Contrasting patterns of evolution following whole genome versus tandem duplication events in *Populus*

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Comparative analysis of multiple angiosperm genomes has implicated gene duplication in the expansion and diversification of many gene families. However, empirical data and theory suggest that whole-genome and small-scale duplication events differ with respect to the types of genes preserved as duplicate pairs. We compared gene duplicates resulting from a recent whole genome duplication to a set of tandemly duplicated genes in the model forest tree *Populus trichocarpa*. We used a combination of microarray expression analyses of a diverse set of tissues and functional annotation to assess factors related to the preservation of duplicate genes of both types. Whole genome duplicates are 700 bp longer and are expressed in 20% more tissues than tandem duplicates. Furthermore, certain functional categories are over-represented in each class of duplicates. In particular, disease resistance genes and receptor-like kinases commonly occur in tandem but are significantly under-retained following whole genome duplication, while whole genome duplicate pairs are enriched for members of signal transduction cascades and transcription factors. The shape of the distribution of expression divergence for duplicated pairs suggests that nearly half of the whole genome duplicates have diverged in expression by a random degeneration process. The remaining pairs have more conserved gene expression than expected by chance, consistent with a role for selection under the constraints of gene balance. We hypothesize that duplicate gene preservation in *Populus* is driven by a combination of subfunctionalization of duplicate pairs and purifying selection favoring retention of genes encoding proteins with large numbers of interactions.

[Supplemental material is available for this article.]

Gene duplication functions as the primary driver of evolutionary novelty within higher eukaryotes (Lynch and Conery 2000; Semon and Wolfe 2007). The recent sequencing of several plant genomes has demonstrated that both whole genome and segmental duplications have played major roles in the expansion of angiosperm gene families (Blanc and Wolfe 2004; Tuskan et al. 2006; Jaillon et al. 2007; Ming et al. 2008; Schnable et al. 2009; Schnutz et al. 2010). Whole genome duplications (WGDs) in particular appear to have occurred recurrently throughout the history of the angiosperm lineage (Blanc and Wolfe 2004; Freeling 2009; Paterson et al. 2010; Jiao et al. 2011). Ancient WGDs in diploid lineages have undergone a fractionation of the polyploid genome, during which chromosomal rearrangements, gene conversions, heightened translposon activity, and epigenetic changes left behind a reduced set of duplicate gene pairs (Chen and Ni 2006; Gaeta et al. 2006; Tate et al. 2009; Wang et al. 2010).

Clusters of duplicated genes have also formed through tandem duplication (TD) processes, which have greatly expanded some gene families, such as the Nucleotide Binding Site-Leucine Rich Repeat (NBS-LRR) subset of plant resistance genes (Meyers et al. 2003; Leister 2004; Kohler et al. 2008). Unequal recombination is thought to be the primary mechanism driving the expansion of these gene clusters (Leister 2004; Babushok et al. 2007; Kane et al. 2010). This occurs when interspersed repetitive elements promote crossing over between nonhomologous segments during meiosis or recombinational repair, resulting in the concomitant introduction of a deletion in one chromosome and an insertion in the other. Tandem duplication can also occur through insertion of retrotransposed genes, although these are thought to insert in a random manner and are often pseudogenized at birth because they lack a promoter and have a processed structure (Zhang et al. 2005; Babushok et al. 2007).

Following duplication, each gene within a paralogous pair may evolve in several ways. For example, it may retain the same set of functions as the ancestral copy (Davis and Petrov 2004), retain only a subset of the original set of functions (subfunctionalization) (Force et al. 1999; Lynch and Force 2000), obtain a new function (neofunctionalization), or degrade into a nonfunctional gene (nonfunctionalization) (Ohno 1970). Notably, the processes of subfunctionalization and neofunctionalization may not be mutually exclusive. Indeed, the degenerative processes leading to subfunctionalization may act upon silencer elements and thereby promote neofunctionalization (Huminiecki and Wolfe 2004). Seminal theory concerning the fates of duplicate genes predicted
that the duplicate copy would be shielded from purifying selection by the ancestral copy, thus promoting pseudogenization in the absence of positive selection for a rare acquired function (Ohno 1970). However, the presence of large numbers of duplicate genes derived from ancient polyploidy events is difficult to reconcile with a model in which null alleles at duplicate loci are easily fixed by genetic drift (Force et al. 1999). Furthermore, duplicate genes show evidence of purifying selection more consistent with buffering of the ancestral gene function than neutralization (Chapman et al. 2006; Hakes et al. 2007; Warren et al. 2010). Force et al. (1999) reconciled Ohno’s original theory with more recent observations by proposing subneutralization as a means of preserving duplicate genes in the presence of degenerative mutations targeting both members of a duplicate pair. This hypothesis, known as the duplication-degeneration-complementation (DDC) process, postulates that degenerative mutations may knock out independent subfunctions encoded by discrete regulatory elements in duplicate genes, thus requiring preservation of both copies in order to maintain the full complement of ancestral gene functions.

More recent models of duplicate gene evolution suggest that rates of duplicate gene retention vary among protein functional groups. Observations of high retention rates among more connected proteins are consistent with the gene balance hypothesis, which predicts that the fate of duplicate genes largely depends on maintaining a stoichiometric balance among members of macromolecular complexes (Freeling 2006; Birchler and Veitia 2007; Edger and Pires 2009; Birchler and Veitia 2010). This hypothesis also predicts that an increasing number of protein-protein interactions should favor retention of WGD pairs while disfavoring the fixation of TD. Indeed, empirical data in yeast and Arabidopsis demonstrate that genes involved in signal transduction and transcription are more likely to be retained following a WGD but less likely to be retained in tandem (Seoighe and Gehring 2004; Davis and Petrov 2005; Maere et al. 2005). Meanwhile, the converse is true for other genes, such as those containing NBS-LRR motifs (Meyers et al. 2003, 2005; Leister 2004; Zhang et al. 2010).

The availability of whole genome sequence and transcriptome data allows us to test the extent to which natural patterns of retention and divergence conform to the predictions of alternative models of duplicate gene evolution. Under the gene balance hypothesis, we expect that WGD and TD genes should have inverse patterns of retention, with retained WGD genes well-conserved and biased toward more central roles in networks (Freeling 2009). Alternatively, a pure subneutralization or neofunctionalization process should lead to extensive divergence of expression between duplicates, with retention patterns primarily driven by stochastic processes.

We used the model forest tree Populus trichocarpa (Torr. & Gray) to examine the factors involved in the preservation of duplicate gene function and expression. P. trichocarpa is an excellent model system for the study of duplicate gene evolution because of the large syntenic regions conserved from the relatively recent Salicoid WGD that is shared across the Salicaceae, containing nearly 8000 similarly aged paralogous gene pairs (Sterck et al. 2005; Tuskan et al. 2006; Berlin et al. 2010). Using a combination of coding sequence annotations and microarray expression data, we aimed to accomplish the following: (1) identify gene characteristics associated with retention following WGD and TD; (2) delineate the factors associated with diversification of expression patterns for gene pairs resulting from the Salicoid WGD; and (3) determine the degree to which whole genome patterns of duplicate gene retention and expression conform to the expectations of the DDC hypothesis.

Results

Overview of gene expression and duplications

We studied gene expression across a diverse set of tissues from field-grown P. trichocarpa trees, including various vegetative tissues and different stages of reproductive development (Supplemental Table S1). We identified 31,445 genes with significant expression levels in at least one of the 14 tissues analyzed (Supplemental Table S2). The vast majority of genes were expressed in both floral and vegetative tissues. However, 4306 transcripts were only detected in floral tissues that ranged from early floral development to early and late fruit/seed development stages, and 1423 transcripts were only detected in vegetative tissues (Supplemental Table S3). Due to source differences between reproductive and vegetative samples, some tissue specificity may be due to sample rather than biological effects. Approximately half of the expressed genes—15,253—have paralogs that, based on the fourfold degenerate transversion rate (4DTV) distances and syntenic positions, presumably date to the Salicoid WGD (Tuskan et al. 2006; Tang et al. 2008b). These are hereafter referred to as “retained” Salicoid duplicates. We also identified 1196 TD genes with pairwise 4DTV distances comparable to those of the Salicoid duplicates. The sizes of tandem arrays ranged from two to 15, with more than half (64%) only containing two duplicates.

Factors associated with occurrence of gene duplicates

We used logistic regression to identify significant predictors of occurrence of duplicate genes resulting from WGD and TD. Candidate variables included gene ontology (GO) functional categories, breadth of expression, and the genomic length of the gene. Sixteen out of the 18 variables we tested were significant predictors of Salicoid duplicate retention (Nagelkerke pseudo-$r^2 = 0.07$, $P < 2 \times 10^{-16}$), and 12 were significant predictors of TD gene presence (Nagelkerke pseudo-$r^2 = 0.121$, $P < 2 \times 10^{-16}$) (Table 1). Interestingly, eight of the 11 predictors that were significant in both sets had contrasting effects on the presence of duplicates in either category (Fig. 1).

Gene length was one factor associated with the occurrence of genes in both duplication categories. Gene length was positively correlated with the odds of Salicoid retention, while it was negatively associated with the odds of TD occurrence. Furthermore, Salicoid duplicates were significantly longer than all other genes in the genome, while TD genes were significantly shorter than all other genes (Table 2).

Expression breadth (i.e., the fraction of tissues in which significant expression occurs) exhibited a similar pattern to that observed for gene length. Higher expression breadth was associated with greater odds of retention as a Salicoid duplicate pair and with lower odds of occurrence in a TD (Fig. 1; Tables 1, 2). Furthermore, Salicoid duplicates had significantly greater expression breadth than all other genes in the genome, while the opposite was true of TD genes (Table 2).

Functional categories of gene duplicates

Genes with transcription factor activity, protein binding activity, kinase activity, phosphatase activity, nucleic acid binding, transporter activity, ligase activity, protease activity, and cation binding activity were associated with significantly higher odds of Salicoid duplicate retention. Conversely, the presence of transmembrane regions, receptor activity, catalytic activity, and stress responsiveness were associated with decreased odds of retention. Remarkably, these
same functional categories had the exact opposite effects on odds of occurrence in a TD (Fig. 1).

Three categories—protein binding, transporter activity, and kinase activity—were associated with increased odds of occurrence for both Salicoid and TD genes. However, closer inspection of the composition of these groups revealed substantial differences between tandem and Salicoid duplicates (Supplemental Tables S4, S5). Among protein binding genes, TDs were highly enriched for genes with leucine rich repeats (LRR), Ankyrin repeats, Toll/Interleukin-1 Receptor (TIR) domains, and stress responsiveness (Fig. 2A). The LRR and TIR domains are primarily components of plant resistance genes (R-genes) which are known to have a tendency to occur in tandem arrays (Meyers et al. 2003; Kohler et al. 2008). Interestingly, proteins containing LRR and TIR domains were under-represented among Salicoid duplicates, while the categories most enriched for Salicoids—RING fingers and DNA-dependent transcriptional regulators—were under-represented among TDs.

There was also a large discrepancy in the composition of the protein kinase groups between the Salicoid and TD genes (Fig. 2B). Ninety-eight percent of tandemly duplicated protein kinases were annotated for a class of receptor-like kinases (PANTHER acc: PTHR23258), while this class was under-represented among Salicoid duplicates. Similarly, 21% of tandem protein kinases, versus 1% of Salicoids, were annotated as S-locus-type glycoproteins, characterized by B-lectin and Pg/Apple/Nematode (PAN) domains. Proteins of this class are primarily known for their roles in self-incompatibility (Bassett et al. 2005; Chen et al. 2006). However, these proteins appear to also have roles in both defense and osmotic stress responses (Bassett et al. 2005; Chen et al. 2006). Salicoid duplicates showed over-retention of a class of protein kinases characterized by the SMART S_TKc annotation (acc. SM00220), which corresponds to the catalytic domain of serine/threonine-specific kinases. Despite its presence within 15% of all protein kinases, no TD genes were annotated for this domain. Closer inspection of this class revealed that most of the proteins were cyclin-dependent or calcium/calmodulin-dependent kinases, suggesting that this annotation primarily identified kinases involved in signal transduction pathways.

Both tandem and Salicoid duplicates were annotated for stress responses. However, genes from the two groups differed with respect to the types of stressors to which they respond (Fig. 2C). Genes involved in defense response and apoptosis were over-represented among TD genes and significantly under-retained among Salicoids. One domain common among plant resistance genes, the TIR domain, was present within 26% of stress-related tandem genes but only in 1% of Salicoids. However, genes responding to oxidative stress were over-represented among Salicoid pairs, comprising 27% of all stress-related genes within this group. Also present were proteins with roles in DNA repair (16%), osmotic stress (4%), and heat shock (3%) responses. In contrast, only 8% of tandem stress-related genes were involved in oxidative stress, and 1% had roles in DNA repair.

Our list of TD genes did not include any that were annotated for osmotic stress or heat-shock responses.

### Table 1. Significant predictors for retention of genes in Salicoid and tandem duplicate pairs

<table>
<thead>
<tr>
<th>Salicoid duplicates</th>
<th>TDs</th>
</tr>
</thead>
<tbody>
<tr>
<td>β</td>
<td>exp(β)</td>
</tr>
<tr>
<td>(Intercept)</td>
<td>-0.57</td>
</tr>
<tr>
<td>Tandem</td>
<td>-1.35</td>
</tr>
<tr>
<td>Salicoid</td>
<td>N/A</td>
</tr>
<tr>
<td>Breadth</td>
<td>0.57</td>
</tr>
<tr>
<td>Gene length</td>
<td>0.10</td>
</tr>
<tr>
<td>Protein binding</td>
<td>0.29</td>
</tr>
<tr>
<td>Transmembrane</td>
<td>-0.10</td>
</tr>
<tr>
<td>Transcription factor</td>
<td>0.66</td>
</tr>
<tr>
<td>Catalytic</td>
<td>-0.19</td>
</tr>
<tr>
<td>Receptor</td>
<td>-1.49</td>
</tr>
<tr>
<td>Stress</td>
<td>-0.74</td>
</tr>
<tr>
<td>Kinase</td>
<td>0.25</td>
</tr>
<tr>
<td>Phosphatase</td>
<td>0.70</td>
</tr>
<tr>
<td>Nucleic acid binding</td>
<td>0.15</td>
</tr>
<tr>
<td>Transporter</td>
<td>0.17</td>
</tr>
<tr>
<td>Ligase</td>
<td>0.61</td>
</tr>
<tr>
<td>Protease</td>
<td>0.24</td>
</tr>
<tr>
<td>Cation binding</td>
<td>0.13</td>
</tr>
</tbody>
</table>

### Figure 1.

Log$_{10}$ of the exponentiated logistic regression coefficients for categories significant for both retention of Salicoid duplicates and the presence of TDs. The log$_{10}$ scales the odds ratios for each category such that those above and below 1 have comparable effects.
Among TD genes associated with transmembrane regions, there was a significant over-representation of protein kinases, LRR and Ankyrin repeat protein binding domains, and S-locus glycoproteins (Fig. 2D). In turn, Salicoid duplicates showed under-representation of protein kinases, Ankyrin repeat, S-locus glycoproteins, and proteins involved in oxidation-reduction. However, RING finger domains were also significantly over-represented among Salicoid duplicates, concordant with the high retention of this class of proteins among genes annotated for protein binding.

**Expression divergence**

There was a complex relationship between expression breadth and the distance between expression patterns for Salicoid duplicates. At low combined breadth (the total fraction of tissues in which at least one duplicate has significant expression), the nonparametric correlation behind the conservation calculation inflates the proportion of pairs with conserved expression due to the small number of tissues in which significant expression occurs (Fig. 3). This artifact disappears above a combined breadth of 0.4, after which pairs from the Salicoid duplication show significantly higher conservation than expected by chance. Furthermore, the proportion of Salicoid

### Table 2. Mean, median, and standard error for the gene length, expression breadth, and d\_N/d\_S (of pairs) within the whole genome, among Salicoid duplicates, and among TDs

<table>
<thead>
<tr>
<th>Group</th>
<th>Mean</th>
<th>Median</th>
<th>SE</th>
<th>P-value*</th>
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</thead>
<tbody>
<tr>
<td>Gene length</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Whole genome</td>
<td>2.587 kb</td>
<td>2.007 kb</td>
<td>0.0126</td>
<td>0.0000</td>
</tr>
<tr>
<td>TDs</td>
<td>2.126 kb</td>
<td>1.767 kb</td>
<td>0.0471</td>
<td>0.0000</td>
</tr>
<tr>
<td>Salicoid duplicates</td>
<td>2.854 kb</td>
<td>2.304 kb</td>
<td>0.0183</td>
<td>0.0000</td>
</tr>
<tr>
<td>Breadth</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Whole genome</td>
<td>0.5396</td>
<td>0.5714</td>
<td>0.00185</td>
<td>0.0000</td>
</tr>
<tr>
<td>TDs</td>
<td>0.3868</td>
<td>0.2857</td>
<td>0.00875</td>
<td>0.0000</td>
</tr>
<tr>
<td>Salicoid duplicates</td>
<td>0.5774</td>
<td>0.6428</td>
<td>0.00264</td>
<td>0.0000</td>
</tr>
<tr>
<td>d_N/d_S</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Salicoid duplicates</td>
<td>0.2728</td>
<td>0.2428</td>
<td>2.29 \times 10^{-3}</td>
<td>0.0000</td>
</tr>
<tr>
<td>TDs</td>
<td>0.4478</td>
<td>0.40295</td>
<td>6.57 \times 10^{-4}</td>
<td>0.0000</td>
</tr>
</tbody>
</table>

*P-values for gene length and breadth are based on van der Waerden tests for independence of gene characteristic and duplicate type.

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**Figure 2.** Relative frequencies of specific annotations within the broad functional categories of protein binding (A), stress response (B), protein kinase (C), and transmembrane regions (D). Annotations were generated by InterproScan, and those shown were among the most common within each broader functional category. Relative protein frequency refers to the fraction of proteins within the broad category that contain the specific annotation. Error bars indicate 95% confidence intervals generated by 1000 bootstrap replicates.
A nonlinear relationship between low expression imbalance (Fig. 5). Distance was most evident at high combined breadth and skew (D’Agostino skewness test, **Furthermore, the observed distribution had a significant positive** correlation of random divergence processes (mean = 0.674, sd = 0.373) were significantly less than expression distances resulting from a simulation of random divergence processes (mean = 0.972, sd = 0.371). Furthermore, the observed distribution had a significant positive skew (D’Agostino skewness test, \(P = 2.074 \times 10^{-14}\)), suggesting that it may be comprised of a mixture of two underlying distributions (Fig. 4). Interestingly, genes annotated for transcription factor and nucleic acid binding activities were significantly over-represented in the left distribution with more conserved expression patterns, while genes with receptor activity or transmembrane regions were over-represented in the right distribution with less expression conservation (Fisher’s exact test, FDR = 0.05) (Table 3).

**Discussion**

Our analyses of duplicate pairs resulting from the Salicoid WGD and TD suggest that duplicate gene retention is driven by multiple processes. We have interpreted these results with respect to the DDC model, or subfunctionalization (Force et al. 1999). As expected under a subfunctionalization or neofunctionalization process, we find extensive divergence of expression among retained Salicoid duplicates, which is associated with increased sequence divergence. Furthermore, although our ability to detect subfunctionalization versus neofunctionalization is limited by the lack of an outgroup, population genetic theory suggests preservation by neofunctionalization should be rare when the effective population size is on the order of that in *Populus* (Lynch et al. 2001). We, therefore, hypothesize that subfunctionalization is the predominant process for this subset of Salicoid duplicates. However, we also find that many Salicoid duplicates have more conserved expression patterns than expected under random divergence and that tandemly duplicated genes strongly contrast with genes from the recent whole genome duplication with respect to both structural and expression characteristics. This suggests the degenerative process of subfunctionalization may be counterbalanced by a selective drive to retain highly connected proteins, as predicted by the gene balance hypothesis (Birchler et al. 2005; Freeling 2009; Birchler and Veitia 2010).

**Salicoid duplicates are longer and more broadly expressed**

Salicoid duplicate genes are significantly longer than other genes in the genome, whereas tandem duplicates are significantly shorter. Although increased length may leave these genes more susceptible to loss-of-function mutations, longer genes may also have an enhanced ability to subfunctionalize within the coding region, as

**Figure 3.** The proportion of Salicoid duplicate genes with conserved expression plotted against the combined breadth of the two duplicate genes for the observed and random distributions. Conserved expression was defined as a Spearman expression distance less than 0.3, which corresponded to the lower 5% cutoff of expression distances for permuted genes. Error bars denote the standard error obtained by 1000 bootstrap replicates.

**Figure 4.** Line plots of the histograms for the observed distribution of Spearman expression distances (solid gray) and the simulated distribution under the assumption of random divergence (solid black). The probability density functions for each of the two truncated normal mixture model distributions are also shown at their mixture proportions (dashed). Parameters for the left mixture distribution were mean (\(\mu_1\)) = 0.3907, standard deviation (\(\sigma_1\)) = 0.2470, mixture proportion (\(\pi_1\)) = 0.5294. Parameters for the right distribution were \(\mu_2\) = 0.9630, \(\sigma_2\) = 0.2944, and \(\pi_2\) = 0.4706.

**Ratio of nonsynonymous to synonymous substitutions and expression imbalance**

Salicoid duplicates had a significantly lower ratio of nonsynonymous to synonymous substitutions (\(d_N/d_S\)) in coding regions compared to TD genes (Table 2). This was true over a range of \(d_S\) values (Supplemental Figs. S1, S2). Furthermore, using a forward selection strategy, we found a nonlinear relationship between \(d_S/d_N\) and expression distance that depended on the interaction between combined breadth and expression imbalance (\(r^2 = 0.188, P < 2.2 \times 10^{-16}\)) (Table 4). A nonlinear relationship between \(d_S/d_N\) and expression distance was found to be superior to a linear fit using an F-test (F = 7.10, P = 0.007725). The relationship between \(d_S/d_N\) and expression distance was most evident at high combined breadth and low expression imbalance (Fig. 5).
functions. Furthermore, the proportion of TD genes for which weaker purifying selection for the retention of ancestral gene of magnitude higher than for Salicoid duplicates (Kong et al. 2007; Freeling 2009; Woodhouse et al. 2010). We contend predicts that TDs of longer genes with more sites of protein-protein cation. These genes would also be more likely to pseudogenize following any single duplication event due to their limited number of subfunctions. Therefore, shorter genes that complete set of ancestral regulatory regions, thereby producing retrotranspositions will entirely remove in- trons. Similarly, these phenomena will also often fail to copy the relatively high expression breadth of Salicoid duplicates appears to contradict the neofunctionalization following the loss of silencer elements (Huminiecki and Wolfe 2004). Furthermore, breadth does not account for quantitative subfunctionalization within tissues. Finally, patterns of expression are only a crude approximation of function: Duplicate pairs may have divergent biochemical or physiological roles due to structural differentiation of the proteins even when expressed in the same tissues.

The shorter lengths and lower breadths of TD genes most likely reflect the stochastic processes leading to their birth—unequal crossing over, transposition, and intrachromosomal recombination—and relatively frequent translocations within the genome (Kong et al. 2007; Freeling 2009; Woodhouse et al. 2010). Each of these may result in the incomplete duplication of genes in the tandem set, and retrotranspositions will entirely remove in- trons. Similarly, these phenomena will also often fail to copy the complete set of ancestral regulatory regions, thereby producing subfunctionalized genes at birth. Therefore, shorter genes that function in a tissue-specific manner may be more likely to produce a viable copy following the degenerative process of tandem duplica- tion. These genes would also be more likely to pseudogenize fol- lowing any single duplication event due to their limited number of cis regulatory elements. However, the gene balance hypothesis also predicts that TDs of longer genes with more sites of protein-protein interaction should be quickly eliminated due to the deleterious ef- fects of dosage imbalance (Birchler and Veitia 2010). We contend that the higher ds/ds observed between TD genes is consistent with weaker purifying selection for the retention of ancestral gene functions. Furthermore, the proportion of TD genes for which there was evidence positive selection (i.e., ds/ds > 1) was an order of magnitude higher than for Salicoid duplicates (P = 7.75 × 10−10). This would suggest that for TDs, maintenance of the full complement of ancestral gene functions may not be the primary factor favoring duplicate retention, as expected under a DDC process (Force et al. 1999).

### Table 3. Proportion of genes annotated with a given functional category in each of the two distributions of the mixture model

<table>
<thead>
<tr>
<th>Category</th>
<th>Mix. dist. 1</th>
<th>Mix. dist. 2</th>
<th>p-value*</th>
<th>q-value</th>
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<tbody>
<tr>
<td>Protein binding</td>
<td>0.11</td>
<td>0.11</td>
<td>0.92</td>
<td>0.51</td>
</tr>
<tr>
<td>Transmembrane</td>
<td>0.28</td>
<td>0.31</td>
<td>0.01</td>
<td>0.03</td>
</tr>
<tr>
<td>Transcription factor</td>
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<td>0.02</td>
<td>0.00</td>
<td>0.00</td>
</tr>
<tr>
<td>Catalytic</td>
<td>0.29</td>
<td>0.32</td>
<td>0.07</td>
<td>0.11</td>
</tr>
<tr>
<td>Receptor</td>
<td>0.00</td>
<td>0.00</td>
<td>0.02</td>
<td>0.04</td>
</tr>
<tr>
<td>Ion</td>
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<td>0.00</td>
<td>0.53</td>
<td>0.38</td>
</tr>
<tr>
<td>Stress</td>
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<td>0.01</td>
<td>0.11</td>
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<tr>
<td>Kinase</td>
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<td>0.05</td>
<td>0.22</td>
<td>0.21</td>
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<tr>
<td>Phosphatase</td>
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<td>0.01</td>
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<td>Nucleic acid binding</td>
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<td>Transporter</td>
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<td>0.04</td>
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<td>Ligase</td>
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<td>0.02</td>
<td>0.75</td>
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<tr>
<td>Protease</td>
<td>0.03</td>
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<tr>
<td>Cation binding</td>
<td>0.12</td>
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<td>0.47</td>
<td>0.38</td>
</tr>
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</table>

*p-values were obtained using Fisher’s exact test, while the q-value is the false discovery rate (FDR) analog.

### Table 4. Regression table for the relationship between expression distance and ds/ds, combined breadth, and expression imbalance

| Term                  | Estimate | Std. error | t value | Pr(>|t|) |
|-----------------------|----------|------------|---------|---------|
| (Intercept)           | 0.5331   | 0.0481     | 11.09   | 0.0000  |
| ds/ds                 | 0.2607   | 0.1829     | 1.43    | 0.1540  |
| (ds/ds)^2             | −0.1915  | 0.0719     | −2.66   | 0.0077  |
| Combined breadth      | −0.2004  | 0.0511     | −3.92   | 0.0001  |
| Exp. imbalance        | 0.0719   | 0.0032     | 22.40   | 0.0000  |
| ds/ds:combined breadth| 0.5982   | 0.1907     | 3.14    | 0.0017  |
| ds/ds:exp. imbalance  | −0.1903  | 0.0281     | −6.78   | 0.0000  |
| ds/ds:combined breadth:exp. simbalance | 0.1324 | 0.0313 | 4.23 | 0.0000 |

*The estimate refers to the linear regression coefficient for each predictor variable.
freed from purifying selection for retention of both duplicates, assuming they are not interaction network hubs. One intriguing possibility is that R-genes may encounter negative selection following WGD because of the trade-off between the increased fitness conferred by resistance when the pathogen is present and decreased fitness in its absence (Tian et al. 2003; Meyers et al. 2005). Thus, retention of the entire complement of functional R-genes during the diploidization process may have had fitness costs in the absence of an onslaught of new pathogens, leading to selection against redundant resistance loci.

Lastly, we observed that TD genes tend to display much more extreme patterns of functional over-representation than Salicoids. This illustrates an important distinction between the modes of duplication. While the set of Salicoid duplicates reflects the pattern of retention following a single duplication event, the tandemly duplicated genes reflect both the propensity of certain genes to duplicate in tandem and the rate of elimination by selection or drift. In the case of R-genes, stress is known to increase rates of somatic recombination, which can facilitate the birth of new genes through unequal crossing over if it occurs within reproductive cell lineages (McDowell and Simon 2006). Moreover, the presence of repeat elements, including the repeated structures of genes within tandem arrays, can also increase the rate of tandem duplication by promoting unequal crossing over (Jelesko et al. 1999). Indeed, an increased rate of birth may explain how gene families can expand through duplication in the absence of purifying selection due to gene balance constraints or the capacity to subfunctionalize with a limited repertoire of ancestral subfunctions. Similarly, the properties of tandem arrays may provide a mechanism for their rapid deletion following WGD, as these regions are known to have high rates of intrachromosomal recombination (Woodhouse et al. 2010).

Expression divergence consistent with two different patterns

Although the expression patterns of Salicoid duplicates were generally more conserved than expected by chance, we interpreted the skewness of the distribution of expression distances for pairs of Salicoid duplicates as an indication that this distribution might be appropriately modeled as a mixture of two distributions. One interpretation based on the estimated parameters of our mixture model is that approximately half (45%) of the duplicate gene pairs diverge in expression following a pattern consistent with a DDC process, wherein regulatory elements randomly degenerate, eventually leading to complete subfunctionalization (Force et al. 1999). The remaining 55% of duplicate gene pairs appear constrained to maintain some redundancy in their expression patterns. Such a constraint is consistent with the expectations of the gene balance hypothesis, in that the drive to retain post-duplication stoichiometric balance should lead to more conserved patterns of expression (Aury et al. 2006; Birchler and Veitia 2010). This is supported by the observation that the GO category for nucleic acid binding proteins—including highly connected transcription factors and ribosomal proteins—was significantly over-represented in the left distribution. Moreover, the proportion of well-conserved Salicoid duplicates is positively associated with the combined breadth of the genes, suggesting that more ubiquitous expression of the putative ancestral gene leads to greater conservation of its descendants.

Expression and sequence divergence correlated for broadly expressed genes

Under the expectations of the DDC model, we would predict that the degradation of regulatory elements would occur independently of coding sequence mutations, as the duplicates are assumed to be initially redundant and unconstrained by requirements to maintain the same set of interactions (MacCarthy and Bergman 2007). In contrast, the gene balance hypothesis predicts that the expression patterns of duplicates will be constrained primarily by their protein-protein interactions (Birchler and Veitia 2010). This implies that changes in expression may accompany nonsynonymous mutations, thereby maintaining interactions within an evolving network. Our results suggest that both processes may be affecting Salicoid duplicates.
We found that there was no discernible relationship between expression distance and $d_{S}/d_{N}$ for gene pairs with low combined expression breadth and high expression imbalance. However, broadly expressed (i.e., combined breadth above 80%) gene pairs did have a significant nonlinear relationship between expression distance and $d_{S}/d_{N}$. Previous studies in *Populus* revealed that more broadly expressed genes tend to be predominantly cis regulated, while trans regulation drives more tissue-specific expression (Quesada et al. 2008; Drost et al. 2010), a finding that is broadly consistent with our results. Our data further suggest that expression divergence in broadly expressed genes may be reflected in coding sequence polymorphisms as well as variation in cis regulatory domains in noncoding regions. Results from previous studies indicated that the most broadly expressed genes tend to have the slowest evolutionary rates (Pal et al. 2001; Ettwiller and Veitia 2007). This is consistent with the negative correlation we observed between combined breadth and $d_{S}/d_{N}$. Moreover, under the constraints of the gene balance hypothesis, we would expect the most highly connected genes to be subject to the greatest selective pressure for maintenance of balanced expression between interacting subunits. Indeed, interacting subunits of macromolecular complexes tend to have positively correlated patterns of expression and evolutionary rates (Ettwiller and Veitia 2007). Therefore, we predict that the strongest relationship between expression distance and $d_{S}/d_{N}$ occurs for the most highly connected genes, which are selected for cis regulatory elements and protein motifs that allow them to maintain balance within the paleopolyploid protein interaction network. This will be the subject of further investigations in our lab.

**Conclusion**

We find the pattern of duplicate gene retention following the Salicoid WGD in *Populus* broadly consistent with the predictions of the gene balance hypothesis. Future investigations should use a network-based approach to directly gauge whether the most connected genes are most highly retained following WGD. Approximately half of the Salicoid duplicate gene pairs showed patterns of divergence that suggest many whole genome duplicates are not subject to constraints for maintenance of redundancy. A future analysis of nonconserved noncoding regions would enable us to more accurately determine the role of degenerative processes in the observed expression patterns of duplicate genes.

We believe our findings are relevant to the evolution of duplicate genes across a wide range of higher eukaryotes. Taken together, our findings imply the patterns of retention and functional conservation observed following duplication events are contingent upon both random and selective forces, although one or the other tends to predominate depending upon the type of duplication and the biochemical function of the gene. This study, therefore, serves as the groundwork for more detailed studies of the relative roles of neutral processes and natural selection in shaping the functional landscape of the paleopolyploid genome.

**Methods**

**Populus whole genome microarray experiments**

The *Populus* whole genome microarray was constructed by Roche NimbleGen (http://www.nimblegen.com/) to target 55,794 nuclear, 69 chloroplast, and 58 mitochondrial gene models predicted from version 1.1 of the *Populus trichocarpa* genome (Tuskan et al. 2006). The array included three 60mer oligonucleotide probes per gene target, which were evaluated for identity to other nontarget gene models using WU-BLASTN as an index of potential cross-hybridization and then further refined based on Nimblegen’s design guidelines. Prior to the final design, all nontarget probes were removed along with probes for targets with high identity to transposable elements.

Tissues for microarray hybridizations were obtained from field-grown *Populus trichocarpa* trees near Corvallis, Oregon, except that one root sample was collected from in vitro-grown plants, and seeds were germinated in vitro. All tissue was obtained from clone Nisqually-1, except for floral tissues and seeds/seedlings, which were collected from wild *P. trichocarpa* trees. Tissues and abbreviations are described in Supplemental Table S1. RNA isolation, labeling, and hybridization were conducted as described in Dharmawardhana et al. (Dharmawardhana et al. 2010). Two biological replicates were used for each tissue sample, and three biological replicates were used for the xylem sample.

**Collection and normalization of microarray data**

The NimbleGen microarray data processing pipeline (NMP) was used to normalize the data using a two-step normalization procedure (Li et al. 2006; Wang et al. 2006). In the first step, quantile normalization was performed among replicates within each tissue, followed by global normalization to adjust all tissues to a similar baseline. ANOVA was used to identify significant differentially expressed genes between tissues. For each gene, the significance of the differences in the mean of the log$_{2}$C of intensities between any two tissues was calculated using t-tests. False discovery rate (FDR) was calculated to correct for multiple testing problems (Benjamini and Hochberg 1995). A gene expression difference with estimated positive (up)- or negative (down)-fold change at alpha > 0.05 and FDR (q) = 0.05 was considered significant.

Thresholds for significant expression within the microarray were set using probes for 3149 transposable elements as negative controls, using the 95th percentile as a cutoff. Tissues were tested for correspondence among replicates (Supplemental Figs. S3, S4), and individual replicates were clustered based on Euclidean distances (Supplemental Fig. S5). Tissue samples that showed strong correspondence between replicates and which clustered together were used for subsequent analyses.

Although the microarray was based on gene models from *Populus* version 1.1, the results were ported to *Populus* version 2.0 using an approach based on synteny and reciprocal best hits following BLASTP of the models against one another. Prior to the analysis, tissue replicates were averaged together, and values were averaged over version 1 gene models if ambiguity existed in the version 1 to version 2 mapping. Average expression values were set to zero if they did not exceed the threshold set by the negative controls. Otherwise, they were set to the observed average value minus the tissue threshold. Only gene models with significant expression in at least one tissue were used in subsequent analyses.

**Identification of duplicate pairs**

Potential duplicate pairs were identified by using BLASTP to compare all version 2 gene models against one another. Models with at least 50% identity over at least half the length of the larger gene were considered potential duplicates. The fourfold degenerate transversion rate (4DTV) was calculated after Smith-Waterman alignment of potential duplicates using an affine gap penalty (gap open = −10, gap extension = −1) and the BLOSUM 60 matrix for scoring. The corresponding CDS sequences were then superimposed onto the alignment, and 4DTV was calculated as the number of transversions within fourfold degenerate sites divided...
by the total number of fourfold degenerate sites (Hellsten et al. 2007).

MCScan was then used to discover intragenomic syntenic blocks (Tang et al. 2008a). Average 4DTV was calculated for syntenic segments by taking the mean across gene pairs found by MCScan in each segment. Potential duplicates on syntenic segments with average 4DTV between 0.08 and 0.15 were considered to be Salicoid duplicates, while gene pairs in these regions with 4DTV greater than 0.2 were filtered out due to concerns that they may have arisen from a more ancient duplication event. Potential duplicates that occurred within 100 kb of each other were defined as tandemly duplicated. In order to limit the set of tandemly duplicated genes to pairs with similar divergence times to the Salicoids, we only selected TDs with 4DTV between 0.06 and 0.16, a range that included most of the Salicoid duplicates but did not include the most recent TDs. Tandemly duplicated genes were permitted to have multiple Salicoid duplicates on a syntenic segment if they conformed to the previously stated criteria in order to avoid biases against tandem duplicates in the Salicoid WGD set.

Functional annotation
Each version 2 peptide sequence was annotated using 14 applications (BLASTProDom, FPrintScan, HMMPIR, HMPfam, HMMsmart, HMMtigr, ProfileScan, HAMAP, patternScan, SuperFamily, SignalPHMM, TMHMM, HMPpanther, and Gene3D) in conjunction with InterProScan (Quevillon et al. 2005). The resulting InterPro annotations were then cross-referenced to GO terms using the Gene Ontology hierarchy: protein binding (GO:0005515), transcription factor activity (GO:0003700), catalytic activity (GO:0003824), receptor activity (GO:0004872), ion channel activity (GO:0005216), stress response (GO:0006950), protein kinase activity (GO:0016301), phosphatase activity (GO:0016791), nucleic acid binding (GO:0003676), transporter activity (GO:0005215), l-gase activity (GO:0016874), protease activity (GO:0008233), and cation binding (GO:0043169).

Measures of expression divergence between duplicate pairs
The expression distance for each duplicate pair was defined as 1 – Sxy, where Sxy denotes the Spearman correlation coefficient between the tissue expression values of each gene following the normalization procedures described above. Combined breadth refers to the number of tissues in which either member of a duplicate pair has significant expression divided by the total number of tissues, and expression imbalance denotes the maximum breadth of the two significant expression divided by the total number of tissues, and the number of tissues in which either member of a duplicate pair has significant expression divided by the total number of tissues. Combined breadth was defined as the proportion of tissues in which either of the duplicate genes had significant expression, while expression imbalance was defined as the breadth of the more broadly expressed gene divided by the breadth of the more narrowly expressed gene.

Distribution of expression distances under random divergence
An empirical distribution of expression distances was constructed for Salicoid duplicates under a model of divergence consistent with a DDC process. This random model assumed the existence of an ancestral gene for each duplicate pair with expression in each tissue corresponding to the maximum of the descendant’s expression levels as well as random divergence from this ancestral expression pattern through purely degenerative processes. For each pair of Salicoid duplicates, a putative ancestral gene expression profile was constructed wherein each tissue was assumed to express the maximum of expression values of the two duplicate genes in that tissue. In order to replicate the observed differences in expression during the simulation, a distribution of expression differences was constructed. The percent difference between the two expression values for each tissue was added to a vector, which was subsequently divided into 50 bins of equal width, including 100% divergence (loss of expression for one of the duplicates in that tissue). Simulated duplicate genes were then made for each putative ancestral gene, wherein the putative ancestral expression level was assigned to each duplicated gene. These duplicates then underwent simulated divergence, during
which one gene would be randomly selected in each tissue to have its expression reduced by a quantity sampled from the distribution of percent differences in the observed data. The expression distance was then calculated for each pair of genes as indicated above.

Modeling of the observed expression distances

The distribution of Spearman correlation coefficients is known to be related to Student’s t distribution (Press et al. 1992), which converges to the normal distribution with large N. Because the expression distance measure has a range between 0 and 2, the observed distribution of expression distances was modeled as a mixture of two truncated normal distributions (Johnson et al. 1994). The Nelder-Mead simplex algorithm provided by the optim function in the R programming language was used to maximize the following log-likelihood function for values of the observed Spearman expression distances between Salicoid duplicates:

\[
\ln(L(\pi_1, \mu_1, \sigma_1, \pi_2, \mu_2, \sigma_2, |D|)) = \sum_{i=1}^{n} \ln(\pi_1 f(x_i; \mu_1, \sigma_1) + \pi_2 f(x_i; \mu_2, \sigma_2)),
\]

where \( D \) denotes the data from the observed distribution of Spearman expression distances, \( \pi \) values denote the mixture proportions of each distribution, and \( \mu \) and \( \sigma \) denote the means and standard deviations of truncated normal distributions between 0 and 2. Initial parameter values were drawn from uniform distributions, wherein means (\( \mu \)) were permitted to range between the fifth and 95th percentiles of the observed distribution, standard deviations (\( \sigma \)) were permitted to range between 0.1 and the entire standard deviation of the observed distribution, and mixture proportions (\( \pi \)) were permitted to range between 0.1 and 0.9, with a sum-to-one constraint. Fitting of the mixture model was performed 100 times until the convergence at a relative tolerance of \( 1 \times 10^{-10} \). Final parameters were estimated by using a weighted average where the final likelihoods for each iteration served as weights.

Salicoid gene pairs were assigned to the mixture distributions using likelihood ratios. Gene pairs with a likelihood ratio for a distribution greater than or equal to 3 were assigned to the corresponding distribution. Otherwise, they were not assigned to any distribution. The over-representation of functional categories within distributions was then assessed using Fisher's exact test, with multiple testing error controlled using the false discovery rate, as implemented in the q-value package for the R programming language.

Data access

Microarray data used in this work have been submitted to the NCBI Gene Expression Omnibus (GEO) (http://www.ncbi.nlm.nih.gov/geo/) under accession numbers GSE21481 and GSE21485.

Acknowledgments

We thank Jennifer Hawkins and Phil Turk for helpful comments on the manuscript. This work was supported by the Office of Science (BER), U.S. Department of Energy (DOE), Grants No. DE-FG02-06ER64185 and DE-FG02-07ER64449, and by funding from the BioEnergy Science Center, a U.S. DOE Bioenergy Research Center supported by the Office of Biological and Environmental Research in the DOE Office of Science.

References


Evolution of tandem and whole genome duplications


