was estimated in 10,000 permutations of the data by non-linear regression.\textsuperscript{32} Singletons were excluded in the LD analyses. This analysis was done both within the three climatic regions and for the complete data set.

2.7. Association analysis

2.7.1. Multi-SNP additive and dominance models

fGWAS, version 2.0 (http://statgen.psu.edu/software/fgwas-soft.html, Li et al.\textsuperscript{33}), in R (http://cran.r-project.org/) was used to conduct multi-SNP association analyses. For each trait, the genotyping data for all common SNPs and their genomic positional information were identified and standardized using PLINK version 1.07 (http://pngu.mgh.harvard.edu/~purcell/plink/res.shtml). This software is a Bayesian hierarchical model and use a MCMC algorithm to simultaneously fit and estimate the possible additive and dominant effects associated with all SNPs. The proportion of the phenotypic variance \((R^2)\) explained by each particular SNP was estimated in the models. Positive dominance represents that the phenotypic mean value observed within the heterozygous class is higher than the average phenotypic mean across both homozygous classes based on Least Squares Means, whereas negative dominance was opposite (not a recessive genetic model). To address the total phenotypic variance accounted for by all trait-associated SNPs on a trait-by-trait basis, we calculated a cumulative \(R^2\) metric. These values were obtained by the difference in \(R^2\) between full and reduced models.\textsuperscript{34} The details were referenced according to McKown et al.\textsuperscript{35}

2.7.2. Multi-SNP epistasis models

A multifactorial dimensionality reduction (MDR) method\textsuperscript{36} that was designed specifically for detecting and characterizing non-additive interactions (i.e. epistasis). The ReliefF algorithm was applied to filter all unlinked SNPs \((r^2 < 0.1\) or different genes) to the best five loci for each trait that provide the greatest signal with 100 nearest neighbours. MDR was applied to assess all two-way to five-way models for the five significant filtered SNPs. Finally, an independent assessment of epistasis was performed using entropy-based measures of information gain.\textsuperscript{37} Specifically, a new measure model of three-way epistasis that adjusts for lower order effects was used to examine high-order non-additive interactions.\textsuperscript{38}

2.7.3. Haplotype-based association models

All high-LD haplotypes \((r^2 \geq 0.70, P \leq 0.001)\) were estimated for each gene, and their frequencies were determined using HaploView version
4.2 (http://www.broad.mit.edu/mpg/haploview.html). Singleton alleles and haplotypes with a frequency <5% were discarded when constructing the haplotypes. Haplotype-based association tests with phenotypic traits were performed using the Haplo.stats package in R, \(^{39}\) and significances of the haplotype associations were identified based on \(10^4\) permutation tests. The input consisted of genotype matrices, phenotype matrices, and structure analysis matrices (Q) to correct for population structure.\(^{40}\) In this analysis, the Q matrix was identified according to the optimal subpopulation structure in our association population.\(^{41}\) Corrections for multiple testing of smoothed \(P\)-values for all associations were performed using the false discovery rate (FDR) through QVALUE.\(^{42}\) A \(q\)-value of 0.10 was considered as the significance threshold.

2.7.4. Single-marker linkage analysis
On the basis of significant SNPs identified above, gene sequences were compared between hybrid parents, and SNP markers that segregated in the 1,200 \(F_1\) progeny were selected. Inheritance tests of these SNPs were examined in the linkage population by performing a \(\chi^2\) test at 0.01 probability, and SNPs following Mendelian expectations were used in single-marker analysis using PLINK version 1.07. The same FDR method was used to correct for multiple tests.\(^{42}\)

3. Results

3.1. Tissue-specific transcript profiling
To provide insights into gene interactions and functions in the biological pathway, we examined transcript accumulation for 11 candidate genes in various tissues and organs of \(P.\) tomentosa, including root, stem, and leaf (Fig. 1A). We found that the 11 genes exhibited distinct, but partially overlapping patterns of expression. Most genes were preferentially expressed in the developing xylem and mature xylem, with moderate expression in the cambium region and expanded leaves (Fig. 1A). For example, in the stem of \(P.\) tomentosa, Cellulose synthase 7 (PtoCesA7), UDP-glucose dehydrogenase 1 (PtoUGDH1), and S-adenosine-L-homocysteine-hydrolase 1 (PtoSAHH1) transcripts were most abundant in the developing and mature xylems, with moderate abundance in the cambium. PtoCesA7

![Figure 1. Tissue-specific transcript profiling and a correlation network of 11 candidate genes in Populus tomentosa. (A) Relative transcript levels of 11 candidate genes in different tissues and organs of P. tomentosa. Expression levels were normalized to the Actin gene; Apex, apical shoot meristem, the error bars represent ±standard deviation (SD). (B) A gene–gene correlation network was constructed, on the basis of comparison and correlation analyses of transcription abundances among these genes. Solid lines represent positive correlations and dashed lines represent negative correlations; the thickness of each line indicates the strength/significance of correlations, \(P<0.05\) level of significance (thin lines), \(P<0.01\) (thick lines). Data were log-transformed before Pearson’s correlation.]

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and PtoUGDH1 were also preferentially expressed in roots (Fig. 1A), whereas PtoCesA3 and UDP-glucuronate decarboxylase 1 (PtoUXS1) were highly expressed in the apex (Fig. 1A). PtoUXS1 were most abundant in expanded leaves.

Next, we constructed gene–gene correlation networks using the patterns of tissue-specific expression of these 11 candidate genes and identified 20 positive or negative gene–gene correlations \( (P \leq 0.05) \) (Supplementary Table S5) that make up a highly interrelated network (Fig. 1B). Of these gene pairs, PtoCesA3 and Sucrose synthase 1 (Pto-SUS1) \( (P \leq 0.01, R = -0.938) \), PtoSAHH1 and PtoSAHH2 \( (P \leq 0.01, R = 0.959) \), and PtoCesA4 and Endo-1,4-β-glucanase 1 (PtoGH9A1) \( (P \leq 0.01, R = 0.952) \) showed highly significant correlations (Supplementary Table S5).

### 3.2. SNP discovery and nucleotide diversity

We first used a re-sequencing approach to identify SNPs in all 11 candidate genes. By sequencing the 11 full-length genes from 40 unrelated individuals encompassing nearly the entire natural range of \( P. tomentosa \), we identified 2,114 SNPs in the \( c. \) 47,300 bp sequenced, with an average density of one SNP per 25 bp. Of these SNPs, \( c. \) 44.0% (930) were common sites \( (\text{frequency} \geq 0.05) \) (Supplementary Table S4). The SNPs were not evenly distributed, with the number of SNPs per gene ranging from 59 to 388 (average, 192), consistent with varying levels of nucleotide diversity among genes (Supplementary Table S4). Using sequences from the \( P. trichocarpa \) reference as an out-group, a survey of the single-base mutation types indicated that \( C : T \) changes were the most abundant (38.6%), followed by \( A : G \) changes (30.4%), and overall SNPs the ratio of transitions to transversions was 2.23.

Within the coding regions of all 11 genes (22,098 bp total sequence), the average non-synonymous nucleotide diversity \( (\pi_N) \) was \( \sim5 \)-fold lower than the synonymous nucleotide diversity \( (\pi_S) \), and the \( d_N/d_S \) ratio was \( <1 \) for all exons, indicating strong purifying selection at non-synonymous sites at these \( Populus \) genes. To investigate the extent of codon bias in \( P. tomentosa \), we calculated RSCU values for all types of codons and found that the 11 candidate genes show similar patterns of RSCU (Supplementary Fig. S1). Most of the preferred codons end with \( U \), but \( A \) and \( G \) were also less commonly found in the third position in these genes. Previous studies indicated that codon bias (with \( C \) or \( G \) ending codons) is expected to be strongest for highly expressed genes. Thus, our observation may be consistent with low expressed values for all these genes (Fig. 1A). More work is needed to sort these issues out in \( P. tomentosa \). The values of \( Fu \) and \( Li \)'s \( D \) statistical tests were negative for all genes, with four significant departures observed in the whole population \( (P \leq 0.05, \text{Supplementary Table S4}) \), revealing an excess of low-frequency polymorphisms in the species-wide samples.

### 3.3. SNP genotyping and LD

We selected a set of 651 common SNPs that met the quality control thresholds and that had genotype frequencies consistent with Hardy–Weinberg equilibrium (Supplementary data S4). The percentage of successfully genotyped SNPs that were found in non-coding regions of the genes was 75.4%, distributed within introns (45.4%), 5′ UTRs (16.9%), and 3′ UTRs (13.1%). Remaining SNPs were located in the coding regions, either at synonymous sites (18.5%) or at non-synonymous sites (6.1%).

Estimates of \( r^2 \) values for all pairwise combinations of SNPs were pooled to assess the overall pattern of LD with physical distance (Fig. 2). The non-linear regression shows a clear and rapid decline of LD from 0.60 to 0.10 at a distance of \( \sim1,100 \) bp in the whole population (Fig. 2). Nevertheless, LD analyses within each geographical climatic region showed a much higher level of LD with the \( r^2 \) values declining to 0.1 within \( c. \) 2,300 bp (Southern and Northeastern regions) and \( c. \) 3,400 bp (Northwestern region) (Fig. 2). We found a clear and rapid decline of LD with distance within each gene in the whole population, respectively \( (r^2 \geq 0.1, \text{within \( c. \) 500–1,800 bp, Supplementary Fig. S2}) \), indicating that LD does not extend over entire gene in the species.

### 3.4. Multi-SNP association under additive, dominance, and epistasis models

We first employed Bayesian hierarchical models, emphasizing multi-SNP additive and dominant effects for each quantitative trait, and uncovered a multitude of genetic associations, including some not previously found by single-marker methods (Supplementary Table S6). Some particular SNP-trait associations detected previously were examined in this case; for example, a non-synonymous substitution in exon3 of PtoCesA4 (PtoCesA4_1914) was shown to be in strong association with \( \alpha \)-cellulose. We then constructed a structural network to visually represent all 123 associations \( (P < 0.01) \) between all 10 growth and wood-property traits and 93 unique SNPs (Fig. 3). The 93 loci were not evenly distributed among all 11 genes, with a range from 1 (PtoBGlu1, \( \beta-1,4 \)-glucosidase) to 19 (PtoUXS1). 37.6% of the 93 loci were distributed in coding regions, and among these, 94.3% of the associations contained a combination of additive and dominant effects (Supplementary Table S6).
Although the association result does not indicate which SNPs may be causal, 11 of these 92 unique SNPs represent amino acid replacement (non-synonymous) polymorphisms (Supplementary Table S6). For each trait, the number of significant SNPs ranged from 8 to 16, which captured a small to moderate fraction (17.0–36.0%) of the additive variation when considered jointly (Table 1, Fig. 4).

Correspondingly, each trait was associated with variation in at least five candidate genes (Table 1). Furthermore, we identified three genes (PtoCesA4, PtoCesA7, and PtoSAHH2) that were associated to all four wood chemical compositions. (Supplementary Table S6). For wood physical properties, SNP-trait associations were identified for four genes (PtoCesA3, PtoCesA4, PtoSUS1, and PtoUXS1) that were associated with all three traits. In addition, PtoCesA7 and PtoSAHH2 were associated with fibre length and microfibre angle (Supplementary Table S6). Additive models found nine genes with SNPs associated with variation in growth traits; of these, two genes (PtoCesA3 and PtoUXS1) with SNPs associated across all three growth traits, as well as identified with all three wood physical properties (Supplementary Tables S6). At least five SNPs within PtoUXS1 had
associations to each growth trait. No significant SNP from P.toBGlu1 and P.toBGlu1 was associated with growth traits.

On the basis of these additive SNPs simultaneously associated to each trait (Supplementary Table S6), we identified phenotypic variation among possible genotype combinations for the same trait and some genotype combinations that may be useful for selection breeding. An example of multi-genotype combinations for lignin is shown in Fig. 5. Eight SNPs from six different genes were significantly associated with lignin content, with two associated SNPs from P.toGH9A1 showing low LD and hence representing independent associations.

3.4.2. Multi-gene associations under the dominance model
Under the dominance model, we detected 121 significant associations, including 72 associations with positive dominance values and 49 with negative values, representing a total of 92 unique SNPs within the 11 genes across all three trait categories (10 traits) (Table 2 and Supplementary data S6). Of the 92 SNPs, ~95% of the significant SNPs had both additive and dominance effects on all traits within each category (Supplementary Table S6). Many genes were repeatedly associated with multiple traits within/across trait categories, and different SNPs with different effects from the same gene were identified. A number of SNP associations with positive versus negative effect across the three trait categories were 25/16 for wood chemical compositions, 25/12 for wood physical properties, and 22/21 for growth traits (Table 2).

When we considered dominant variation per trait by combining the positive and negative types, the total number of significant SNPs varied from 8 to 18, with the total $R^2$ explained ranging from 1.8 to 13.0% (Fig. 4).

The 72 associations showing positive dominant effects represented 59 SNPs from 10 genes associating with all 10 traits. Of these, 22 SNPs were located in the coding regions, including 7 non-synonymous and 15 synonymous changes. A number of significant SNPs per gene varied from 1 to 11. For each trait, the number of SNPs ranged from 3 to 10, with total $R^2$ explained ranging from 9.0 to 30.0% (Table 2).

For the 49 associations showing negative dominant effects, we identified 44 loci from 10 genes that were significantly associated with 10 traits, with a small $R^2$ explained per locus ranging from 0.6 to 5.4%, respectively. The number of significant SNP within each gene varied from 1 (P.toBGlu1) to 11 (P.toUXS1). A range of 2–8 SNPs with the negative dominant effects was found for each trait, with total variation explained ranging from 4.4% ($\alpha$-cellulose) to 17.3% (microfibre angle) (Table 2). Ten SNPs from six genes that simultaneously have positive and negative dominant effects for different traits (Supplementary Table S6). For example, the synonymous SNP P.toSAHH2_285 had a negative dominant effect for lignin, while having a positive effect for holocellulose. Table 2 shows detailed descriptions of the genetic parameters for all traits under positive and negative dominant effects.

3.4.3. Multi-gene associations under the epistasis model
We used the MDR method (http://phg.mc.vanderbilt.edu/Software/MDR) to detect and characterize SNP–SNP epistasis associated with these traits (Supplementary Tables S7 and S8). We first applied the Relief algorithm to filter all unlinked SNPs ($r^2 < 0.1$ or located in different genes) to identify significant 5 loci per trait. After correcting for multiple testing, we found 42 associations ($Q \leq 0.10$) across nine traits. These associations represent 34 unique SNPs from seven genes with main effects varying from 0.1 to 7.5% (Supplementary Table S7). We found that 9 of the 34 unique SNPs observed in the epistasis models also showed significant additive and dominant effects. Using multi-way SNP–SNP interaction models for these filtered SNPs, we identified 38 significant SNP–SNP pairs where the epistatic interactions ranged from −7.0 to 0.9% (Supplementary Table S8). The majority of epistatic effects were found between unlinked loci located in different genes. We also detected epistatic effects among different SNPs within the same gene. For instance, two epistasis pairs consisting of three SNPs within P.toCesA4 were identified for fibre length, all with negative interaction values (Supplementary Table S8).

Three-way interaction model provided a combination of three non-synonymous loci (P.toCesA4_1914, P.toGH9A1_1936, and P.toGA20ox_632) from three unique genes, representing a significant epistatic interaction for DBH and stem volume, with three-way epistatic effects of −11.7 and −12.1% (Fig. 6A). Negative epistatic effects suggest redundancy between loci, meaning that these loci provide, in part,
Figure 5. Phenotypic variations among the possible genotype combinations for lignin under multi-SNP additive models. (A) These eight allelic variants have significant additive effect with a small proportion of phenotypic variation ($R^2$), ranging from 0.80 to 5.48%. Pairwise LD plots among multiple loci within each candidate genes were estimated using TASSEL Ver. 2.0.1 (http://www.maizegenetics.net/). Genotype effect and the position/haplotype block of each allelic variant were shown for lignin content. (B) Twenty possible common combinations with a frequency $\geq 5\%$ from the eight allelic variants were identified with various phenotypic variations on lignin content in the association population of Populus tomentosa. Some combinations were discarded, because the sample size was <10 individuals.

Table 2. Summary of the dominant effect and phenotypic contribution rate ($R^2$) of all significant SNPs for each trait in the Populus tomentosa association population.

| Trait                  | Number of candidate genes | Number of SNPs | Range of dominance effect (%) | Range of dominant effect (%) | Range of $R^2$ (%) | Total $R^2$ (%)$^a$ | Negative/positive$^b$ |
|------------------------|---------------------------|----------------|-------------------------------|-------------------------------|--------------------|----------------------|------------------------|
| Lignin (%)             | 6                         | 6              | −1.347 to −0.125              | −1.347 to −0.125              | 0.6 to 2.0         | 7.2                  | N                      |
|                        | 3                         | 3              | 1.579 to 2.192                | 1.579 to 2.192                | 2.4 to 3.5         | 9.0                  | P                      |
| Holocellulose (%)      | 4                         | 4              | −2.482 to −0.08               | −2.482 to −0.08               | 0.8 to 3.6         | 10.2                 | N                      |
|                        | 3                         | 8              | 0.941 to 2.371                | 0.941 to 2.371                | 1.7 to 4.0         | 22.6                 | P                      |
| Hemicellulose (%)      | 2                         | 2              | −2.110 to −1.527              | −2.110 to −1.527              | 3.3 to 4.9         | 8.1                  | N                      |
|                        | 3                         | 6              | 0.623 to 2.221                | 0.623 to 2.221                | 0.6 to 5.1         | 17.6                 | P                      |
| α-Cellulose (%)        | 4                         | 4              | −0.973 to −0.170              | −0.973 to −0.170              | 0.7 to 2.0         | 4.4                  | N                      |
|                        | 5                         | 8              | 0.073 to 3.347                | 0.073 to 3.347                | 0.6 to 3.9         | 16.6                 | P                      |
| Fibre length (mm)      | 4                         | 4              | −0.043 to −0.009              | −0.043 to −0.009              | 0.9 to 2.5         | 6.6                  | N                      |
|                        | 4                         | 8              | 0.008 to 0.071                | 0.008 to 0.071                | 0.6 to 4.5         | 17.1                 | P                      |
| Fibre width (µm)       | 2                         | 2              | −1.420 to −0.515              | −1.420 to −0.515              | 1.1 to 3.5         | 4.6                  | N                      |
|                        | 5                         | 7              | 0.179 to 1.301                | 0.179 to 1.301                | 0.6 to 3.1         | 15.9                 | P                      |
| Microfibre angle (MFA, °) | 4                        | 6              | −3.882 to −0.433              | −3.882 to −0.433              | 1.2 to 5.4         | 17.3                 | N                      |
|                        | 6                         | 10             | 0.180 to 3.697                | 0.180 to 3.697                | 0.6 to 5.3         | 27.9                 | P                      |
| Diameter at breast height (DBH, cm) | 5                        | 6              | −3.028 to −0.452              | −3.028 to −0.452              | 0.9 to 3.0         | 12.0                 | N                      |
|                        | 4                         | 6              | 0.694 to 2.785                | 0.694 to 2.785                | 1.2 to 3.2         | 14.4                 | P                      |
| Tree height (m)        | 6                         | 8              | −2.341 to −0.126              | −2.341 to −0.126              | 0.7 to 4.4         | 17.0                 | N                      |
|                        | 7                         | 10             | 0.202 to 2.303                | 0.202 to 2.303                | 1.0 to 4.9         | 30.0                 | P                      |
| Stem volume (m³)       | 4                         | 7              | −0.350 to −0.085              | −0.350 to −0.085              | 1.7 to 3.9         | 17.0                 | N                      |
|                        | 3                         | 6              | 0.044 to 0.389                | 0.044 to 0.389                | 1.0 to 4.1         | 12.6                 | P                      |

Significance level for association (significance is $P \leq 0.01$).

$^a$Total $R^2$ was calculated according to McKown et al. 35

$^b$N: negative dominant effect; P: positive dominant effect; see Supplementary Table S6 for further details of these association results under dominance models.
the same information for the traits. Figure 6B shows the mean values of phenotypic variances among all three genotype combinations. We investigated the nature of epistatic interactions among non-synonymous mutations, contributing to the phenotypic variation, in the three genes, and discovered that most of the variation is captured by combinatorial permutations of allelic variants at three loci (Fig. 6C). We identified the specific combinatorial patterns of amino acid replacements at these three loci, by comparing the expected values of eight combinations with high (H) or low (L) phenotypic values with the observed values of the specific combinatorial patterns (Fig. 6).

Overall, using multi-gene association models, we captured all three components of genetic variation conferred by multiple variants within candidate genes in cellulose biosynthesis pathway both within and across these three trait categories. In total, we identified 118, 121, and 43 associations consistent with additive, dominant, and epistatic effects, respectively (Supplementary Tables S6–S8). Also, we observed no differences in the magnitudes of additive and dominant effects between three trait categories (Tables 1 and 2). Some genes have extensive complexity in both genetic variation and the resulting phenotype, and numerous SNPs from different genes were associated across these three trait categories, suggesting that tree growth and wood biosynthesis are complex, dynamic processes that require the coordinate regulation of diverse sets of genes. Notably, our present investigation suggests that CesA members may have additional functional roles in tree growth and development, beyond the direct effects of the cellulose synthase subunits in biosynthesis of the cell wall. CesA members represent an interesting set of candidates for further study in trees.

3.5. Validation of association in the linkage population

To validate the associations identified above, we examined associations in a linkage population. From the 118 unique significant SNPs obtained by multi-SNP and MDR models (Supplementary Tables S6 and S7), we identified 109 common SNP markers that segregated in the 1,200 F1 progeny following Mendelian expectations (P ≥ 0.01, Supplementary data S5). We conducted 1,090 tests for association of these SNP with growth and wood-quality traits (109 SNPs × 10 traits) in the linkage population. After correction for multiple testing errors, we found 61 associations at the threshold of Q ≤ 0.10, representing 38 SNPs from nine genes that show associations with all 10 traits (Supplementary Table S9). These 38 unique SNPs included 5 non-synonymous, 15 synonymous, and 18 non-coding SNPs (Supplementary Table S9). Thirty SNPs were similarly detected by multi-SNP additive and dominance models in the discovery population, and nine loci were shared between the epistasis models and the single-marker
analysis (Supplementary Tables S6 and S7). Most of the significant markers were identified for multiple traits with different genotypic effects; for example, PtoCesA4_1914 is associated with seven traits, and PtoGA20ox_632 with six (Supplementary Table S9). In addition, 28 (46.0%) associations identified in the validation population were similarly detected in the multi-SNP models (Supplementary Tables S6 and S9). For each trait, the number of SNPs ranged from 3 to 9, with marker effects that varied from 2.2 to 13.5% (Supplementary Table S9). Effect sizes for the SNPs analyzed using single-marker linkage test were nearly 2-fold higher than effects identified using multi-SNP models (average $R^2 = 3.8\%$).

Correspondingly, each trait is associated with at least three candidate genes (Supplementary Table S9) using single-marker linkage analysis. With wood chemical composition, we identified two genes (PtoCesA4 and PtoCesA7) that were simultaneously associated with the content of all four polysaccharide traits. In addition, we identified that PtoGH9A1 was significantly associated with hemicellulose and lignin contents, whereas PtoCesA3 was associated with hemicellulose and $\alpha$-cellulose content (Supplementary Table S9). Among wood physical properties and growth traits, SNP-trait associations identified two genes (PtoCesA4 and PtoGA20ox) that were associated with all six traits. Furthermore, PtoUXS1 has SNPs associated with both DBH and stem volume traits, as well as two wood physical properties (fibre width and microfibre angle), consistent with the associations under multi-SNP models in our discovery population (Supplementary Tables S6 and S9). No SNPs from PtoUGDH1 and PtoBGlu1 were associated with any of the 10 phenotypic traits.

3.6 Haplotype-based association analysis

We identified 138 high-LD blocks ($r^2 \geq 0.70$, $P \leq 0.001$), including 429 common haplotypes (frequency $\geq 5\%$) from the 11 candidate genes, with block sizes from 2 to 564 bp. The number of common haplotypes per block varied from 2 to 7, with an average of 3.0 (Table 3). The number of LD blocks and common haplotypes per gene varied from 4 to 20 and 15 to 60, respectively (Table 3). Haplotype-based association tests identified a total of 162 common haplotypes from 66 blocks at $P \leq 0.05$ (Table 3 and Supplementary data S10). Using multiple test corrections, we found 39 significant blocks from 10 candidate genes, including 90 haplotypes, that associated with nine traits excluding stem volume (139 associations, $Q \leq 0.10$); no significant haplotypes were found for PtoSAHH2. The number of blocks and haplotypes for each trait ranged from 2 to 12 and 5 to 33, respectively (Supplementary Table S11).

Most of the significantly associated genes/blocks were shared among traits (Supplementary Table S11). For example, haplotypes belonging to PtoUXS1 were associated with seven traits. PtoCesA7 had haplotypes that were simultaneously associated with all four chemical composition traits. In addition, PtoUGDH1 with haplotypes were significantly associated with three chemical compositions, excluding $\alpha$-cellulose content (Supplementary Table S11). PtoUXS1 had haplotypes that were associated with all three wood physical properties. Within the growth trait category, several significant haplotypes were identified within PtoUXS1 and PtoSUS1 that were associated with tree height and DBH. Examination of haplotypes may provide support for the SNP-based association.

On the basis of the significant blocks we identified within several genes for each trait (Supplementary Table S11), we identified four exclusive multi-haplotype combinations with numerous genetic variants for three traits—holocellulose, lignin, and tree height (Supplementary Fig. S3). Taking holocellulose content as example, we identified several combinations of multiple haplotypes from two significant blocks in PtoCesA4 and PtoGH9A1, with mean values from 66.725 to 75.321%, consistent with the two genes having significant correlation ($P \leq 0.01$, $R = 0.952$) (Fig. 1B). In addition, eight unique haplotypes from four genes, showing specific (presence/absence) for different climatic regions of the natural population of P. tomentosa, were associated with five traits, respectively (Table 4). For fibre width, two haplotypes (C-A-G and T-A-G) from PtoSUS1 and PtoBGlu1 were exclusively found in the Northwestern region, and were associated with higher fibre width (Supplementary Table S11).

| Table 3. Summary of haplotype-based association analysis within 11 candidate genes for each trait in the Populus tomentosa association population |
|---------------------------------|---------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| Abbreviation of gene            | Gene name                       | Number of LD block | Number of common haplotypes | Length range of haplotype (bp) | Number of haplotypes ($P \leq 0.05$) | Number of haplotypes, FDR corrected ($Q \leq 0.10$) | Number of associated traits |
|---------------------------------|---------------------------------|---------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|-------------------------------|
| PtoBGlu1                         | $\beta$-1,4-glucosidase         | 4                   | 15                            | 35–564                        | 4                             | 4                             | 4                             |
| PtoCesA3                         | Cellulose synthase              | 20                  | 58                            | 31–431                        | 20                            | 12                            | 3                             |
| PtoCesA4                         | Cellulose synthase              | 19                  | 57                            | 2–320                         | 16                            | 11                            | 4                             |
| PtoCesA7                         | Cellulose synthase              | 16                  | 60                            | 12–484                        | 23                            | 9                             | 5                             |
| PtoGA20Ox                       | GA20-oxidase                    | 9                   | 30                            | 52–246                        | 10                            | 7                             | 2                             |
| PtoGH9A1                        | Endo-1,4-$\beta$-glucanase      | 14                  | 41                            | 27–274                        | 19                            | 9                             | 5                             |
| PtoSAHH1                        | S-adenosine-1-homocystene-hydrolase | 8                 | 30                            | 60–202                        | 11                            | 6                             | 1                             |
| PtoSAHH2                        | S-adenosine-1-homocystene-hydrolase | 5               | 21                            | 77–355                        | 6                             | 0                             | 0                             |
| PtoSUS1                         | Sucrose synthase                | 20                  | 46                            | 8–395                         | 25                            | 14                            | 6                             |
| PtoUGDH1                        | UDP-glucose dehydrogenase       | 7                   | 28                            | 47–192                        | 8                             | 8                             | 3                             |
| PtoUXS1                         | UDP-gluconate decarboxylase     | 16                  | 43                            | 8–203                         | 20                            | 10                            | 7                             |
| Total                           | /                               | 138                 | 429                           | 2–564                         | 162                           | 90                            | /                             |

a: not applied.
b: $P$-value = significance level for association (significance is $P \leq 0.05$).
c: $Q$-value = a correction for multiple testing [FDR ($Q \leq 0.10$); see Supplementary Tables S10 and S11 for further details of these haplotype-based association results.]
4. Discussion

4.1. Functional interpretations of genetic associations

Here we have used multi-gene association approaches, combining extensive full-length candidate genes from a shared biological pathway and with detailed, replicated phenotypic information from natural populations of *Populus tomentosa* to uncover numerous loci underlying variation in wood chemical composition, physical properties, and growth traits. We further validated these associations in an independent linkage population. Significant additive, dominant, and epistatic effects were observed for tree growth and lignocellulosic composition. The number of associations we detected is larger, while the average $R^2$ explained individually is lower, than in previous single-marker association for similar traits in the association population. Also, it is possible that linkage drag in regions around the candidate genes causes larger $R^2$ values of allelic variants in the linkage population than in the association population. More statistics are needed. Markers found to be associated with traits span both coding and non-coding portions of genes, with a predominance for additive effects and small effect sizes (Tables 1 and 2), suggesting that this best describe the genetic architecture of complex traits in tree species.

<i>Table 4</i>. List of climatic region-specific haplotypes significantly associated with wood quality and growth traits in *Populus tomentosa* after correction for multiple testing (FDR ($Q ≤ 0.10$)).

| Trait                | Probability (FDR) | $Q$-value | Significant haplotypes | Amplicon | Phenotypic variation (mean ± SE) |
|----------------------|------------------|-----------|-----------------------|----------|----------------------------------|
| Holocellulose        | 0.0010           | 0.0373    | C-G-G-C               | PtoBGlu1_4972-4990-5014-5164 | 0.005  |
| Hemicellulose        | 0.0010           | 0.0275    | A-C                   | PtoCesA3_555-571 | 0.325  |
| Fibre length         | 0.0001           | 0.0160    | C-A-G                 | PtoUXS1_1678-1685-1707-1848-1862-1884-1899 | 0.111  |
| Fibre width          | 0.0010           | 0.0373    | T-A-G                 | PtoUXS1_3000-3005-3043-3054-3059 | 0.111  |
| Tree height          | 0.0001           | 0.0160    | C-G-T-G-C             | PtoUXS1_3199-3242-3301 | 0.111  |

These AMPs were selected based on the significance squared correlation of allele frequencies ($R^2$) at $P < 0.001$. The phenotype was found in the climatic region: NE, northeastern region; S, southern region; NW, northwestern region; H, standard error.
Genetic variation conferred by key genes in cellulose biosynthesis pathway in *Populus*.

Taking α-cellulose content as an example, we identified six common genes significantly related to this trait, specifically *PtoBGlu1*, *PtoCesA3*, *PtoCesA4*, *PtoCesA7*, *PtoGA20Ox*, and *PtoSUS1* (Table 5). When taken together, multi-SNP models explained a small to moderate portions of the variation from both additive and dominant effects (31.7 and 12.2%, respectively). Previous evidence suggests that these six genes may play key roles in cellulose biosynthesis. Supporting these observations, SNPs in *PtoSUS1* showed high additive and dominant effects (Supplementary Table S6), and two LD blocks, including five significant haplotypes (Supplementary Table S11), were associated with α-cellulose. In plants, SUS catalyzes a reversible reaction that preferentially converts sucrose into fructose and UDP-glucose. UDP-glucose is the immediate precursor for the synthesis of cellulose. Another enzyme involved in cellulose formation during xylem development is β-glucosidase (BGlu), which hydrolyzes cellulose to glucose, and is a rate-limiting factor during enzymatic hydrolysis of cellulose. Here, we identified two haplotypes and a non-coding SNP in *PtoBGlu1* (Supplementary Tables S6 and S11). Furthermore, korrigan cellulase (KOR), an endo-β-1,4-glucanase, belonging to the glycosyl hydrolase family 9 (GH9), can hydrolyze the β-1,4 linkage of the cellulose chain. *PtoGH9A1*, an orthologue of KORRIGAN1, contains three haplotypes associated with α-cellulose (Supplementary Table S6), and we also found epistatic interactions for the trait between a non-synonymous SNP (PtoGH9A1_1936) and SNPs in *PtoCesA3* and *PtoCesA4*. These findings were supported by significant correlations in gene expression among these six genes (Fig. 1B).

A similar situation was also observed for hemicellulose, the second most abundant polysaccharide after cellulose, where we identified several haplotypes and eight SNPs from five of the candidate genes (Supplementary Tables S6 and S11). Of these, *PtoUGDP1* and *PtoUXS1* are key enzymes in hemicellulose biosynthesis. Five common candidate genes were shared between wood-property and growth traits (Table 5). *Pto SAHH2* is associated with hemicellulose, tree height, and stem volume. The observation is consistent with SAHH being a cytokinin-binding protein participating in xylogenesis. Interestingly, associations with lignin were found within genes directly or indirectly related to cellulose or xylan biosynthesis, such as *PtoUXS1*, *PtoCesA4*, and *PtoCesA7* (Table 5). This finding was supported by the strong genetic interrelations between the lignin biosynthetic pathway and polysaccharide biosynthesis. Applying biological pathway-based association to examine many more candidate genes involved in the diverse metabolic pathways of tree is needed in future investigation.

The numerous genes that we detected have multiple significant associations across all three trait categories (Tables 1–3, and 5), and the finding was possible to be explained by two hypothesis. On one hand, significant genetic correlations were frequently estimated among multiple related trait fractions or percentages of the whole tree body, especially within each trait category in our present study (Supplementary data S3). On the other hand, this observation may be considered to be an indication of pleiotropy in the broad sense, and the potential pleiotropic loci identified in this study provide novel candidates for explaining phenotypic variation in *Populus*. Pleiotropic loci span multiple functions and may have effects by acting upstream in signalling pathways that affect many traits or by directly targeting multiple genes for regulation, which is consistent with our discoveries that *PtoGA20Ox*, *PtoSAHH1*, and *PtoSAHH2* are all associated with multiple traits (Tables 1–5). More work should be focused on transcription factors/regulators in future.

Table 5. Identification of the common genes significantly related to growth and wood properties in *Populus*.

| Lignin | Holocellulose | Hemicellulose | Fiber length | Fiber width | Microfibre angle | DBH | Tree height | Stem volume |
|--------|---------------|---------------|--------------|-------------|-----------------|-----|-------------|-------------|
| + + + + + + | + + + | + + + + + | + + + + + | + + + + + | + + + + + + | + + + + + | + + + | + + + + + |

| *PtoBGlu1* | *PtoCesA3* | *PtoCesA4* | *PtoCesA7* | *PtoGA20Ox* | *PtoSAHH1* | *PtoSAHH2* | *PtoSUS1* |
|------------|------------|------------|------------|-------------|-------------|-------------|-----------|
| +          | +          | +          | +          | +           | +           | +           | +         |
4.2. Epistasis for growth and wood-property traits
Non-additive interactions between separate mutations (epistasis) can significantly influence the rate and direction of evolutionary change.33 Also, epistasis is an important but often-ignored component of genetic variation in breeding for quantitative traits, due to limited statistical power to detect and estimate epistasis. In our study, we detected and characterized SNP–SNP epistasis associated with the phenotypic traits by using a MDR method (Supplementary Tables S7 and S8). The detection of epistasis hence provides necessary information complementary to that gained through single-locus analysis. We observed that 11 of the 34 unique SNPs identified through the MDR analyses were located in coding regions, with the remaining 23 being located in non-coding regions (Supplementary Table S7), supporting the notion that most associations exhibited pervasive epistatic effects and that it involves both non-coding and coding polymorphisms.36

We identified three non-synonymous loci from three unique genes with significant epistatic effects for DBH and stem volume (Fig. 6 A–D). The similar epistatic pattern for these two traits may be due to a significant positive phenotypic correlation (Supplementary data S3).20 Also, the observation may be supported by previous studies reporting that examinations of interactions between amino acid mutations in the same/shared proteins underlying variation in a measurable phenotype may best reveal the mechanisms of epistasis.37 Without the expectations of an additive null model, the phenotypic effects of allelic substitutions (L → H and H → L) at specific combinatorial patterns of amino acid replacements at these three non-synonymous mutations were highly dependent on genetic background.19 We observed two specific combinatorial patterns: first, in the presence of each amino acid type (H for Ser and L for Thr) in PtoGH9A1_1936, the Asn (L) in Pro-GA20Ox_632 conferred an increased phenotypic value on the Asn (L) background in ProCesA4_1914, and a decrease on the His (H) background in ProCesA4_1914 (Fig. 6B and C). Similarly, the L-type amino acid type in PtoCesA4_1914 conferred a decrease on the Ser (H) background in ProGA20Ox_632 and an increase on the Asn (L) background in ProGA20Ox_632 (Fig. 6B and C). Second, without consideration of amino acid types present in PtoCesA4_1914, the L-type in PtoGH9A1_1936 conferred an increased phenotypic value in the L background in ProGA20Ox_632 and a decrease on the H background in ProGA20Ox_632 (Fig. 6B and C). Similarly, the L-type amino acid in PtoGA20Ox_632 conferred a decrease on the H background in PtoGH9A1_1936 and an increase on the L background in ProGH9A1_1936 (Fig. 6B and C). These findings are examples of sign epistasis38 and suggest the co-selection of epistatic alleles or protein–protein interactions, although the mechanisms for their generation and maintenance are not clear. These observations were supported by the significant positive correlations in gene expression for each pair of these three genes in our tissue-specific transcript profiling (Fig. 1B).

4.3. Future perspectives for MAS breeding
The future of tree improvement will depend on the co-selection of favourable alleles for traits of interest and on the application of combination of haplotypes/genotypes in a manner that effectively overcomes the low levels of LD observed in forest trees.39 Also, genomic selection (GS) is expected to cause a paradigm shift in tree breeding by improving the efficiency and speed of the breeding process.45 Therefore, examining the phenotypic differences among combinations of multiple haplotypes/genotypes for a specific trait (Fig. 5 and Supplementary data S3) provides a foundation for dissecting complex traits through multi-gene or GS in trees. Only 63% of SNPs in haplotypes were identified in the multi-SNP models (Supplementary Tables S6 and S11), consistent with previous observations that haplotype-based tests can detect previously unknown QTL and are thus well supported for unraveling the genetic basis of complex traits.39 Populus tomentosa is distributed across a wide geographical area of northern China20 and thus can provide a potential source of region-specific haplotypes (Table 4). These region-specific signatures may reflect, as well as affect, strong regional differentiation in these traits. Probably these fixed loci are likely key to understanding geographic adaptation and may also be targeted in region-based breeding.

The present work has identified a number of candidate alleles variants that could be exploited to alter key economic traits for industrial applications. However, the direct use of QTL or association information for MAS breeding has not been successful, chiefly because of the difficulty in transferring associations across populations and species of forest trees.45 Future work should focus on replicated sampling for a larger number of broadly representative ecotypes especially across different environments, as well as considering genetic background or genotype × environment interactions.1 Also, joint mapping with association populations and multiple specific biparental crosses is likely to be extremely powerful for improving map resolution and to improve identification of new alleles. The incorporation of more genes (such as gene family members, transcription factors/regulators, transferases, transporters, kinases, and hydrolases) in shared biosynthetic pathways13 would provide a more complete dissection of the effects of genetic variation on growth and lignocellulosic traits. We believe these studies will gradually drive the application of MAS to selection breeding of trees at the early seedling stage.

Acknowledgements
We thank Prof. Ronald R. Sederoff (NC State University, Raleigh, NC, USA) for detailed comments and specific suggestions for improving the manuscript, and also Prof. Rongling Wu and Dr Zhong Wang (Center for Statistical Genetics, Pennsylvania State University, Hershey, PA, USA) for advice on statistical analysis.

Supplementary data
Supplementary data are available at www.dnaresearch.oxfordjournals.org.

Funding
This work was supported by the State Key Basic Research Program of China (No. 2012CB114506), the National ‘863’ Plan Project (No. 2013AA102702), and the 111 Project (No. B13007). Funding to pay the Open Access publication charges for this article was provided by the State Key Basic Research Program of China (No. 2012CB114506), the National ‘863’ Plan Project (No. 2013AA102702), and the 111 Project (No. B13007).

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