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The genome of black cottonwood, *Populus trichocarpa* (Torr. & Gray)

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ABSTRACT

3 We report the draft genome of the black cottonwood tree, *Populus trichocarpa*.
Integration of shotgun sequence assembly with genetic mapping enabled chromosome-
scale reconstruction of the genome. Over 45,000 putative protein-coding genes were
6 identified. Analysis of the assembled genome revealed a whole-genome duplication
event, with approximately 8,000 pairs of duplicated genes from that event surviving in
the *Populus* genome. A second, older duplication event is indistinguishably coincident
9 with the divergence of the *Populus* and *Arabidopsis* lineages. Nucleotide substitution,
tandem gene duplication and gross chromosomal rearrangement appear to proceed
substantially slower in *Populus* relative to *Arabidopsis*. *Populus* has more protein-coding
12 genes than *Arabidopsis*, ranging on average between 1.4-1.6 putative *Populus*
homologs for each *Arabidopsis* gene. However, the relative frequency of protein
domains in the two genomes is similar. Overrepresented exceptions in *Populus* include
15 genes associated with disease resistance, meristem development, metabolite transport
and lignocellulosic wall biosynthesis.

18 **KEYWORDS:** Whole-genome shotgun sequencing, genome-wide duplication, perennial
habit, woody plant, poplar, *Salix*, *Arabidopsis*, angiosperm evolution

Forests cover thirty percent of the earth's terrestrial surface (ca., 3.8 billion hectares), harbor large amounts of biodiversity, and provide humanity with benefits, including clean air and water, lumber, fiber and fuels. Worldwide, one quarter of all industrial feedstocks have their origins in forest-based resources(1). Occurring in extensive wild populations across continents, large and long-lived forest trees have evolved under selective pressures unlike those of annual herbaceous plants. Their growth and development involves extensive secondary growth, coordinated signaling and distribution of water and nutrients over great distances, and strategic storage and re-distribution of metabolites in concordance with inter-annual climatic cycles. The need to survive and thrive in fixed locations over centuries under continually changing physical and biotic stresses also sets them apart from short-lived plants. Many of the features that distinguish trees from other organisms, especially their large sizes and long-generation times, present challenges to the study of the cellular and molecular mechanisms that underlie their unique biology. To enable and facilitate such investigations in a relatively well-studied model tree, we describe here the draft genome of black cottonwood, *Populus trichocarpa* (Torr. & Gray) and its comparison with other sequenced plant genomes.

Populus trichocarpa was selected as the model forest species for genome sequencing not only because of its modest genome size, but also because of its rapid growth, relative ease of experimental manipulation, and range of available genetic tools(2, 3). The genus is phenotypically diverse and interspecific hybrids facilitate the genetic mapping of economically important traits related to growth rate, stature, wood properties and paper quality. Dozens of quantitative trait loci (QTL) are already mapped(4) and methods of genetic transformation have been developed(5). Under appropriate conditions, *Populus* can reach reproductive maturity in as few as 4-6 years, permitting selective breeding for large-scale sustainable plantation forestry. Finally, rapid growth of trees coupled with thermochemical or biochemical conversion of the lignocellulosic portion of the plant has the potential to provide a renewable energy resource with a concomitant reduction of greenhouse gases(6-8).

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SEQUENCING and ASSEMBLY

A single female genotype, 'Nisqually-1', was selected and used in a whole-genome shotgun sequence and assembly strategy(9). Approximately 7.6 million end-reads representing 4.2 billion high-quality (i.e., Q20 or higher) base pairs were

assembled into 2,447 major scaffolds containing an estimated 410 Mb of genomic DNA (SOM T1 & T2). On the basis of the depth of coverage of major scaffolds (~7.5X), and the total amount of non-organellar shotgun sequence that was generated, the *Populus* genome size was estimated to be 485±10 Mb, in rough agreement with previous cytogenetic estimates of approximately 550 Mb(10). The near completeness of the shotgun assembly in protein-coding regions is supported by the identification of more than 95% of known *Populus* cDNA in the assembly (see Gene Content section below).

The ~75 Mb of unassembled genomic sequence is consistent with cytogenetic evidence for ~30% of the genomic being heterochromatic(9). The amount of euchromatin contained within the *Populus* genome was estimated in parallel by subtraction on the basis of direct measurements of DAPI-stained prophase and metaphase chromosomes (SOM F4). On average, 69.5±0.3% of the genome consisted of euchromatin, with a significantly lower proportion of euchromatin in linkage group I (66.4±1.1%), compared to the other 18 chromosomes (69.7±0.03%, $p \leq 0.05$). In contrast, *Arabidopsis* chromosomes contain roughly 93% euchromatin(11). The unassembled shotgun sequences were derived from variants of organellar DNA, including recent nuclear translocations, highly repetitive genomic DNA, haplotypic segments that were redundant with short subsegments of the major scaffolds (separated due to extensive sequence polymorphism, *i.e.*, allelic variants), and contaminants of the template DNA, including endophytic microbes inhabiting the leaf and root tissues used for template preparation(12) (SOM F1 & T3). The end reads corresponding to chloroplast (SOM F5) and mitochondrial genomes were assembled into circular genomes of 157 and 803 kb, respectively(9).

We anchored the 410 Mb of assembled scaffolds to a sequence-tagged genetic map (SOM F3). In total, 356 microsatellite markers were used to assign 155 scaffolds (335 Mb of sequence) to the 19 *P. trichocarpa* chromosome-scale linkage groups (LG) (13). The vast majority (91%) of the mapped microsatellite markers were colinear with the sequence assembly. At the extremes, the smallest chromosome, LGIX (79 cM), is covered by two scaffolds containing 12.5 Mb of assembled sequence; whereas the largest chromosome, LGI (265 cM), contains 21 scaffolds representing 35.5 Mb (SOM F3). We also generated a physical map based on BAC fingerprint contigs using a Nisqually-1 BAC library representing an estimated 9.5-fold genome coverage (SOM F2). Paired BAC-end sequences from most of the physical map were linked to the large-scale assembly, permitting 2,460 of the physical map contigs to be positioned on the genome

assembly. Combining the genetic and physical map, nearly 385 Mb of the 410 Mb of assembled sequence is placed on a linkage group.

3 Unlike Arabidopsis, where predominantly self-fertilizing ecotypes maintain low levels of allelic polymorphism, *Populus* species are predominantly dioecious, which results in obligate outcrossing. This compulsory outcrossing, along with wind pollination
6 and wind dispersed plumose seeds, results in high levels of gene flow and high levels of heterozygosity (*i.e.*, within-individual genetic polymorphisms). Within the heterozygous Nisqually-1 genome, we identified 1,241,251 single nucleotide polymorphisms (SNP) or small indel polymorphisms for an overall rate of approximately 2.6 polymorphisms per kb. Of these polymorphisms the overwhelming majority (83%) occur in non-coding portions of the genome (Table 1). Short insertion/deletion polymorphisms (indels) and
9 SNP polymorphisms within exons resulted in some frameshifts and nonsense stop codons within predicted exons, respectively, suggesting that null alleles of these genes exist in one of the haplotypes. Some of the polymorphisms may be artifacts from the assembly process; though these errors were minimized by using stringent criteria for
12 SNP identification(9).

18 GENE ANNOTATION

We tentatively identified a first-draft reference set of 45,555 protein-coding gene loci in the *Populus* nuclear genome (www.jgi.doe.gov/poplar) using a variety of *ab initio*,
21 homology-based and EST-based methods(14-17) (SOM T5). Similarly, 101 and 52 genes were annotated in the chloroplast and mitochondrial genomes, respectively⁹. To aid the annotation process, 4,664 full-length sequences, from full-length enriched cDNA
24 libraries from Nisqually-1, were generated and used in training the gene-calling algorithms. Prior to gene prediction, repetitive sequences were characterized (SOM F15 & T14) and masked; additional putative transposable elements were identified and
27 subsequently removed from the reference gene set(9). Given the current draft nature of the genome, we expect that the gene set in *Populus* will continue to be refined.

Approximately 89% of the predicted gene models had homology ($E\text{-value} \leq 1e^{-8}$)
30 to the non-redundant (NR) set of proteins from NCBI, including 60% with extensive homology over 75% of both model and NR protein lengths. Nearly 12% (5,248) of the predicted *Populus* genes had no detectable similarity to Arabidopsis genes ($E\text{-value} \leq 1e^{-3}$); conversely, in the more refined Arabidopsis set, only 9% (2,321) of the predicted
33 genes had no similarity to the *Populus* reference set. Of the 5,248 *Populus* genes

without *Arabidopsis* similarity, 1,883 have expression evidence from the manually-curated *Populus* EST dataset, and of these, 274 have no hits ($E\text{-value} \geq 1e^{-3}$) to the NR database(9). Whole-genome oligonucleotide microarray analysis provided evidence of tissue-based expression for 53% for the reference gene models (Fig. 1). In addition, signal was detected from 20% of genes that were initially annotated and excluded from the reference set, suggesting that as many as 4,000 additional genes (or gene fragments) may be present. Within the reference gene set, 13,019 pairs of orthologs were identified between genes in *Populus* and *Arabidopsis* using the best bi-directional BLAST hits, with average mutual coverage of these alignments equal to 93%; 11,654 pairs of orthologs had coverage greater than 90% of gene lengths, with only 156 genes with less than 50% coverage. As of June 1, 2006, ~10% (4,378) gene models have been manually validated and curated.

GENOME ORGANIZATION

15 Genome Duplication in the Salicaceae

Populus and *Arabidopsis* lineages diverged ca. 100-120 Mya. Analysis of the *Populus* genome provided evidence of a more recent duplication event that impacted roughly 92% of the *Populus* genome. Nearly 8,000 pairs of paralogous genes of similar age (excluding tandem or local duplications) were identified (Fig. 2). The relative age of the duplicate genes was estimated by the accumulated nucleotide divergence at four-fold synonymous third-codon transversion position (4DTV) values. A sharp peak in 4DTV values, corrected for multiple substitutions, representing a burst of gene duplication, is evident at 0.0916 ± 0.0004 (Fig. 3A). Comparison of 1,825 *Populus* and *Salix* orthologous genes derived from *Salix* EST suggests that both genera share this whole-genome duplication event (Fig. 3B). Moreover, the parallel karyotypes and collinear genetic maps(18) of *Salix* and *Populus* also support the conclusion that both lineages share the same large-scale genome history.

If we naively calibrate the molecular clock using synonymous rates observed in the Brassicaceae(19) or derived from the *Arabidopsis*-*Oryza* divergence(20), we would conclude that the genome duplication in *Populus* is very recent (8-13 Mya as reported by 19). Yet the fossil record shows that the *Populus* and *Salix* lineages diverged 60-65 million years ago(22-25). Thus the molecular clock in *Populus* must be ticking at only one sixth the estimated rate for *Arabidopsis* (i.e., 8-13 Mya/60-65 Mya). Qualitatively similar slowing of the molecular clock is found in the *Populus* chloroplast and

mitochondrial genomes(9). As a long-lived vegetatively propagated species *Populus* has the potential to successfully contribute gametes to multiple generations. A single
3 *Populus* genotype can persist as a clone on the landscape for millennia(26), and we propose that recurrent contributions of “ancient gametes” from very old individuals could account for the dramatically reduced rate of sequence evolution. As result of the slowing
6 of the molecular clock, the *Populus* genome most likely resembles the ancestral eurousid genome.

To test if the burst of gene creation 60-65 Mya was due to a single whole-
9 genome event or independent but near-synchronous gene duplication events we used a variant of the algorithm of Hokamp *et al.* (27) to identify segments of conserved synteny within the *Populus* genome. The longest conserved syntenic block from the 4DTV ~0.09
12 epoch spanned 765 pairs of paralogous genes. In total, 32,577 genes were contained within syntenic blocks from the salicoid epoch; half of these genes were contained in segments longer than 142 paralogous pairs. The same algorithm, when applied to
15 randomly shuffled genes, typically yields duplicate blocks with fewer than 8-9 genes, indicating that the *Populus* gene duplications occurred as a single genome-wide event. Through the remainder of this paper this duplication event will be referred to as the
18 “salicoid” duplication event.

Nearly every mapped segment of the *Populus* genome had a parallel
“paralogous” segment elsewhere in the genome as a result of the salicoid event (Fig. 2).
21 The “pinwheel” patterns can be understood as a whole-genome duplication followed by a series of reciprocal tandem terminal fusions between two separate sets of four chromosomes each; the first involving LGII, V, VII and XIV and the second involving LGI,
24 XI, IV and IX. In addition, several chromosomes appear to have experienced minor reorganizational exchanges. Furthermore, LGI appears to be the result of multiple rearrangements involving three major tandem fusions. These results suggest that the
27 progenitor of *Populus* had a base chromosome number of 10 which, following the whole-genome duplication event, experienced a genome-wide reorganization and diploidization of the duplicated chromosomes into four pairs of complete paralogous chromosomes
30 (LGVI, VIII, X, XII, XIII, XV, XVI, XVIII & XIX), two sets of four chromosomes each containing a terminal translocation (LGI, II, IV, V, VII, IX & XI) and one chromosome containing three terminally joined chromosomes (LGIII with I or XVII with VII). The
33 colinearity of genetic maps among multiple *Populus* species suggests that the genome reorganization must have occurred prior to the evolution of the modern taxa of *Populus*.

Genome Duplication in a Common Ancestor of *Populus* and *Arabidopsis*

3 The distribution of 4DTV values for paralogous pairs of genes also shows that a large
fraction of the *Populus* genome falls in a set of duplicated segments anchored by gene
6 pairs with 4DTV at 0.364 ± 0.001 , representing the residue of a more ancient, large-scale,
apparently synchronous duplication event (Fig 3A). This relatively older duplication event
covers approximately 59% of the *Populus* genome with 16% of genes in these segments
9 present in two copies. Since this duplication preceded and is therefore superimposed
upon the salicoid event, each genomic region is potentially covered by four such
segments. Similarly, the *Arabidopsis* genome experienced an older “beta” duplication
that preceded the Brassicaceae-specific “alpha” event(28-32).

12 We next asked if the *Arabidopsis* “beta” (30, 32) and *Populus* 4DTV~0.36
duplication events were (i) independent genome-wide duplications that occurred after
the split from the last common eucosid ancestor (H_1) or (ii) a single shared duplication
15 event that occurred in an ancestral lineage (*i.e.*, prior to the eucosid I/II divergence) (H_2).
These two hypotheses have very different implications for the interpretation of homology
between *Populus* and *Arabidopsis*. Under H_1 each genomic segment in one species is
18 homologous to four segments in the other, while under H_2 each segment is homologous
to only two segments in the other species. These hypotheses were tested by comparing
the relative distances between gene pairs sampled within and between *Populus* and
21 *Arabidopsis*. H_2 was generally supported(9), but, we could not reject H_1 . We can only
conclude that the *Populus* genome duplication occurred very close to the time of
divergence of the Eucosid I and II lineages(9), with slight support for a shared
24 duplication. This coincident timing raises the possibility of a causal link between this
duplication and rapid diversification early in eucosid (and perhaps core eudicot) history.
Through the remainder of this paper this older *Populus/Arabidopsis* duplication event will
27 be referred to as the “eucosid” duplication event. We note that the salicoid duplication
occurred independently of the eucosid duplication observed in the *Arabidopsis* genome.

30 GENE CONTENT

Although *Populus* has substantially more protein-coding genes than *Arabidopsis*,
the relative frequency of domains represented in protein databases (Prints, Prosite,
33 Pfam, ProDom & SMART) in the two genomes is similar(9). However, the most common
domains occur in *Populus* in a 1.4-1.8 to one ratio compared with *Arabidopsis*.

Noteworthy outliers in *Populus* include genes and gene domains associated with disease and insect resistance (e.g., leucine rich repeats, 1,271 vs. 527; NB-ARC domain, 302 vs. 141; thaumatin, 55 vs. 24, *Populus* vs. *Arabidopsis*, respectively), meristem development (e.g., NAC transcription factors, 157 vs. 100, respectively) and metabolite/nutrient transport (e.g., oligopeptide transporter of the POT and OPT families, 129 vs. 61; potassium transporter, 30 vs. 13, respectively).

Some domains were underrepresented in *Populus* compared to *Arabidopsis*. For example, the F-box domain was twice as prevalent in *Arabidopsis* as in *Populus* (624 vs. 303, respectively). The F-box domain is involved in diverse and complex interactions involving protein degradation via the ubiquitin-26S proteasome pathway(33). Many of the ubiquitin-associated domains are underrepresented in *Populus* compared to *Arabidopsis* (e.g., *Ulp1* protease family, C-terminal catalytic domain, 10 vs. 63, respectively). Moreover, the RING finger domains are nearly equally present in both genomes (503 vs. 407, respectively), suggesting that protein degradation pathways in the two organisms are metabolically divergent.

The Common Eurosid Gene Set

The *Populus* and *Arabidopsis* gene sets were compared to infer the conserved gene complement of their common eurosid ancestor, integrating information from nucleotide divergence, synteny and mutual best BLAST-hit analysis(9). The ancestral eurosid genome contained at least 11,666 protein-coding genes, along with an undetermined number that were either lost in one or both of the lineages or whose homology could not be detected. These ancestral genes were the progenitors of gene families of typically 1-4 descendents in each of the complete plant genomes and account for 28,257 *Populus* and 17,521 *Arabidopsis* genes. Gene family lists are accessible at: www.phytozome.net. The gene predictions in these two genomes that could not be accounted for in the eurosid clusters were often fragmentary or could not be confidently assigned orthology and may include novel or rapidly evolving genes in the *Populus* and/or *Arabidopsis* lineages, as well as poorly predicted genes.

Non-Coding RNAs

Based on a series of publicly available RNA detection algorithms(34), including tRNAScan-SE, INFERNAL and snoScan, we identified 817 putative transfer RNAs (tRNA), 22 U1, 26 U2, 6 U4, 23 U5, 11 U6 spliceosomal RNAs (snRNA), 339 putative

C/D small nucleolar RNAs (snoRNA) and 88 predicted H/ACA snoRNAs in the *Populus* assembly. All 57 possible anti-codon tRNA were found. One selenocysteine tRNA was detected and two possible suppressor tRNA (anticodons which bind stop codons) were also discovered. *Populus* has nearly 1.3 times as many tRNA genes as Arabidopsis. In contrast to Arabidopsis (SOM F7A), the copy number of tRNA in *Populus* was significantly and positively correlated with amino acid occurrence in predicted gene models (SOM F7B). *Populus* has a 1.3 to 1.0 ratio in the number of snRNA compared with Arabidopsis, yet U1, U2 and U5 are overrepresented in *Populus* while U4 is underrepresented. Furthermore, U14 was not detected in Arabidopsis. The snRNA and snoRNA have not been experimentally verified in *Populus*.

There are 169 identified microRNA (miRNA) genes representing 21 families in *Populus* (SOM T7). In Arabidopsis, these 21 families contain 91 miRNA genes, representing a 1.9X expansion in *Populus*, primarily in miR169 and miR159/319. All 21 miRNA families have regulatory targets that appear to be conserved among Arabidopsis and *Populus* (SOM T8). Like the miRNA genes themselves, the number of predicted targets for these miRNA is expanded in *Populus* (147) compared to Arabidopsis (89). Similarly, the genes that mediate RNAi are also overrepresented in *Populus* (21) compared to Arabidopsis (11) (e.g., AGO1 class, 7 vs. 3; RNA helicase 2 vs.1; HEN, 2 vs.1; HYL1-like (dsRNA binding proteins) 9 vs. 5, respectively).

Tandem Duplications

In *Populus* there were 1,518 tandemly duplicated arrays of two or more genes based on a Smith-Waterman alignment $E\text{-value} \leq e^{-25}$ and a 100 kb window. The total number of genes in such arrays was 4,839 and the total length of tandemly duplicated segments in *Populus* was 47.9 Mb or 15.6% of the genome (SOM F8). By the same criteria, there are 1,366 tandemly duplicated segments in Arabidopsis, covering 32.4 Mb or 27% of the genome. By far the most common number of genes within a single array was two, with 958 such arrays in *Populus* and 805 in Arabidopsis. Arabidopsis had a larger number of arrays containing six or more genes than did *Populus*. Tandem duplications thus appear to be relatively more common in Arabidopsis than in *Populus*. This may in part be due to difficulties in assembling tandem repeats from a whole-genome shotgun sequencing approach, particularly when tandemly-duplicated genes are highly conserved. Alternatively, the *Populus* genome may be undergoing rearrangements at a slower rate than the Arabidopsis genome, which is consistent with

our observations of reduced chromosomal rearrangements and slower nucleotide substitution rates in *Populus*.

3 In some cases, genes were highly duplicated in both species, with some tandem duplications predating the *Populus*-*Arabidopsis* split(9). The largest number of tandem repeats in *Populus* in a single array was 24 and contained genes with high homology to
6 S-locus specific glycoproteins. Genes of this class also occur as tandem repeats in *Arabidopsis*, with the largest segments containing 14 tandem duplicates on chromosome 1. One of the InterPro domains in this protein, [IPR008271](#), a serine/threonine protein
9 kinase active site, was the most frequent domain in tandemly repeated genes in both species (SOM F8). Other common domains in both species were the leucine-rich repeat ([IPR007090](#), primarily from tandem repeats of disease resistance genes), the
12 pentatricopeptide repeat RNA-binding proteins ([IPR002885](#)), and the UDP-glucuronosyl/UDP-glucosyltransferase domain ([IPR002213](#)) (SOM T9).

In contrast, some genes were highly expanded in tandem duplicates in one
15 genome and not in the other (SOM F8). For example, one of the most frequent classes of tandemly duplicated genes in *Arabidopsis* was F-box genes, with a total of 342 involved in tandem duplications, of which the largest segment contained 24 F-box
18 genes.. *Populus* contains only 37 F-box genes in tandem duplications, with the largest segment containing only three genes.

21 **POST-DUPLICATION GENE FATE**

Functional expression divergence

In *Populus*, 20 of the 66 salicoid-event duplicate gene pairs contained in 19
24 *Populus* EST libraries (2.3% of the total) showed differential expression(9) (displayed significant deviation in EST frequencies per library, e.g., Fig. 4). Eleven of 18 eurosid-event duplicate gene pairs (2.7% of the total) also displayed significant deviation in EST
27 frequencies per library. Many of the duplicate gene pairs that displayed significant overrepresentation in one or more of the 19 sampled libraries were involved in protein-protein interactions (e.g., annexin) or protein folding (e.g., cyclophilins). In the eurosid
30 set, there was a greater divergence in the best BLAST hit among pairwise sets of genes. These results support functional expression divergence among some duplicated gene pairs in *Populus*.

33 As a further test for variation in gene expression among duplicated genes we examined whole-genome oligonucleotide microarray data containing the 45,555

promoted genes(9). There was significantly lower differential expression in the salicoid duplicated pairs of genes (mean: 5%) relative to eurosid duplications (mean: 11%), again suggesting that differential expression patterns for retained paralogous gene pairs is an ongoing process that has had more time to occur in eurosid pairs (Fig. 5). This difference could also be due to absolute expression level, which may vary systematically between the two duplication events. Moreover, differential expression was more evident in the wood-forming organs. Almost 14% and 13% (2,632 pairs of genes) of eurosid duplicated genes in the nodes and internodes, respectively, displayed differential expression compared to 8% or lower in roots and young leaves (Fig. 5).

Single Nucleotide Polymorphisms

Populus is a highly polymorphic taxon and substantial numbers of SNP are present even within a single individual (Table 1). The ratio of nonsynonymous to synonymous substitution rate ($\omega=dN/dS$) was calculated as an index of selective constraints for alleles of individual genes(9). The overall average dN across all genes was 0.0014, while dS value was 0.0035, for a total ω of 0.40, suggesting that the majority of coding regions in the *Populus* genome are subject to purifying selection. There was a significant, negative correlation between ω and the 4DTV distance to the most closely related paralog ($r=-0.034$, $p=0.028$), which is consistent with the expectation of higher levels of nonsynonymous polymorphism in recently duplicated genes due to functional redundancy(20, 35). Similarly, genes with recent tandem duplicates (4DTV \leq 0.2) had significantly higher ω than genes with no recent tandem duplicates (Wilcoxon Rank Sum $Z=8.65$, $p\leq 0.0001$) (SOM T10).

The results for tandemly duplicated genes were consistent with expectations for accelerated evolution of duplicated genes(20). However, this expectation was not upheld for paralogous pairs of genes from the whole-genome duplication events. Relative rates of nonsynonymous substitution were actually lower for genes with paralogs from the salicoid and eurosid whole-genome duplication events than for genes with no paralogs (SOM T11). One possible explanation for this discrepancy is that the apparent single-copy genes have a corresponding overrepresentation of rapidly-evolving pseudogenes. However, this does not appear to be the case, as demonstrated by an analysis of gene size, synonymous substitution rate and minimum genetic distance to the closest paralog as covariates in an analysis of variance with ω as the response variable (SOM T11). Therefore, genes with no paralogs from the salicoid and eurosid duplication events seem

to be under lower selective constraints and purifying selection is apparently stronger for genes with paralogs retained from the whole-genome duplications. Chapman *et al.* (36) have recently proposed the concept of functional buffering to account for similar reduction in detected mutations in paralogs from whole-genome duplications in Arabidopsis and *Oryza*. The vegetative propagation habit of *Populus* may also favor the conservation of nucleotide sequences among duplicated genes, in that complementation among duplicate pairs of genes would minimize loss of gene function associated with the accumulation of deleterious somatic mutations.

Gene Family Evolution

Lignocellulosic Wall Formation Among the processes unique to tree biology, one of the most obvious is the yearly development of secondary xylem from the vascular cambium. *Populus* orthologs of the approximately 20 Arabidopsis genes/gene families involved in or associated with cellulose biosynthesis were identified. The *Populus* genome has 93 cellulose synthesis-related genes vs. 78 in Arabidopsis. Arabidopsis genome encodes 10 *CesA* genes belonging to six classes known to participate in cellulose microfibril biosynthesis(37). *Populus* has 18 *CesA* genes(38), including duplicate copies of *CesA7* and *CesA8* homologs. *Populus* homologs of Arabidopsis *CesA4*, *CesA7* and *CesA8* are coexpressed during xylem development and tension wood formation(39). Furthermore, one pair of *CesA* genes appears unique to *Populus*, with no homologs found in Arabidopsis(40). Many other types of genes associated with cellulose biosynthesis, *e.g.*, *KOR*, *SuSY*, *COBRA* and *FRA2*, occur in duplicate pairs in *Populus* relative to single-copy Arabidopsis genes(39). For example *COBRA*, a regulator of cellulose biogenesis(41), is a single-copy gene in Arabidopsis yet in *Populus* there are four copies.

The repertoire of acknowledged hemicellulose biosynthetic genes in *Populus* is generally similar to that in Arabidopsis. However, *Populus* has more genes encoding α -L-fucosidases and fewer genes encoding α -L-fucosyltransferases than does Arabidopsis, which is consistent with the lower xyloglucan fucose content(42) in *Populus* relative to Arabidopsis.

Lignin, the second most abundant secondary cell wall polymer after cellulose, is a complex polymer of monolignols (hydroxycinnamyl alcohols) that encrusts and interacts with the cellulose/hemicellulose matrix of the secondary cell wall(43). The full set of 34 *Populus* phenylpropanoid and lignin biosynthetic genes (SOM T13) were

identified by sequence alignment to the known Arabidopsis phenylpropanoid and lignin genes(44, 45). The size of *Populus* gene families encoding these enzymes is generally larger than in Arabidopsis (34 vs. 18, respectively). The only exception is *CAD* (cinnamyl alcohol dehydrogenase), which is encoded by a single gene in *Populus* and two genes in Arabidopsis (Fig. 6C); *CAD* is also encoded by only a single gene in *Pinus taeda*(46, 47). Two lignin-related *Populus C4H* genes are strongly co-expressed in tissues related to wood formation while the three *Populus C3H* genes show reciprocally exclusive expression patterns(48).

Secondary Metabolism *Populus* produce a broad array of non-structural, carbon-rich secondary metabolites that exhibit wide variation in abundance, stress inducibility, and effects on tree growth and host-pest interactions(49-53). Shikimate-phenylpropanoid derived phenolic esters, phenolic glycosides and condensed tannins and their flavonoid precursors comprise the largest classes of these metabolites. Phenolic glycosides and condensed tannins alone can constitute up to 35% leaf dry weight and are abundant in buds, bark and roots of *Populus*(50, 54, 55).

The flavonoid biosynthetic genes are well annotated in Arabidopsis(56) and almost all (with the exception of flavonol synthase) are encoded by single-copy genes. In contrast, all but three such enzymes (chalcone isomerase, flavonoid 3'-hydroxylase and flavanone 3-hydroxylase) are encoded by multiple genes in *Populus*(53). For example, the chalcone synthase (*CHS*), controlling the committed step to flavonoid biosynthesis, has expanded to at least six genes in *Populus*. In addition, *Populus* contains two genes each for flavone synthase II (*CYP98B*) and flavonoid 3', 5'-hydroxylase (*CYP75A12* and *CYP75A13*) which are absent in Arabidopsis. Furthermore, three *Populus* genes encode leucoanthocyanidin reductase, required for the synthesis of condensed tannin precursor 2,3-*trans*-flavan-3-ols, a stereochemical configuration also lacking in Arabidopsis(57). In contrast to the 32 terpenoid synthases (*TPS*) genes of secondary metabolism identified in the Arabidopsis genome(58), the *Populus* genome contains at least 47 *TPS* genes, suggesting a wide-ranging capacity for the formation of terpenoid secondary metabolites.

A number of phenylpropanoid-like enzymes have been annotated in the Arabidopsis genome(44, 45, 59-61). One example is the family encoding cinnamyl alcohol dehydrogenase (*CAD*). In addition to the single *Populus CAD* gene involved in lignin biosynthesis, several other clades of *CAD-like* (*CADL*) genes are present, most of

which fall within larger sub-families containing enzymes related to multifunctional alcohol dehydrogenases (Fig. 6). This comparative analysis makes it clear that there has been selective expansion and retention of *Populus CADL* gene families. For example, *Populus* contains seven *CADL* genes (*PoptrCADL1-7*; Fig. 6C) encoding enzymes related to the Arabidopsis *BAD1* and *BAD2* enzymes with apparent benzyl alcohol dehydrogenase activities(62). *BAD1* and *BAD2* are known to be pathogen-inducible, suggesting that this group of *Populus* genes, including *Populus SAD* gene, previously characterized as encoding a sinapaldehyde-specific CAD enzyme(63), may be involved in chemical defense.

Disease Resistance The likelihood that a perennial plant will encounter a pathogen or herbivore before reproduction is near unity. The long-generation intervals for trees make it difficult for such plants to match the evolutionary rates of a microbial or insect pest. Aside from the formation of thickened cell walls and the synthesis of secondary metabolites that constitute a first line of defense against microbial and insect pests, plants use a variety of disease-resistance (*R*) genes.

The largest class of characterized *R* genes encodes intracellular proteins that contain a nucleotide-binding site (NBS) and carboxy-terminal leucine-rich-repeats (LRR) (64). The NBS-coding *R* gene family is one of the largest in *Populus*, with 399 members, approximately 2-fold higher than in Arabidopsis. The NBS family can be divided into multiple subfamilies with distinct domain organizations, including 64 TIR-NBS-LRR genes, 10 truncated TIR-NBS that lack an LRR, 233 non-TIR-NBS-LRR genes and 17 unusual TIR-NBS-containing genes not identified previously in Arabidopsis (TNLT, TNLN or TCNL) (Table 2). Five gene models coding for TNL proteins contained a predicted N-terminal nuclear localization signal (NLS) (65). The number of non-TIR-NBS-LRR genes in *Populus* is also much higher than that in Arabidopsis (209 vs. 57, respectively). Intriguingly, 40 non-TIR-NBS genes, not found in Arabidopsis, carry an N-terminal BED DNA-binding zinc finger domain that was also found in the *Oryza Xa1* gene. These findings suggest that domain cooption occurred in *Populus*. Most NBS-LRR (ca. 65%) in *Populus* occur as singletons or in tandem duplications and the distribution of pairwise genetic distances among these genes suggests a recent expansion of this family. That is, only 10% of the NBS-LRR genes are associated with the eurosid and salicoid duplication events, compared with 55% of the extracellular LRR receptor-like kinase genes, for example (SOM F10).

Several conserved signaling components such as *RAR1*, *EDS1*, *PAD4* and *NPR1*, known to be recruited by *R* genes, also contain multiple homologs in *Populus*.
3 For example, two copies of the *PAD4* gene, which functions upstream of salicylic acid (SA) accumulation, and five copies of the *NPR1*, an important regulator of responses downstream of SA, are found in *Populus*. Nearly all genes known to control disease
6 resistance signaling in Arabidopsis have putative orthologs in *Populus*. *Populus* has a larger number of β -1, 3-glucanase and chitinase genes than Arabidopsis (131 vs. 73, respectively). In summary, the structural and genetic diversity that exists among *R* genes
9 and their signaling components in *Populus* is remarkable and suggests that unlike the rest of the genome, contemporary diversifying selection has played an important role in the evolution of disease resistance genes in *Populus*. Such diversification suggests that
12 enhanced ability to detect and respond to biotic challenges via *R* gene-mediated signaling may be critical over a decades-long life span of this genus.

Membrane Transporters Attributes of *Populus* biology such as massive interannual,
15 seasonal and diurnal metabolic shifts and re-deployment of carbon and nitrogen may require an elaborate array of transporters. Investigation of gene families coding for transporter proteins (<http://plantst.genomics.purdue.edu/>) in the *Populus* genome
18 revealed a general expansion relative to Arabidopsis (1,722 vs. 959, *Populus* vs. Arabidopsis, respectively) (SOM T12). Five gene families, coding for ATP-binding cassette proteins (ABC transporters, 226 gene models), major facilitator superfamily
21 proteins (MFS, 187 genes), drug/metabolite transporters (DMT, 108 genes), amino acid/auxin permeases (AAP, 95 genes) and proton-dependent oligopeptide transporters (POT, 90 genes), accounted for more than 40% of the total number of
24 transporter gene models (SOM F14). Some large families such as those encoding POT (4.3X relative to Arabidopsis), glutamate-gated ion channels (3.7X), potassium uptake permeases (2.3X) and ABC transporters (1.9X) are expanded in *Populus*. A novel
27 subfamily of five putative aquaporins, lacking in the Arabidopsis, was identified. *Populus* also harbors seven transmembrane receptor genes only found so far in fungi, and two genes, identified as mycorrhizal-specific phosphate transporters, confirming that the
30 mycorrhizal symbiosis may have a significant impact on the mineral nutrition of this long-lived species. This expanded inventory of transporters could conceivably play a role in adaptation to nutrient-limited forest soils, long-distance transport and storage of water

and metabolites, secretion and movement of secondary metabolites, and/or mediation of resistance to pathogen-produced secondary metabolites or other toxic compounds.

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Phytohormones Both physiological and molecular studies have indicated the importance of hormonal regulation underlying plant development. Auxin, gibberellin, cytokinin and ethylene responses are of particular interest in tree biology.

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Many auxin responses(66-71) are controlled by auxin response factor (*ARF*) transcription factors, which work together with cognate AUX/IAA repressor proteins to regulate auxin-responsive target genes(72, 73). A phylogenetic analysis using the known and predicted *ARF* protein sequences showed that *Populus* and Arabidopsis *ARF* gene families have expanded independently since they diverged from their common ancestor.

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Six duplicate *ARF* genes in *Populus* encode paralogs of *ARF* genes that are single-copy Arabidopsis genes, including *ARF5* (*MONOPTEROS*), an important gene required for auxin-mediated signal transduction and xylem development. Furthermore, five Arabidopsis *ARF* genes have four or more predicted *Populus* *ARF* gene paralogs. In contrast to *ARF* genes, *Populus* does not contain a dramatically expanded repertoire of *AUX/IAA* genes relative to Arabidopsis (35 vs. 29, respectively) (74). Interestingly, there is a group of four Arabidopsis *AUX/IAA* genes with no apparent *Populus* orthologs, suggesting Arabidopsis-specific functions.

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Gibberellins are thought to regulate multiple processes during wood and root development, including xylem fiber length(75). Among all gibberellin biosynthesis and signaling genes, the *Populus* GA20-oxidase gene family is the only family with approximately 2-fold increase in gene number relative to Arabidopsis, indicating that most of the duplicated genes that arose from the salicoid duplication event have been lost. GA20-oxidase appears to control flux in the biosynthetic pathway leading to the bioactive gibberellins GA₁ and GA₄. The higher complement of GA20-oxidase genes may have biological significance in *Populus* with respect to secondary xylem and fiber cell development.

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Cytokinins are thought to control the identity and proliferation of cell types relevant for wood formation as well as general cell division(67). The total number of members in gene families encoding cytokinin homeostasis related isopentenyl transferases (*IPT*) and cytokinin oxidases is roughly similar between *Populus* and Arabidopsis, although there appears to be lineage-specific expansion of *IPT* subfamilies. The cytokinin signal transduction pathway represents a two-component phosphorelay

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system, where a two-component hybrid receptor initiates a phosphotransfer via histidine-containing phosphotransmitters (*HPT*) to phospho-accepting response regulators (*RR*).

3 One family of genes, encoding the two-component receptors (*i.e.*, *CKI1*), is notably expanded in *Populus* (4 vs. 1, respectively) (**76**). Gene families coding for recently identified pseudo *HPT* and atypical *RR* are overrepresented in *Populus* relative to
6 *Arabidopsis* (2.5X and 4.0X, respectively). Both of these gene families have been implicated in the negative regulation of cytokinin signaling(**67, 77**), which is consistent with the idea of increased complexity in regulation of cytokinin signal transduction in
9 *Populus*.

Populus and *Arabidopsis* genomes contain almost identical number of genes for the three enzymes of ethylene biosynthesis, whereas the number of genes for proteins
12 involved in ethylene perception and signaling is higher in *Populus*. For example, *Populus* has seven predicted genes for ethylene receptor proteins and *Arabidopsis* has five; the constitutive triple response (*CTR1*) kinase that acts just downstream of the receptor is
15 encoded by four genes in *Populus* and only one in *Arabidopsis*(**78**). The number of ethylene-responsive element binding factor (*ERF*) proteins (a subfamily of *AP2/ERF* family) is higher in *Populus* than in *Arabidopsis* (172 vs. 122, respectively). The
18 increased variation in the number of *ERF* transcription factors may be involved in the ethylene-dependent processes specific to trees, such as tension wood formation(**68**) and the establishment of dormancy(**71**).

21

CONCLUDING REMARKS

Our initial analyses provide a flavor of the opportunities for comparative plant
24 genomics made possible by the generation of the *Populus* genome sequence. A complex history of whole-genome duplications, chromosomal rearrangements and tandem duplications has shaped the genome that we observe today. The differences in
27 gene content between *Populus* and *Arabidopsis* have provided some tantalizing insights into the possible molecular bases of their strongly contrasting life histories, though it is important to note that factors unrelated to gene content (*e.g.*, regulatory elements,
30 miRNA, post-translational modification, or epigenetic modifications) may ultimately be of equal or greater importance. With the sequence of *Populus*, researchers can now go beyond what could be learned from *Arabidopsis* alone to explore hypotheses to linking
33 genome sequence features to wood development, nutrient and water movement, crown development, and disease resistance in perennial plants. The availability of the *Populus*

genome sequence will enable continuing comparative genomics studies among species that will shed new light on genome reorganization and gene family evolution.

3 Furthermore, the genetics and population biology of *Populus* make it an immense source of allelic variation. Because *Populus* is an obligate outcrossing species, recessive alleles tend to be maintained in a heterozygous state. Informatics tools enabled by the

6 sequence, assembly and annotation of the *Populus* genome will facilitate the characterization of allelic variation in wild *Populus* populations adapted to a wide range of environmental conditions and gradients over large portions of the northern

9 hemisphere. Such variants represent a rich reservoir of molecular resources useful in biotechnological applications, development of alternative energy sources, and mitigation of anthropogenic environmental problems. Finally, the keystone role of *Populus* in many

12 ecosystems provides the first opportunity for the application of genomics approaches to questions with ecosystem-scale implications(79, 80).

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