Early genome duplications in conifers and other seed plants

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Polyploidy is a common mode of speciation and evolution in angiosperms (flowering plants). In contrast, there is little evidence to date that whole genome duplication (WGD) has played a significant role in the evolution of their putative extant sister lineage, the gymnosperms. Recent analyses of the spruce genome, the first published conifer genome, failed to detect evidence of WGDs in gene age distributions and attributed many aspects of conifer biology to a lack of WGDs. We present evidence for three ancient genome duplications during the evolution of gymnosperms, based on phylogenomic analyses of transcriptomes from 24 gymnosperms and 3 outgroups. We use a new algorithm to place these WGD events in phylogenetic context: two in the ancestry of major conifer clades (Pinaceae and cypressophyte conifers) and one in Welwitschia (Gnetales). We also confirm that a WGD hypothesized to be restricted to seed plants is indeed not shared with ferns and relatives (monilophytes), a result that was unclear in earlier studies. Contrary to previous genomic research that reported an absence of polyploidy in the ancestry of contemporary gymnosperms, our analyses indicate that polyploidy has contributed to the evolution of conifers and other gymnosperms. As in the flowering plants, the evolution of the large genome sizes of gymnosperms involved both polyploidy and repetitive element activity.

INTRODUCTION

Polyploidy, or whole genome duplication (WGD), is one of the most important forces in vascular plant evolution. Nearly 25% of vascular plants are recent polyploids (1), with approximately 15% of angiosperm and 31% of fern speciation events due to genome duplication (2). Ancient polyploidy is found in the ancestry of all extant seed and flowering plants (3), and many angiosperm lineages have experienced additional rounds of genome duplication (4–10). Changes in the rates of molecular evolution and turnover in genome content following polyploidy may have provided novel genetic variation that was important for the evolution of plant diversity (3, 8, 11–16).

Despite the prevalence of polyploidy in the history of flowering plants, the role of polyploidy in gymnosperm evolution is less clear. The extant gymnosperms appear to be the sister clade of angiosperms (17), and they diverged from their most recent common ancestor (MRCA) as much as 310 million years ago (18). Most evidence indicates that polyploid speciation is relatively rare among extant gymnosperms (2), although in some genera (for example, Ephedra), polyploidy is prevalent (19, 20). Previous analyses of conifer genome sizes and chromosomes suggested that paleopolyploidy occurred in Pinaceae (19, 21). Although there was evidence of an ancient polyploidy shared by all seed plants (3), no evidence of a gymnosperm or conifer ancient polyploidy was found in the genome of Norway spruce (Picea abies), the first published gymnosperm genome. However, this conclusion was based on only a single plot of the relative ages of duplicate genes, presumably because the genome assembly was not of high enough quality (N50 = 4.87 kb) for syntenic analyses. Based on the pattern of accumulation of paralogs seen in this plot, they suggested that the large genomes of conifers originated by mechanisms exclusive of WGD, in particular through proliferation of long terminal repeat retrotransposons (LTR-RTs). Given that paleopolyploidy has been repeatedly observed among flowering plants and is also hypothesized to occur among conifers (19, 21), our goal was to test more thoroughly for evidence of ancient polyploidy in gymnosperms, using a phylogenetically diverse data set and a new phylogenomic method for determining the phylogenetic placement of WGDs.

We assembled transcriptomes for 24 gymnosperms and 3 outgroup species, including representatives of all major gymnosperm and vascular plant clades (table S1). Three of these transcriptomes—Ophioglossum petiolatum, Gnetum gnemon, and Ephedra frustillata—were newly sequenced to cover phylogenetic gaps in our data set. For each transcriptome, we used our DupPipe bioinformatic pipeline to generate age distributions of paralogs to identify shared bursts of gene duplication that are indicative of ancient WGD (7, 22, 23). We also introduce a newly developed algorithm, Multi-tAxon Paleopolyploidy Search (MAPS), to place inferred paleopolyploid events in phylogenetic context. For each node in a phylogeny, MAPS evaluates the percentage of gene duplications shared by all taxa descended from that node. Ancient WGDs are identified and located as peaks in plots of duplication events shared among a set of species (Materials and Methods; figs. S1 and S2). We used MAPS to confirm and locate genome duplication events in the history of the gymnosperms and seed plants.

RESULTS

Phylogenetic position of the ancient seed plant polyploidy

Most seed plant species contained evidence of a gene duplication peak consistent with previous evidence for a WGD in the ancestry of all seed plants (3). With the exception of the Gnetales taxa, each gymnosperm Ks plot (fig. S3) had a peak with a median Ks = 0.75 to 1.5, which, in some of these taxa, has previously been correlated with a WGD shared by all seed plants (3). Among the Gnetales, we only observed a peak with a median Ks = 1.05 in Welwitschia mirabilis, which is consistent with a Welwitschia-specific WGD (4). All three Gnetales taxa do not
contain clear evidence of the putative seed plant WGD, perhaps due to elevated substitution or gene birth/death rates among these species.

To place this ancient WGD in the vascular plant phylogeny, we implemented a new multispecies paleopolyploid search tool, MAPS. Previous analyses found evidence for an ancient polyploidy in the ancestry of all extant seed plants, Jiao et al. (3). However, a major clade of vascular plants, the monilophytes (ferns), was not included in that analysis. It was therefore unclear if this WGD is shared among all euphyllophytes (seed plants and monilophytes) or restricted to only seed plants. To better place this WGD in the vascular plant phylogeny, we analyzed new transcriptome data from the eusporangiate fern Ophioglossum with data from Araucaria (gymnosperm), Ginkgo (gymnosperm), Amborella (angiosperm), and Selaginella (lycophyte, the sister lineage to euphyllophytes). Gene trees were constructed for 3235 gene families with at least one gene copy present in each species. Among these gene families, MAPS identified 544 subtrees that included the MRCA of Araucaria, Ginkgo, and Amborella, which were consistent with the species tree. Nearly 64% of these subtrees contained evidence for a shared duplication in the MRCA of the seed plants that was not shared with Ophioglossum (Fig. 1A, fig. S4A, and table S2). This result demonstrates that the unclearly delimited euphyllophyte genome duplication (3) is indeed limited to seed plants as a whole and not shared with ferns and other vascular plants (Fig. 2).

Independent paleopolyploidies in Pinaceae and Cupressaceae

Most gymnosperm lineages only contained evidence for a single, ancient WGD, but some species had multiple signals. The Ks plots for most of the conifers contained a younger peak consistent with a WGD since the seed plant genome duplication (fig. S3). Among Pinaceae, we observed a younger peak with a median Ks = 0.2 to 0.4 for each taxon in our data set. Similarly, gene age distributions for taxa in Cephalotaxaceae, Cupressaceae, and Taxaceae contained a younger peak with a median Ks = 0.2 to 0.5. Araucaria was the only conifer in our data set without an unambiguous younger peak. Thus, the Ks plots suggest that there may have been one shared conifer WGD or independent WGDs in the history of different conifer families.

We conducted two different MAPS analyses to resolve the placement and number of WGDs among the conifers. For one analysis, we selected the transcriptomes of Pinus, Larix, and Cedrus to represent Pinaceae, and the transcriptome of Taxus to represent Taxaceae; we chose Ginkgo, Ophioglossum, and Selaginella as outgroups. We recovered 2175 gene family phylogenies with at least one gene copy from each taxon. MAPS identified 625 subtrees among these gene family phylogenies that included the MRCA of Pinaceae. More than 52% of the subtrees supported a shared duplication in the ancestry of Pinaceae (Fig. 1B, fig. S4B, and table S3). In contrast, only 9% of 535 subtrees supported a gene duplication shared between Pinaceae and Taxaceae. In the second analysis, we selected Taxus (Taxaceae), Cephalotaxus (Cephalotaxaceae), Cryptomeria (Cupressaceae), and Pinus (Pinaceae), with Ginkgo, Ophioglossum, and Selaginella as outgroups. Among 1886 gene family phylogenies for these taxa, MAPS identified 469 subtrees that included the MRCA of the cupressophytes. More than 42% of the subtrees supported a shared gene duplication in the MRCA of Cupressaceae and Taxaceae (Fig. 1C, fig. S4C, and table S4). Only 10% of the subtrees supported a duplication event shared by Pinaceae, Cupressaceae, and Taxaceae. We found similar results with MAPS using only gene trees with >50% bootstrap support for all branches (table S5). These results suggest that there are two ancient WGDs in the conifers: one shared by Cupressaceae and Taxaceae (the cupressophytes), and one in the ancestry of Pinaceae (Fig. 2).

Analyses of ortholog divergence corroborated our MAPS results and supported independent WGDs among the conifers. We identified 3266 orthologs by reciprocal best BLAST hit (22) from representatives...
of Pinaceae and Cupressaceae, *Picea glauca* and *Cryptomeria japonica*. Excluding poorly aligned orthologs with Ks >5, the median orthologous divergence between *P. glauca* and *C. japonica* was Ks = 0.78. In contrast, their most recent WGDs occurred at median Ks = 0.35 and 0.24, respectively (Fig. 3), much later than the divergence of their lineages. Orthologous divergence and phylogenomic approaches both support independent WGDs in Pinaceae and cupressophytes. Consistent with this interpretation is an absence of evidence for these WGDs in Araucariaceae (fig. S3). Overall, these results are consistent with previous analyses of chromosomes and genome sizes that hypothesized no paleopolyploidy in Araucariaceae, but likely ancient WGD in Pinaceae (19, 21).

**DISCUSSION**

In contrast to the recently published study of the Norway spruce genome (24), our analyses find evidence for at least two independent WGDs in the ancestry of major conifer clades. Why did analyses of the spruce genome not recover similar evidence of this WGD? Visual
ences in turnover rates of genome content likely contribute more to
gynosperm genome evolution also holds true for gymnosperms; differ-
sperm genomes indicate that the degree of synteny conservation
Araucariaceae (\textit{Li et al}. [Supplementary Fig. 2.6 of Nystedt
fection events have played a role in generating some of the largest
conifer genome evolution clearly did involve WGDs, and genome dup-
conservation. Nonetheless, our analyses do not con-
medians peaks for these plots are highlighted. Analyses
ieder and loss of genes \cite{33}. If most fractionation and genome rearrange-
emissions occur quickly after polyploidy, descendant polyploids may also
mogenomes may also inherit a largely common synteny \cite{34, 35}. The lack of reciprocal ge-
arrangement events following WGDs, such as in Poaceae \cite{36},
would also reduce syntenic diversity in descendant lineages. For deca-
des, the broad ancestry of polyploidy in the flowering plants was un-
dected in linkage mapping studies. Thus, relatively conserved
synteny, especially from linkage map data, is not evidence against a
paleopolyploidy in Pinaceae.

One of the most intriguing evolutionary questions raised by our
analyses is, why are there so few polyploid species among extant con-
ifers and other gymnosperms? Our analyses indicate that polyploid
speciation contributed to their diversity. Perhaps these WGDs thrived
at a climatically favorable time for polyploid species, as was proposed
to explain the apparent clustering of angiosperm WGDs near the K-Pg
mass extinction event \cite{37}. Based on our phylogenetic placements of
WGDs and existing estimates for the ages of gymnosperm lineages
\cite{38}, the conifer WGDs occurred ca. 210 to 275 million years ago (Cu-
pressaceae + Taxaceae) and ca. 200 to 342 million years ago (Pinaceae).
Many major events in Earth’s history occurred during this time frame,
including Earth’s most severe mass extinction event, the Permain-Triassic
extinction. Did polyploid conifers survive the end-Permain event bet-
ter than did their diploid contemporaries? Given that many of these
conifer clades originated during this period, these WGDs may have
uniquely contributed to the morphological and biological diversity of
these lineages. Polyploidy may differentially influence the evolution of
dosage-sensitive genes and pathways \cite{16, 39–41} or generate novelty by sub-
- or neofunctionalization \cite{42}. Examining further data sets to more
precisely pinpoint these WGDs in the conifer phylogeny and to explore
the effects of duplication on specific gene families will be critical to
further answer how polyploidy has contributed to conifer evolution.

**MATERIALS AND METHODS**

**Sampling and sequencing**

Leaf material of \textit{O. petiolatum} (PRJNA257107), \textit{G. gnemon} (PRJNA283231),
and \textit{E. frustillata} (PRJNA283230) was collected in liquid nitrogen from the
University of British Columbia (UBC) Botanical Gardens and Greenhouse and then stored in a −80°C freezer (table S1). We extracted
total RNA using the TRIzol reagent (Invitrogen)/RNasey (Qiagen) ap-
proach as described by Lai \textit{et al}. \cite{43}. For 454 sequencing (454 Life
Sciences), we used modified oligo-dT primers for complementary DNA
cDNA) synthesis to reduce the length of mononucleotide runs associated
with the polyadenylate [poly(A)] tail of mRNA. We used a “broken chain”
short oligo-dT primer to prime the poly(A) tail of mRNA during first-
strand cDNA synthesis \cite{44}. cDNA was amplified and normalized with the
TRIMMER-DIRECT cDNA Normalization Kit. After normalization, we
fragmented the cDNA to 500–800 base pair fragments by either
sonication or nebulization and removed small fragments through size
selection using AMPure SPRI beads (Angencourt). Then, the fragmented
ends were polished and ligated with adaptors. The optimal ligation products
were selectively amplified and subjected to two rounds of size selection by
gel electrophoresis and AMPure SPRI bead purification \cite{45}. Normal-
ized cDNA was prepared for sequencing following the standard genomic
DNA shotgun protocol recommended by 454 Life Sciences.
Additional data sets were downloaded from the GenBank Sequence Read Archive (SRA) (table S1). These included Sanger and Illumina data from 22 species. Data sets were selected to provide broad phylogenetic coverage of the gymnosperms. We also obtained the annotated coding DNA sequences of Amborella trichopoda (46) and Selaginella moellendorfii (47) from Phytozome (www.phytozome.net/).

Transcriptome assembly
Raw read quality filtering and trimming were performed by SnoWhite (48) before assembly. Three different assembly strategies were used for our three different data types. Sanger expressed sequence tags (EST) were cleaned using the SeqClean pipeline and assembled using TGICL. For 454 data, we used a combination of MIRA and CAP3 to assemble contigs. We used MIRA version 3.2.1 (49) using the “accurate.est. denovo.454” assembly mode. Because MIRA may split high-coverage contigs into multiple contigs, we used CAP3 at 94% identity to further assemble the MIRA contigs and singletons (50). SOAPdenovo-Trans (51) was used to assemble Illumina sequenced transcriptomes using a k-mer of about $\frac{7}{3}$ s read length. All other parameters were set to default. Assembly statistics for the 26 assemblies are given in table S1.

Age distribution of paralogs
For each species data set, we used our DupPipe pipeline to construct gene families and estimate the age of gene duplications (7, 22, 23, 47, 52). Translations and reading frames were estimated by Genevise alignment to the best hit protein from a collection of proteins from 25 plant genomes on Phytozome. As in other DupPipe runs, we used protein-guided DNA alignments to align our nucleic acids while maintaining the reading frame. For each node in our gene family phylogenies, we estimated synonymous divergence (Ks) using PAML with the F3X4 model (53). Summary plots of the age distribution of gene duplications were evaluated for each gymnosperm species for peaks of gene duplication as evidence of ancient WGDs. Taxa with peaks suggesting ancient WGDs were further analyzed using a multispecies approach (described below) to assess what fraction of gene families show a shared gene duplication and simultaneously place potential WGDs in phylogenetic context.

Estimating the orthologous divergence of Pinaceae and Cupressaceae
To estimate the average ortholog divergence of conifer taxa and compare it to observed paleopolyploidy peaks, we used our previously described RBH Ortholog pipeline (22). Briefly, we identified orthologs as reciprocal best blast hits in the transcriptomes of P. glauca (Pinaceae) and C. japonica (Cupressaceae). Using protein-guided DNA alignments, we estimated the pairwise synonymous (Ks) divergence for each pair of orthologs using PAML with the F3X4 model (53). We plotted the distribution of ortholog divergences and compared the median divergence against the synonymous divergence of paralogs from inferred WGDs in these lineages.

Inference of gene family phylogenies
Each transcriptome was translated into amino acid sequences using the TransPipe pipeline (22). We performed reciprocal protein BLAST (blastp) searches of selected transcriptomes with an e-value of $10^{-5}$ as a cutoff. Gene families were clustered from these BLAST results using OrthoMCL v2.0 with default parameters (54). Using a custom perl script, we filtered for gene families that contained at least one gene copy from each taxon and discarded the remaining OrthoMCL clusters. SATé was used for automatic alignment and phylogeny reconstruction of gene families (55). For each gene family phylogeny, we ran SATé until five iterations without an improvement in score using a centroid breaking strategy. MAFFT was used for alignments (56), Opal for mergers (57), and RAXML for tree estimation (58). The best SATé tree for each gene family was used to infer and locate WGDs by our MAPS algorithm.

Multi-taxon Paleopolyploidy Search (MAPS)
To infer and locate ancient WGDs in our data sets, we developed a gene tree sorting and counting algorithm, MAPS. This algorithm uses a given species tree to filter for subtrees within complex gene trees consistent with relationships at each node in the species tree. For each node of the species tree, MAPS parses the species tree into subtrees with a sister species and an outgroup, for example, ((A,B),C). MAPS iteratively searches for each of these subtrees in the gene tree and will ignore subtrees that do not have the expected relationship. In-paralogs are collapsed by MAPS to simplify the search. We filter for these subtrees, rather than filtering on entire topologies, because ancient WGDs may yield phylogenies with many nested and/or orthologous clades. Filtering for a simple gene tree that matches the species tree would eliminate many of the trees that support WGDs. By filtering for subtrees of the species tree, MAPS captures the evidence for polyploidy in complex gene family topologies. Using this filtered set of gene trees, MAPS records the number of subtrees that support a gene duplication at a particular node in the species tree (fig. S1). To infer and locate a potential WGD in the species tree, we plot the percentage of gene duplications shared by descendant taxa by node (fig. S2). A WGD will produce a large burst of shared duplications across taxa and gene trees. This burst of duplication will appear as an increase in the percentage of shared gene duplications in our MAPS analyses.

To evaluate if a WGD occurred before the divergence of taxa A and B, MAPS requires gene trees with at least a sister group A and B and an outgroup C (fig. S1). The basic algorithm of MAPS has two steps. In step 1, MAPS collapses in-paralogs that evolved after the divergence of A and B to a single copy in each gene tree (fig. S1). In step 2, MAPS counts subtrees from all gene trees that are consistent with a duplication event in the MRCA of A and B. In our ABC example, subtrees with a topology consistent with duplication before the divergence of A and B (for example, (((A,B),(A,B)),C)) will be recorded as a duplication at their MRCA node (fig. S1, 1.6). Additionally, subtrees with a topology consistent with duplication before the divergence of A and B followed by independent gene loss [for example, ((A,−),(A,B)),C] or (((A,B),(−,B)), C]) will also be recorded as a duplication at their MRCA node (fig. S1, 1.7 to 1.10). If gene trees do not have a topology consistent with any gene duplication among the ingroup taxa, then no duplications will be recorded at the internal nodes (fig. S1, 1.1 to 1.5). When searching for ancient WGDs in a collection of gene trees that contain more than three taxa, MAPS will repeat the same algorithm on each node of the tree (fig. S2). WGDs are inferred by searching for evidence of a large number of shared duplications at a particular node(s) of the species tree (fig. S2).

To evaluate the phylogenetic placement of the putative “seed plant” WGD, we used MAPS to analyze gene families from representatives of each vascular plant lineage (Fig. 1A and fig. S4A). We selected Araucaria angustifolia and Ginkgo biloba to represent gymnosperms because our Ks plots suggest that they only experienced the seed plant WGD. We also analyzed the Amborella genome to represent angiosperms (46). The
newly sequenced *O. petiolatum* transcriptome and the *S. moellendorffii* genome (47) were chosen to represent ferns and lycophytes, respectively.

We conducted two MAPS analyses to evaluate numbers and placements of WGDs among conifers (Fig. 1, B and C, and fig. S4, B and C). Two analyses were conducted instead of one because the MAPS algorithm works best with simple, ladderized species trees. To maximize the numbers of gene trees in the MAPS analysis and have good coverage of the Pinaceae phylogeny, we selected the transcriptomes of *Pinus monticola*, *Larix gmelinii*, and *Cedrus atlantica* to represent Pinaceae. We also selected *Taxus mairei* to represent the cupressophytes. Likewise, we chose *T. mairei*, *Cephalotaxus hainanensis*, and *C. japonica* to represent cycadophytes and *P. monticola* to represent Pinaceae. For both Pinaceae and cupressophyte analyses, the transcriptomes of *G. biloba* and *O. petiolatum* as well as the *S. moellendorffii* genome were selected as outgroups.

**SUPPLEMENTARY MATERIALS**

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/full/1/10/e1501084/DC1.

Fig. S1. Example topologies processed by MAPS to identify a gene duplication (red star) or not (black dot) in a given gene family phylogeny.

Fig. S2. Example MAPS summary results for a four-taxon phylogeny.

Fig. S3. Histograms of the age distribution of gene duplications from 24 gymnosperm genome datasets.

Supplementary material for this article is available at http://advances.sciencemag.org/cgi/content/101/4/552/DC1.

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