A physical map for the *Amborella trichopoda* genome sheds light on the evolution of angiosperm genome structure

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Abstract

**Background:** Recent phylogenetic analyses have identified *Amborella trichopoda*, an understory tree species endemic to the forests of New Caledonia, as sister to a clade including all other known flowering plant species. The *Amborella* genome is a unique reference for understanding the evolution of angiosperm genomes because it can serve as an outgroup to root comparative analyses. A physical map, BAC end sequences and sample shotgun sequences provide a first view of the 870 Mbp *Amborella* genome.

**Results:** Analysis of *Amborella* BAC ends sequenced from each contig suggests that the density of long terminal repeat retrotransposons is negatively correlated with that of protein coding genes. Syntenic, presumably ancestral, gene blocks were identified in comparisons of the *Amborella* BAC contigs and the sequenced *Arabidopsis thaliana*, *Populus trichocarpa*, *Vitis vinifera* and *Oryza sativa* genomes. Parsimony mapping of the loss of synteny corroborates previous analyses suggesting that the rate of structural change has been more rapid on lineages leading to *Arabidopsis* and *Oryza* compared with lineages leading to *Populus* and *Vitis*. The gamma paleohexaploidy event identified in the *Arabidopsis*, *Populus* and *Vitis* genomes is shown to have occurred after the divergence of all other known angiosperms from the lineage leading to *Amborella*.

**Conclusions:** When placed in the context of a physical map, BAC end sequences representing just 5.4% of the *Amborella* genome have facilitated reconstruction of gene blocks that existed in the last common ancestor of all flowering plants. The *Amborella* genome is an invaluable reference for inferences concerning the ancestral angiosperm and subsequent genome evolution.

**Background**

The origin and rapid diversification of the angiosperms (flowering plants) were pivotal events in the evolutionary history of Earth’s biota. Over the past 130 to 150 million years angiosperms have diversified to include approximately 350,000 species occupying nearly all habitable terrestrial and many aquatic environments. Angiosperms generate the vast majority of human food either directly or indirectly as animal feed, and they account for a huge proportion of land-based photosynthesis and carbon sequestration. Comparative analyses of genome sequences and gene function for a growing number of species are shedding light on how gene and genome duplications have contributed to the diversification within major flowering plant lineages (for example, Rosidae, Asteridae, Monocotyledoneae [1]), but elucidation of the genetic and genomic processes underlying the key innovations associated with the origin of flowering plants (for example, typically
b bisexual flowers, endosperm formation, double fertilization, ovules with two integuments, seed development within the carpel) requires comparisons between lineages that diverged from the last common ancestor of all extant angiosperms [2,3].

Recent phylogenetic analyses have identified *Amborella trichopoda*, an understory tree or shrub species endemic to the forests of New Caledonia, as the sister species to all other extant angiosperms [4-8]. *Amborella* is no more ‘ancient’ or ‘primitive’ than any other extant flowering plant species, but comparisons between *Amborella* and other angiosperms are allowing researchers to triangulate on characteristics of their last common ancestor. Using a similar approach, researchers have used the complete genome sequence of platypus, *Ornithorhyncus anatinus*, representing the sister group of all other extant mammals, to elucidate mammalian genome evolution [9].

Previous comparisons of transcriptome content [10], gene expression patterns [11-13], and gene function [14,15] between *Amborella* and other flowering plant species have suggested that much of the floral development program that has been characterized in *Arabidopsis*, snapdragon and maize existed in the last common ancestor of extant angiosperms. While gene duplications in the MADS-box transcription factor family likely contributed to the earliest floral development regulatory networks [11,12,16-19], it is not clear whether these were single gene duplications or the product of polyploidization. Genome duplications have occurred repeatedly throughout angiosperm history [20-23] but there is uncertainty in the timing of polyploidy events relative to the origin of the angiosperms and important innovations in flowering plant history [24].

Here we describe a BAC-based draft physical map for *A. trichopoda* and use BAC end sequences (BESs) to compare the structure of the *Amborella* genome to representative eudicot (*Vitis, Populus* and *Arabidopsis*) and grass (*Oryza*) genomes. Comparative analyses of sequences for two large contiguous regions (487.3 and 629.7 kb in the *Amborella* genome) were also performed. In addition we use a large transcriptome assembly to identify BAC ends matching protein-coding sequences [25]. Our aim here is to begin to investigate whether regions of these genomes have remained syntenic throughout angiosperm history, and determine whether ancient genome duplications discovered in eudicot and grass genomes [26-29] occurred before or after the divergence of these lineages from the *Amborella* lineage. In addition, the physical map and sequence analyses establish a framework for future studies of all flowering plant genomes, including the *Amborella* genome itself.

**Results and discussion**

**BAC library and physical map**

The structure and composition of the 870 Mbp/C [30] *A. trichopoda* genome was investigated through physical mapping of clones from a 5.2 x coverage BAC library. The library was constructed after partial digest of high-molecular-weight DNA with HindIII. The library, which comprises 36,684 BAC clones with an estimated average insert size of 123 kb, is available through the Arizona Genomics Institute [31]. The BAC library was double spotted in high density onto Hybond N+ filters. All 36,684 clones were end-sequenced, and a physical map was constructed after high information content fingerprinting (HICF) [32,33]. A total of 32,719 fingerprinted BACs was assembled into 3,106 contigs and 1,356 singletons using the program FPC version 7.2 [34].

The quality of the physical map was assessed by screening the arrayed library with probes developed for *Amborella* homologs for eight genes that have been found to be single-copy in sequenced plant genomes [35,36]. Probes derived from *Amborella* cDNA clones or PCR amplicons were putative homologs of following single-copy *Arabidopsis* genes: *ASD* (At1g14810), *DWARF1* (At3g19820), *GIANTTEA* (At1g22770), *LEAFY* (At5g61850), a diene-lactone hydrolase gene (At2g32520), a cytochrome-C- oxidase-related gene (At4g37830), *EIF3K* (At4g33250) and a hypothetical protein-coding gene with strong similarity to *rice* gene Os02g0593400 (At5g63135). All verified positive clones mapped to the same FPC contig for six of the eight probes (Figure S1 in Additional file 1). Positive clones for the *EIF3K* and the hypothetical protein-coding gene probes were each distributed between two FPC contigs and inspection of the HICF bands for these contigs suggests that the genes have been duplicated in the *Amborella* lineage. In accordance with the expected library coverage, the single copy nuclear gene probes hybridized to 3 to 13 clones (mean 6.9).

The correlation between HICF bands and the number of BACs included in each FPC contig was 0.655 for all contigs and 0.917 after removing two contigs derived from the chloroplast and mitochondrial genomes and one contig composed largely of repetitive elements (Figure S2 in Additional file 1). We used a calibration of average insert size (123 kb) over the average number of HICF bands per BAC clone to obtain a rough estimate of FPC contig lengths. Of 77 FPC contigs with 39 or more BACs (not including the contigs with the plastome and repetitive elements), estimated lengths ranged from 308 to 1,429 kb.

BAC end sequencing was performed on all fingerprinted BACs producing 69,466 Sanger reads with an average length of 695 bp after quality and vector trimming. This corresponds to 48.25 Mbp, or roughly 5.4% of the *Amborella* genome. BESs were related to the
physical map and used to identify regions of synteny between regions of the Amborella genome and the sequenced Arabidopsis, Populus, Vitis (grape), and Oryza (rice) genomes (see below). In addition, end sequences were used to verify the identity of the three excluded FPC contigs described above. All BESs mapping at least 100 bp apart on the plastid genome [37] were found in the same FPC contig. This contig included just 532 BACs, indicating very low (1.6%) plastid DNA contamination.

Characterization of repeats in BAC end and shotgun sequences

Repet composition and frequency in the Amborella genome were characterized through analysis of the BAC end and whole genome sequence sequences. Reads were first compared with sequences in Repbase (v.15.08) [38] using BLASTN [39]. In order to minimize the effect of divergence between Amborella genera and homologous repeats from other species, we used relaxed BLASTN settings (-q -4 -r 5) to accommodate an estimated 160 million years of sequence divergence since the last common ancestor of extant flowering plants [8,40-42] while maintaining rigorous support for significant hits (E-value threshold was set at 1e-10). All BAC end sequences without significant hits were then compared with the non-redundant protein database in GenBank using BLASTX and an E-value threshold of e-5. Finally, the remaining sequences without matches in Repbase or the GenBank nr database were compared with sequences that did have matches in either database using BLASTN with an E-value threshold of 1.0e-10. We report results both excluding these `internal’ BLAST searches and including them (I). Together these results provide estimates of transposable element (TE) content based on conservative and more comprehensive (and possibly more permissive; I) search strategies.

With the more comprehensive strategy (I), slightly more than half of all the Amborella BESs matched known TE sequences. Not surprisingly, the most highly represented TE class was long terminal repeat (LTR) retrotransposons, accounting for 7.65% (I: 30.01%) of all BESs and 57.5% (I: 56.58%) of all those with hits to Repbase. Hits to Ty1-copia type sequences were slightly more common (3.11%; I: 13.79%) than matches to Ty3-gypsy-like LTRs (3.50%; I: 12.09%); the remaining LTR retrotransposon matches (1.04%; I: 4.13%) were not classified. LINEs also represented a significant fraction of Amborella BAC ends: 2.70% (I: 11.60%) of the total, 19.98% of all the repeats (I: 22.22%). This is noteworthy because LINEs are usually significantly less numerous than LTR retrotransposons in plant genomes [43-47] with some notable exceptions, such as the element del2 in Lilium speciosum [48]. The complete set of DNA TE-related BESs accounts for just 1.63% (I: 4.51%) of the total, and the most represented classes are those of hAT and MuDR elements: 0.92% (I: 2.41%) and 0.49% (I: 1.04%) of the total BESs, respectively. Results from the same analyses replicated on the set of 2,695 random sheared Sanger sequences (Table 1) and 648,519 454 reads (Table S1 in Additional file 1) are generally in very good agreement with those obtained using BES data.

A de novo search for novel miniature inverted repeat transposable elements (MITEs) overlooked by the similarity search approach was carried out using the pipeline MUST [49]. The most abundant candidates identified by the pipeline were manually inspected to confirm features typical of MITEs, such as small size, terminal inverted repeats, high A+T nucleotide content and target site duplications. Three putative high-copy MITEs were identified. All of these were small elements (174 to 500 bp) with terminal inverted repeats, target site duplications, and A+T content greater than 65% (Figure S3 in Additional file 1). Repeat copy numbers estimated from the BESs and random sheared sequences were extrapolated to obtain genome-wide estimates using the procedure developed by Hawkins et al. [50]. Copy number ranges from 3,300 copies for MITE_2 to 17,000 copies for MITE_1. The estimates inferred from BESs were generally consistent with those calculated for random sheared reads (with the possible exception of MITE_3; Table 2).

The conserved reverse transcriptase domains of LTR retrotransposons and LINEs were collected and used to estimate maximum likelihood trees (Figure 1). In the case of LTR retroelements, the trees indicate substitution rate heterogeneity (that is, variation in root-to-tip distances) and no evidence for recent retrotranspositional bursts of single families (that is, short terminal branches). In the case of LINEs, the phylogenetic tree displays very long branches suggestive of an ancient diversification or very rapid substitution rates. As has been described for other plants [51], Amborella LINEs exhibit high sequence divergence and extreme heterogeneity.

The Amborella BESs were also searched for microsatellites (that is, simple sequence repeats (SSRs)); for comparison, the search was also conducted on the Amborella random sheared reads and on BESs (from other HindIII BAC libraries) from Glycine (soybean) and Oryza rufipogon. In comparison to the other two species, Amborella shows a higher frequency of SSRs, particularly mono- and dinucleotide repeats, with a particularly high frequency of ‘AG’ dinucleotide microsatellites. The results of SSR analysis in BESs were confirmed by those obtained from the randomly sheared Amborella sequences (Table 3).

Repeat profiles in the shotgun sequences were also assessed using Tallymer to characterize K-mer frequencies [52]. The Amborella K-mer frequency profiles were compared with those of Arabidopsis thaliana, Oryza sativa (rice), Sorghum bicolor and Zea mays (maize).
While the *Amborella* genome size is closest to Sorghum's (870 and 740 Mbp/C, respectively), its K-mer frequency profiles were more similar to those of *Arabidopsis* and rice, with much smaller genome sizes (157 and 490 Mbp/C, respectively [53]) (Figure 2).

Distribution of BESs with matches to protein-coding regions of reference genomes

All BESs and shotgun sequences were compared to the GenBank nr database using BLASTX [39] with an e-value threshold of 1e-5. After the removal of sequences similar to TEs, the overall frequencies of sequences finding matches in the protein database were 11.9% and 8.05% for the BES and Sanger shotgun sequences, respectively. For BESs from FPC contigs with ten or more BACs, we found a negative correlation between the frequencies of BESs matching protein-coding genes and LTR retrotransposons (r = -0.423, P < 0.0001). As has been described for other genomes [54-56], gene density seems to be negatively correlated with retrotransposon density in the *Amborella* genome.

Identification of syntenic blocks between *Amborella*, *Arabidopsis*, rice, poplar and grape

Taking advantage of the availability of a phase I physical map assembly, we mapped the *Amborella* contigs onto the genomes of *A. thaliana*, *Populus trichocarpa*, *Vitis vinifera*, and *O. sativa*. We focused on the 77 largest contigs with at least 39 clones. BLAST analyses of BESs were done within the context of their linkages within FPC contigs. All of the contig BESs classified as repeats (see above) were discarded. Those remaining were compared against the four reference genomes. Because of the large evolutionary time that separates *Amborella* from the other four sequenced genomes [41,42,57], the comparisons were carried out at the protein level using tBLASTX; only the best hits were taken into account. *Amborella* FPC contigs were considered for further analyses if at least two BESs had matches with bit scores greater than 80 (typically a maximum e-value of 1.0E-20 over 100 amino acidic residues) to loci separated by less than 500 kb within one of the four genomes being compared. Positive matches were used as anchors to circumscribe 4-Mbp tracts within the genome.

| DNA TEs | Absolute number in BESs | Percentage BESs | Percentage repeats in BESs | Absolute number in SGSs | Percentage SGSs | Percentage repeats in SGSs |
|---------|-------------------------|-----------------|---------------------------|------------------------|----------------|---------------------------|
| hAT     | 642 (1,671)             | 0.92 (2.41)     | 6.84 (4.61)               | 20 (41)                | 0.74 (1.52)   | 5.73 (2.94)               |
| MuDR    | 343 (724)               | 0.49 (1.04)     | 3.65 (2.00)               | 7 (30)                 | 0.26 (1.11)   | 2.00 (2.15)               |
| CACTA   | 27 (75)                 | 0.04 (0.11)     | 0.29 (0.21)               | 0 (4)                  | 0 (0.15)      | 0 (0.29)                  |
| Helitrons | 12 (69)               | 0.02 (0.10)     | 0.13 (0.19)               | 0 (3)                  | 0 (0.11)      | 0 (0.22)                  |
| Other   | 108 (595)               | 0.15 (0.86)     | 1.15 (1.64)               | 1 (24)                 | 0.04 (0.89)   | 0.29 (1.72)               |
| Total   | 1,132 (3,134)           | 1.63 (4.51)     | 12.06 (8.64)              | 28 (102)               | 1.04 (3.78)   | 8.02 (7.31)               |

| Retrotransposons | Absolute number in BESs | Percentage BESs | Percentage repeats in BESs | Absolute number in SGSs | Percentage SGSs | Percentage repeats in SGSs |
|------------------|-------------------------|-----------------|---------------------------|------------------------|----------------|---------------------------|
| LTR Ty1-copia    | 2,162 (9,578)           | 3.11 (13.79)    | 23.02 (26.42)             | 64 (314)               | 2.37 (11.65) | 18.34 (22.51)             |
| LTR Ty3-gypsy    | 2,431 (8,395)           | 3.50 (12.09)    | 25.89 (23.15)             | 129 (377)              | 4.78 (13.98) | 36.96 (27.03)             |
| LTR not classified | 720 (2,686)            | 1.04 (4.13)     | 7.67 (7.91)               | 51 (139)               | 1.89 (5.16)  | 14.61 (0.96)              |
| LINEs            | 1,876 (8,055)           | 2.70 (11.60)    | 19.98 (22.22)             | 55 (294)               | 2.04 (10.91) | 15.76 (21.08)             |
| SINEs            | 11 (183)                | 0.02 (0.26)     | 0.12 (0.50)               | 0 (4)                  | 0 (0.15)     | 0 (0.29)                  |
| Retro not classified | 1,058 (4,046)         | 1.52 (5.82)     | 11.27 (11.16)             | 23 (165)               | 0.85 (6.12)  | 6.59 (11.83)              |
| Total            | 8,258 (33,125)          | 11.89 (47.69)   | 87.94 (91.36)             | 321 (1,293)            | 11.91 (47.96) | 91.98 (92.69)             |

| Total | 9,390 (36,259) | 13.52 (52.20) | 100 (100) | 349 (1,395) | 12.95 (51.74) | 100 (100) |

Results in parentheses include internal BlastN searches. Repbase v.15.08 was used [38]. SINE, short interspersed element; SGS, Sanger shot gun sequence.

| Table 2 Putatively high-copy MITEs identified in the BESs and Sanger shot gun sequences using MUST pipeline |
|-------------------------------------------------|-----------------|-----------------|-----------------|-----------------|-----------------|-----------------|
| Length | Inverted repeat length | BES hits | Copy number estimate | SGS hits | Copy number estimate | AT% |
|--------|------------------------|----------|---------------------|----------|---------------------|-----|
| MITE_1 | 358                    | 26       | 542                 | ~17,000  | 18                  | ~17,200 68.80 |
| MITE_2 | 190                    | 19       | 140                 | ~3,300   | 8                   | ~3,100 68.70  |
| MITE_3 | 516                    | 47       | 394                 | ~17,000  | 8                   | ~11,300 75.20  |

Copy number estimates based on procedure of Hawkins et al. [50]. SGS, Sanger shot gun sequence.
reference genomes and a second, more focused tBLASTX search was performed comparing the BESs with these regions. An e-value threshold of 1.0E-4 was used for the second set of tBLASTX searches and all significant hits were used to identify syntenic regions. We considered a contig as anchored if the contig had at least four positive hits (e-value lower than 1.0e-4) to at least three distinct genes.

Non-repetitive BESs were also compared to a database of 246,196 Amborella cDNA unigene assemblies with lengths greater than 100 bp. These cDNAs were derived from comprehensive sequencing of nine cDNA libraries (Table 4) [25]. Sixty-six percent of the non-repetitive BESs matched cDNA sequences in BLASTN searches with an e-value cutoff of 1.0e-10.

Using the search strategy described above, 29 large Amborella BAC contigs (>39 BAC clones) showed

### Table 3 Simple sequence repeats identified in BESs and Sanger shot gun sequences

| Repeat | Amborella (BES)* | Amborella (RS)* | Soybean* | Oryza rufipogon* |
|--------|------------------|-----------------|----------|-----------------|
| Mono   | 149.66           | 152.89          | 72.74    | 50.79           |
| Di     | 225.03           | 211.00          | 77.89    | 63.94           |
| Tri    | 72.49            | 78.96           | 110.01   | 144.06          |
| Tetra  | 89.88            | 90.70           | 100.67   | 102.25          |
| Penta  | 74.85            | 89.73           | 64.54    | 56.00           |
| Total  | 611.92           | 623.28          | 425.85   | 417.04          |

*Values are presence per million base pairs. RS, Random sheared

Figure 1 Maximum likelihood trees for reverse transcriptase genes classified as Copia-type and Gypsy-type LTR and LINE elements. (a) Copia-type; (b) Gypsy-type LTRs; (c) Gypsy-type LINEs. The maximum likelihood trees show rate heterogeneity and no recent expansive radiations (that is, short terminal branches). Reverse transcriptase sequences were mined from BAC end sequence set.

![Figure 1](http://genomebiology.com/2011/12/5/R48)

Figure 2 K-mer analyses of Sanger shotgun sequences reveal low frequencies of short repeats in the Amborella genome relative to the sorghum and maize genomes.

![Figure 2](http://genomebiology.com/2011/12/5/R48)
synteny with at least one of the four sequenced genomes, and nine of these showed synteny with at least one region in all four genomes. All BESs mapping to these syntenic regions also exhibited significant matches to the sequences in the Amborella cDNA assembly (Table 4; Table S2 in Additional file 1). Whereas 25 of these Amborella BAC contigs mapped to at least one tract in the Vitis genome, 15, 16, and 24 contigs were found to be syntenic with one or more tracts in the Oryza, Arabidopsis, and Populus genomes, respectively (Table S2 in Additional file 1). These results provide a novel, albeit coarse, first view of the ancestral genome for all flowering plants and the timing of rearrangements and other structural changes (for example, genome duplications, fractionation, chromosomal fissions and fusions) that have reduced synteny between the monocot and eudicot genomes analyzed here (Figure 3). Parsimony mapping of synteny loss onto a phylogeny consisting of Amborella and the other four species indicates variation in rates of change in genome structure. In agreement with previous studies [29,45], Vitis seems to have been the most stable of the sequenced genomes, and the rate of change slowed in the lineage leading to Populus following divergence from the lineage leading to Arabidopsis (Figure 3).

Paleopolyploidy in angiosperm genomes
Paleopolyploidy events have been well characterized in all four sequenced genomes analyzed here [29,45,58-60], and the syntenic Amborella FPC contigs described above often match multiple regions in these genomes. The most ancient of these paleopolyploidy events is the so-called g tripllication that has been inferred to have occurred before the divergence of the Asteridae (represented by tomato, Solanum lycopersicon) and the Rosidae, including Vitis, Populus and Arabidopsis [29]. Given the very incomplete view of the Amborella genome that is available in the BES data, we are not able to assess synteny between Amborella FPC contigs. Nevertheless, comparisons between the Amborella contigs and sets of syntenic blocks in the Vitis genome indicate that the γ tripllication most likely occurred sometime after the divergence of all other angiosperms from the lineage leading to Amborella.

All BESs were compared to all annotated protein-coding genes in the Vitis genome placed within the context of the pre-tripllication ancestral gene blocks and post-tripllication syntenic segments identified by Tang et al. [29]. A total of 328 Amborella FPC contigs had between two and eight genes with significant best BLASTX matches (e-values ≤1.0E-6) to Vitis genes corresponding to pre-tripllication gene blocks in the ancestral genome. In most of these cases (199 of 328; Additional file 2), best hits were distributed between two or three homeologous (that is, post-tripllication) syntenic Vitis genome segments. Of the remaining 129 Amborella FPC contigs with BESs showing significant BLASTX hits to a single Vitis subgenome (that is, single copy of a triplicated ancestral block), most (113) included just 2 genes mapping to the ancestral Vitis gene blocks (14 including 3 genes, and 2 including 4 genes) (Additional file 2). All 21

Table 4 Statistics for cDNA sequences included in multi-library transcriptome assembly of 246,196 unigenes with lengths greater than 100 bp

| Tissue - library name                  | Sequencing method | Number of reads | Unscreened reads | Total passing bases (MB) |
|---------------------------------------|-------------------|----------------|------------------|--------------------------|
| Apical meristem - Atr12               | 454 FLX Titanium  | 794,746        | 688,305          | 201.90                   |
| Male flowers - Atr15                  | 454 FLX Titanium  | 277,023        | 255,213          | 73.49                    |
| Old leaves - Atr14                    | 454 FLX Titanium  | 280,097        | 260,563          | 73.49                    |
| Old stem - Atr13                      | 454 FLX Titanium  | 259,431        | 238,156          | 68.70                    |
| Pre-meiotic female flower buds - Atr10| 454 FLX GS        | 895,000        | 812,325          | 176.97                   |
| Pre-meiotic female flower bud - Atr02 | Sanger            | 13,263         | 13,141           | 7.17                     |
| Pre-meiotic male flower bud - Atr01   | Sanger            | 25,343         | 25,006           | 14.17                    |
| Root - Atr11                          | 454 FLX GS        | 324,070        | 300,275          | 64.88                    |
| Stem - Atr16                          | 454 FLX Titanium  | 410,098        | 388,436          | 120.03                   |

Assemblies and raw data can be downloaded from the Ancestral Angiosperm Genome Project website [25]. A BLAST portal for the assembly is also available at the project website.

Figure 3 Variation in rates of structural evolution evident in parsimony mapping of losses of synteny with 29 gene blocks inferred for the last common ancestor of all extant flowering plant lineages
FPC contigs with best BLASTX matches to five or more genes within the ancestral *Vitis* blocks were distributed among two or three post-triplication subgenomes. Complete sequences for the *Amborella* BAC contigs may reveal more even distribution of segments among *Vitis* subgenomes, but the results described here suggest that triplcation, fractionation and divergence of homoeologous segments in the *Vitis* genome postdate the divergence between lineages leading to *Vitis* and *Amborella* (that is, the last common ancestor of all extant angiosperms).

**Analysis of complete sequences for two *Amborella* BAC contigs**

Two of the larger (approximately 500 kb) BAC contigs (IDs 431 and 1003) mapping to multiple segments in all four sequenced reference genomes were identified for further investigation. A minimum tiling path was constructed for each contig, and florescence *in situ* hybridizations were performed to verify that the BACs mapped to a single contiguous region in the *Amborella* genome (Figure 4). Each BAC in the tiling paths was subcloned and sequenced to 8× coverage on an ABI 3730xl sequencer. Gaps were closed for each scaffold, and contiguous 487,318 and 629,678 bp phase II sequences were assembled for contigs 431 and 1003, respectively.

The DAWGPAWS suite of scripts was used to organize *ab initio* gene predictions, BLAST results and the output of repeat identification tools [61,62]. *Ab initio* gene predictions were generated using FGENESH [63], AUGUSTUS [64], SNAP [65], GeneID [66] and GenScan [67]. In addition, *Amborella* EST sequences produced by the 454 Titanium platform (2,943,273 reads; total read size of approximately 776 Mbp; average read length of 263.60 bp) and Sanger sequencing (38,147 reads; total read size of approximately 21.3 Mbp; average read length of 559.57 bp) were splice-aligned to the contigs using GMAP (Genomic Mapping and Alignment Program) [68] with the PASA (Program to Assemble Spliced Alignments) genome annotation tool [69]. All predictions were manually compared with BLASTX results against gene annotations from *Arabidopsis* [70], *Vitis* [45], Z. mays [56], *Medicago* [71], *Oryza* [72,73], and *Sorghum* [55] as well as tBLASTx results against the *Amborella* transcript assemblies. GBrowse views of gene annotations and BLAST results for each contig are available at the Ancestral Angiosperm Genome Project website [25].

Rigorous assessments of synteny between these *Amborella* contigs and the aforementioned four angiosperm genomes were performed using LASTZ [74,75]. Dotplots comparing the *Amborella* contigs and the *Vitis*...
At least two genome duplications (ρ and σ) have been inferred to have occurred within the monocot lineage leading to rice since divergence of monocots and eudicots [28]. These duplications were evident in comparisons with both Amborella contigs. Regions of contig 1003 were found to be syntenic with portions of rice chromosomes 2 and 4 derived from the ρ duplication and a portion of chromosome 10 (Figure 5) that is related to these two regions through the earlier σ duplication [28]. The LASTZ analysis of contig 431 revealed synteny with seven regions in the rice genome (Figure 6) and one of the ‘putative ancestral regions’ (PAR 17) characterized by Tang et al. [28]. These PARs were defined as regions of synteny between the rice and Vitis genomes. Phylogenetic analyses of genes in Amborella contig 431 and syntenic regions of the rice and Vitis genomes may elucidate the timing of the γ triplication and genome duplications.
evident in synteny analyses of the rice genome relative to the divergence of monocots and eudicots.

**Phylogenetic analyses of gene families represented in sequenced Amborella contigs**

While the fractionation process has resulted in the loss of most duplicated genes following the ancient polyploidy events evident in the syntenic *Vitis* and rice segments shown in Figures 5 and 6, duplicate *Vitis* genes have been retained for homologs of three *Amborella* genes located on contig 431 (Figures 6a). These genes were used to search the PlantTribes gene family database [35]. The three gene sets identified in the synteny analysis correspond to three gene families (auxin-independent growth promoter, ceramidase and plant uncoupling mitochondrial protein) circumscribed through OrthoMCL clustering [76] of gene annotations from the available *Arabidopsis*, *Carica* (papaya), *Populus*, *Medicago* (alfalfa), *Glycine*, *Cucumis* (cucumber), *Vitis*, *Mimulus*, *Oryza*, *Sorghum*, *Selaginella* (spike moss) and *Physcomitrella* genomes. Homologous genes sampled from exemplar asterid, ranunculid, non-grass monocot and gymnosperm species were obtained from EST assembly databases [25,77,78] and were added to each gene family set. Sequences in each gene family set were aligned using MUSCLE [79], and RAxML [80] run with the GTRGAMMA substitution model was used to obtain maximum likelihood estimates of gene trees.

Inspection of the resulting gene trees shows support for the inference drawn from the BAC end sequence analysis. The γ triplation (hexaploidy event) clearly occurred after *Amborella* diverged from other extant angiosperm lineages (Figure 7). The placement of the γ triplation with respect to the divergence of monocots and eudicots or core eudicots and the Ranunculales varies among the three gene trees. This incongruence among gene trees is likely due to artifacts associated with substitution rate variation and insufficient taxon sampling. Analyses of additional gene families with broader taxon sampling will be necessary to obtain better resolution for the timing of the γ triplation with respect to the divergence of monocot, eudicots, Ranunculales (that is, ‘basal’ eudicots) and core eudicots.

**Conclusions**

*A. trichopoda* is the sister species to the large clade encompassing all other extant flowering plants. As such,
comparative analyses of *Amborella* and other flowering plants offer a uniquely informative perspective on the most recent common ancestor of all extant angiosperms. The physical map and BAC end sequences described in this study provide a low-resolution view of the *Amborella* genome. Nonetheless, these data shed light on genomic features of the last common ancestor of flowering plants. Moreover, the *Amborella* genome provides a unique reference for understanding genome evolution throughout angiosperm history. When placed in the context of the physical map, BESs representing just 5.4% of the *Amborella* genome allowed reconstruction of ancestral gene blocks in regions represented by 29 BAC contigs and inference of the timing of structural mutations that disrupted these blocks (Figure 3).

Analyses of BESs and BAC contigs also indicate that the ancient γ polyploidy event inferred from the *Arabidopsis* [58], *Carica* [81], *Populus* [60], and *Vitis* [45] genomes occurred after the *Amborella* lineage diverged from the rest of the angiosperms. Therefore, if the origin of angiosperms was associated with a genome duplication as has been hypothesized elsewhere [16,20,23], that polyploidy event predated the γ event.

### Materials and methods

#### BAC library construction

Protocols for DNA megabase preparation, library construction, picking and arraying proposed in Luo and Wing [82] were followed.

#### Fingerprinting

The SNaPshot fingerprinting technique was adopted [32] with the modifications described by Kim et al. [83]. Snapshot reactions were loaded into ABI 3730xl DNA sequencers. Analysis of data for each contig was carried out using the ABI Data Collection Program.

#### Physical map construction

Fingerprints were assembled into contigs using the program FPC version 7.2 [34]. The initial assembly was carried out using a Sulston score threshold of e-50 followed by three rounds of dequeuing at the same stringency and auto-merging of contigs at e-21.

#### BAC end extraction and sequencing

BAC DNA was extracted and end sequenced from 36,684 clones using the methods described by Ammiraju et al. [83,84]. Sequence quality assessment and trimming were carried out using the programs Phred [85] and Lucy [86].

#### Random sheared library

A random sheared library was constructed as previously described [87].

#### cDNA sequencing and assembly

Additional Sanger ESTs were generated from available male and female flower bud cDNA libraries [10] (Table 4). Libraries for 454 sequencing were constructed from the tissues listed in Table 4 using the Mint cDNA synthesis kit (Evrogen, Moscow, Russia). Total RNAs for cDNA synthesis were isolated using a combination of CTAB extraction and the RNeasy Plant Mini kit (Qiagen Valencia, CA USA) as previously described for basal angiosperms [11]. Two rounds of messenger RNA isolation were performed with the Poly(A)Purist™ mRNA Purification Kit (Ambion Inc. Austin, TX USA) according to the manufacturer’s recommendation. Contaminant DNA was removed with DNA-free™ (Ambion Inc.) and mRNA quality was verified using a Bioanalyzer (Agilent Inc. Santa Clara, CA, United States). Vector and adaptor sequences were trimmed from 454 Titanium (2,943,273 reads; total read size of approximately 776 Mbp; average read length of 263.60 bp) and Sanger sequences (38,147 reads; total read size of approximately 21.3 Mbp; average read length of 559.57 bp) using seqclean [88] and assembled using MIRA [89].

#### Similarity searches, repeat classification and contig anchoring

Similarity searches were carried out using the programs BLASTN and BLASTX [39]. BLASTN was run under relaxed settings (-q -4 -r 5) in order to accommodate the evolutionary distance between *Amborella* and the species included in the repeat databases used; the significance threshold was set at 1e-10. In the case of BLASTX searches the threshold was set at 1e-5 or 1e-4 for the BES synteny analysis. tBLASTX was used to anchor the contigs to the reference genomes (see Results for details).

#### Databases

The databases used in similarity searches were RepBase version 15.08 [38], the GenBank non-redundant (nr) database, and the *Oryza, Arabidopsis, Vitis* and *Populus* genome sequences.

#### Validation of repeat searches and MITE identification

The program MUST [49] was used for *de novo* characterization of highly repeated sequences; results were then inspected for the presence of MITE features. Inverted repeats were identified manually parsing the results of dot-plot comparisons made using the program ‘Dotter’ [90].

#### Simple sequence repeat searches

Microsatellites were identified using the program Sputnik [91]. SSR composition, length and distribution were parsed and analyzed using the tools and the strategy used by Morgante et al. [92].
Fluorescence in situ hybridization

FPC contigs were validated by hybridizing BAC DNAs to Amborella chromosome squashes. DNA was prepared for BAC mapping to the middle and both ends of BAC contigs 431 and 1003 and used to prepare fluorescently labeled BAC-FISH probes. Chromosome squashes were prepared from root tips and labeled BAC-FISH probes were prepared as described by Xiong et al. [93].

Contig sequencing and annotation

Minimum tiling paths of seven and six BACs were identified for contigs 1003 and 431, respectively, by the visual inspection of the FPC assemblies. Adjacent clones were chosen based on their reciprocal position and probability value associated to their overlapping fingerprinted bands as shown by FPC. Sequencing of selected minimum tiling path BACs was done to phase II quality as previously described [73]. Phase II BAC sequences were then assembled into 1003 and 431 contig sequences based on dot plot comparisons and overlap similarity between adjacent clones.

Perl scripts available from the DAWGPAWS package [61,62] were used to convert computational annotation results from multiple sources into a single GFF3 file for combined evidence annotation in Apollo [94] and publication in Gbrowse [95]. Ab initio gene annotation programs used in this process included FGENESH [63] AUGUSTUS [64], SNAP [65], GenElD [66] and GenScan [67]. Because Amborella-specific gene model parameterizations were not available for these programs, multiple plant models were used for each ab initio program. The sequence of the entire contig was BLASTx (e < 1 x 10^-5) searched against gene annotations from Arabidopsis [70], Vitis [45], Z. mays [56], Medicago [71], Oryza [72], and Sorghum [55] as well as tBLASTx (e < 1 x 10^-5) searched against a database of comprehensive Amborella transcript assemblies [25]. In addition, Amborella EST sequences (reads and assemblies; Table 4) were splice-aligned to the contigs using GMAP (Genomic Mapping and Alignment Program) [68] with the PASA (Program to Assemble Spliced Alignments) genome annotation tool [69]. The gene models and BLAST search results were manually combined into gene models using the Apollo genome annotation curation tool [94].

Synteny analysis of sequenced BAC contigs with Vitis and Oryza genomes

Sequenced Amborella BAC contigs 431 (487,318 bp) and 1003 (629,678 bp) were compared to the International Rice Genome Sequencing Project (IRGSP) rice genome assembly (version 5) and the Genoscope 12 × Vitis genome assembly using LASTZ and default parameters. Prior to LASTZ comparisons, all genomic sequences were masked using NCBI’s WindowMasker to remove simple repeats. Significant matches after repeat masking were visualized as dot plots. Gene annotations for the rice and Vitis genomes were obtained from the Rice Annotation Project [96] and Genoscope [97], respectively, and plotted on the vertical axes of the dot plots (Figures 5 and 6). FGENESH [63] annotations for the Amborella contigs were included on the horizontal axes of the dot plots. LASTZ scores were summed for all aligned Amborella-rice or Amborella-Vitis blocks within 100 kb of each other in sequenced genomes. All regions with summed scores >100,000 were considered as syntonic and included in Figures 5 and 6.

Phylogenetic analysis

All alignments were carried out using the program ‘MUSCLE’ [79] run under default settings. Maximum likelihood analyses were run on aligned DNA and amino acid sequences using RAxML [80] and the GTRGAMMA nucleotide substitution model.

Submission of data to GenBank databases

BE5s (HR616970 to HR686434), full-length BAC sequences (AC243594.1 to AC243606.1), Sanger shotgun sequences (HR614237 to HR616931), 454 shotgun sequences (SRP006044), Sanger ESTs (FD425831.1 to FD443502.1) and 454 cDNA sequences (SRX018174, SRX018165, SRX018164, SRX018163, SRX018157, SRX018156) have been deposited in the appropriate NCBI GenBank sequence databases. All sequences are also available at the Ancestral Angiosperm Genome Project website [25].

Additional material

### Additional file 1: Supplemental tables and figures cited with additional details for the physical map and shotgun sequences

### Additional file 2: Synteny analysis of Amborella BAC ends and Vitis genes

Abbreviations

bp: base pair; BAC: bacterial artificial chromosome; BE5: BAC end sequence; EST: expressed sequence tag; FISH: fluorescence in situ hybridization; HICF: high information content fingerprinting; LINE: long interspersed element; LTR: long terminal repeat; MITE: miniature inverted repeat transposable element; SSR: simple sequence repeat; TE: transposable element.

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References
1. Cantino P, Doyle JJ, Graham S, Iudd W, Olmstead R, Soltis D, Soltis P, Donoghue MJ. Towards a phylogenetic nomenclature of Tracheophyta. Taxon 2007, 56:822-846.
2. Leebens-Mack JH, Wall PK, Duarte J, Zheng Z, Oppenheimer D, dePamphilis CW. A genomics approach to the study of floral developmental genetics: strengths and limitations. Adv Bot Res 2006, 42:527-549.
3. Soltis DE, Albert VA, Leebens-Mack J, Palmer JD, Wing RA, dePamphilis CW, Ma H, Carlson JE, Atman N, Kim S, Wall PK, Zuccolo A, Soltis PS. The Amborella genome: an evolutionary reference for plant biology. Genome Biol 2008, 9:402.
4. Mathews S, Donoghue MJ. The root of angiosperm phylogeny inferred from duplicate phytochromosome genes. Science 1999, 286:947-950.
5. Qiu YL, Lee J, Bernasconi-Quadroni F, Soltis PS, Soltis DE, Zanis M, Zimmer EA, Chen Z, Savolainen V, Chase MW. The earliest angiosperms: evidence from mitochondrial, plastid and nuclear genomes. Nature 1999, 402:404-407.
6. Soltis PS, Soltis DE, Chase MW. Angiosperm phylogeny inferred from multiple genes as a tool for comparative biology. Nature 1999, 402:402-404.
7. Jansen RK, Cai Z, Raubeson LA, Danielh H, dePamphilis CW, Leebens-Mack J, Muller KF, Gisinger-Bellain M, Haberer RC, Hansen AK, Chumley TW, Lee SB, Peery R, McNeal JR, Kuehl JV, Boore JL. Analysis of 81 genes from 64 plastid genomes reveals relationships in angiosperms and identifies genome-scale evolutionary patterns. Proc Natl Acad Sci USA 2007, 104:1960-1967.
8. Moore MJ, Bell CD, Soltis PS, Soltis DE. Using plastid genome-scale data to resolve enigmatic relationships among basal angiosperms. Proc Natl Acad Sci USA 2007, 104:1963-1968.
9. Warren WC, Hiller LW, Marshall Graves JA, Birney E, Ponting CP, Grutzner F, Belokov K, Miller W, Clarke L, Chinnwalla AT, Yang SP, Heger A, Locke DP, Mitteki P, Waters PD, Veyrunes F, Fulton L, Fulton B, Graves T, Wallis J, Fuenter XS, Lopez-Otin C, Ordonez GR, Elchier EE, Chen L, Cheng Z, Deakin Je, Alope A, Thomas-Knight K, Kittby, P et al. Genome analysis of the platypus reveals unique signatures of evolution. Nature 2008, 453:175-183.
10. Albert VA, Soltis DE, Carlson JE, Farmerie WG, Wall PK, Ilu DC, Solow TM, Mueller LA, Landherde LL, Hu Y, Buzgo M, Kim S, Yoo MJ, Frohlich MW, Perl-Trevres S, Schlarbaum SE, Bliss BJ, Zhang X, Tankely SD, Oppenheimer DG, Soltis PS, Ma H, dePamphilis CW, Leebens-Mack JH. Floral gene resources from basal angiosperms for comparative genomics research. BMC Plant Biol 2005, 5:5.
11. Kim S, Koh J, Yoo MJ, Kong H, Hu Y, Ma H, Soltis PS, Soltis DE. Expansion of floral MADS-box genes in basal angiosperms: implications for the evolution of floral regulators. Plant J 2005, 43:724-744.
12. Soltis DE, Chanderbali AS, Kim S, Buzgo M, Soltis PS. The ABC model and its applicability to basal angiosperms. Ann Rev Plant 2007, 10:155-163.
13. Vialette-Giraud AC, Adam H, Finet C, Jasinski S, Jauaniss S, Scott CP. Insights from ANA-grade angiosperms into the early evolution of CUP-SHAPED COTYLEDON genes. Ann Bot 2011, 107:1511-1519.
14. Fourquin C, Vinaugier-Douard M, Chambrier P, Berne-Dedieu A, Scott CP. Functional conservation between CRABS CLAW orthologues from widely diverged angiosperms. Ann Bot 2007, 100:651-657.
15. Fourquin C, Vinaugier-Douard M, Fogliani B, Dumas C, Scott CP. Evidence that CRABS CLAW and TOUSLED have conserved their roles in carpel development since the ancestor of the extant angiosperms. Proc Natl Acad Sci USA 2005, 102:4649-4654.
16. Zahn LM, Long H, Leebens-Mack JH, Kim S, Soltis PS, Landherde LL, Soltis DE, dePamphilis CW, Ma H. The evolution of the SEPARATA subfamily of MADS-box genes: a preangiosperm origin with multiple duplications throughout angiosperm history. Genetics 2005, 169:2209-2223.
17. Zahn LM, Leebens-Mack J, dePamphilis CW, Ma H, Theissen G. To B or Not to B a flower: the role of DEFICIENS and GLOBOSA orthologs in the evolution of the angiosperms. J Hered 2003, 95:225-240.
18. Zahn LM, Leebens-Mack JH, Arrington JM, Hu Y, Landherde LL, dePamphilis CW, Becker A, Theissen G, Ma H. Conservation and divergence in the AGAMOUS subfamily of MADS-box genes: evidence of independent sub- and neofunctionalization events. Evol Dev 2006, 8:30-45.
19. Shan H, Zahn L, Guindon S, Wall PK, Kong H, Ma H, dePamphilis CW, Leebens-Mack J. Evolution of plant MADS box transcription factors: evidence for shifts in selection associated with early angiosperm diversification and concerted gene duplications. Mol Biol Evol 2009, 26:2229-2244.
20. Cui L, Wall PK, Leebens-Mack JH, Lindsay BG, Soltis DE, Doyle JJ, Soltis PS, Carlson JE, Arnuganathan K, Barakat A, Albert VA, Ma H, dePamphilis CW. Widespread genome duplications throughout the history of flowering plants. Genome Res 2006, 16:758-769.
21. Van de Peer Y, Fawcett JA, Proost S, Sterck L, Vandepoele K. The flowering world: a tale of duplications. Trends Plant Sci 2009, 14:680-688.
22. Wood TE, Takebayashi N, Barker MS, Mayrose I, Greenspoon PB, Rieseberg LH. The frequency of polyploid speciation in vascular plants. Proc Natl Acad Sci USA 2009, 106:13875-13879.
23. De Bodt S, Maere S, Van de Peer Y. Genome duplication and the origin of angiosperms. Trends Ecol Evol 2005, 20:591-597.
24. Soltis DE, Albert VA, Leebens-Mack J, Bell CD, Paterson AH, Zheng C, Sankoff D, dePamphilis CW, Wall PK, Soltis PS. Polyploidy and angiosperm diversification. Annu Rev Bot 2009, 60:336-348.
25. Ancestral Angiosperm Genome Project. (http://ancaigno.uag.edu/).
26. Lyons E, Pederson BC, Kene C, Alam M, Bing R, Tang H, Wang X, Bowers J, Paterson A, Lisch D, Freeling M. Finding and comparing syntenic regions among Arabidopsis and the outgroups papaya, poplar, and grape: CoGe with rosids. Plant Physiol 2008, 148:1772-1781.
27. Tang H, Bowers JE, Wang X, Ming R, Alam M, Paterson AH. Synteny and collinearity in plant genomes. Science 2008, 320:485-488.
28. Tang H, Bowers JE, Wang X, Paterson AH. Angiosperm genome comparisons reveal early polyploidy in the monocot lineage. Proc Natl Acad Sci USA 2010, 107:472-477.
29. Tang H, Wang X, Bowers JE, Ming R, Alam M, Paterson AH. Unraveling ancient hexaploidy through multiply-aligned angiosperm gene maps. Genome Res 2008, 18:1944-1954.
30. Leitch I, Hanson L: DNA C-values in seven families fill phylogenetic
basal angiosperms. Bot J Linn Soc 2002, 140:175-179.
31. Arizona Genome Institute. (http://www.genome.arizona.edu/orders/direct.
html?library=AT_38a).
32. Luo MC, Thomas C, You FM, Hsiao J, Ouyang S, Buell CR, Malandro M,
McGuire PE, Anderson OD, Dvorak J: High-throughput fingerprinting of
bacterial artificial chromosomes using the snapshot labeling kit and
sizing of restriction fragments by capillary electrophoresis. Genomics 2003,
82:378-389.
33. Nelson WM, Bharti AK, Butler E, Wei F, Fuks G, Kim H, Wing RA, Messing J,
Soderlund C: Whole-genome validation of high-information-content
fingerprinting. Plant Physiol 2005, 139:27-38.
34. Soderlund C, Humphray S, Dunham A, French L: Contigs built with
fingerprinters, markers, and FPC V2.7. Genome Res 2000, 10:1772-1787.
35. Wall PK, Leebens-Mack J, Muller KF, Field D, Altman NS, dePamphilis CW:
PlantTribes: a gene and gene family resource for comparative genomics
in plants. nucAcidS Res 2008, 36:D90-D96.
36. Duarte JM, Wall PK, Edger PP, Landherr LL, Ma H, Pires JC, Leebens-Mack J,
dePamphilis CW: Identification of shared single copy nuclear genes in
Arabidopsis, Populus, Vitis and Oozyo and their phylogenetic utility
across various taxonomic levels. BMC Evol Biol 2010, 10:61.
37. Goremykin VV, Hirsch-Ernst KI, Wolfl S, Hellwig FH: Analysis of the
Amborella trichopoda chloroplast genome sequence suggests that
amborella is not a basal angiosperm. Mol Biol Evol 2003, 20:1499-1505.
38. Jurka J, Kapitonov VV, Pavlicek A, Klonowski P, Kohany O, Walichiewicz J:
Comparative physical mapping links conservation of microsynteny to chromosome
structure and recombination in grasses. Proc Natl Acad Sci USA 2005,
102:13206-13211.
39. Paterson AH, Bowers JE, Bruggmann R, Dubchak I, Grimwood J,
gundlach H, Haberer G, Hellsten U, Mitros T, Poliaikov A, Schmutz J,
Spannagl M, Tang H, Wang X, Wicker T, Bharti AK, Chapman J, Feltus FA,
Glowik G, Grigoriev IV, Lyons E, Maher CA, Marts M, Narechania A, Ottill RP,
Penning BH, Salanov AA, Wang Y, Zhang L, Carpita NC, et al. The
Sorghum bicolor genome and the diversification of grasses. Nature 2009,
457:551-556.
40. Schnable PS, Ware D, Fulton RS, Stein JC, Wei F, Fesstern S, Liang C,
Zhang J, Fulton L, Graves TA, Minx P, Reily AD, Courtem Y, Krakowski SS,
Tomlinson C, Strong C, Delehaunty K, Friesen C, Courtemey B, Rock SM,
Bellet E, Du F, Kim K, Tuch SM, Cotman M, Levy A, Marchetto P, Ochoa K,
Jackson SM, Gillam R, et al. The B73 maize genome: complexity, diversity,
and dynamics. Science 2009, 326:1112-1115.
41. Bell CD, Solits DE, Solits P. The age and diversification of angiosperms re-
visited. Am J Bot 2010, 97:296-1303.
42. Bowers JE, Chapman BA, Rong J, Paterson AH: Unravelling angiosperm
genome evolution by phylogenetic analysis of chromosomal duplication
events. Nature 2003, 422:433-438.
43. Paterson AH, Bowers JE, Chapman BA: Ancient polyploidization pre
divergence of the cereals, and its consequences for comparative
genomics. Proc Natl Acad Sci USA 2004, 101:9903-9908.
44. Tuskan GA, Difazio S, Jansson S, Bohlmann J, Grigoriev I, Hellsten U.,
Putnam N, Ralph S, Rombouts S, Salanov Schein AJ, Stert A, Aerts A,
Blahara RR, Blahara RP, Blahoe D, Boerjan W, Brun A, Brunner A, Busov V,
Campbell M, Carlson J, Chalot M, Chapman J, Chen GL, Cooper D,
Coutinho PM, Couturier J, Covert S, Cunningham R, et al. The
genome of black cottonwood, Populus trichocarpa (Toru. & Gray). Science
2006, 313:1596-1604.
45. Estill JC, Henriksen JL. The DAWGPAWLS pipeline for the annotation of
genes and transposable elements in plant genomes. Plant Methods 2009,
5:8.
46. DAWGPAWLS. (http://dawgpawlsourceforge.net).
47. FGNEISH. (http://softberry.com).
48. Stanke M, Schoffmann O, Morgendorf B, Waack S. Gene prediction in
eukaryotes with a generalized hidden Markov model that uses hints from
external sources. BMC Bioinformatics 2006, 7:62.
49. Korf I: Gene finding in novel genomes. FGENESH... Bioinformatics
2005, 21:403-410.
50. J., Leebens-Mack J, Rabiosis LA, Cu L, Keal J, Fournace MH, Chmurel YH,
Boore JL, Jansen RK, dePamphilis CW: Identifying the basal angiosperm
node in chloroplast genome phylogenies: sampling one’s way out of the Felsenstein zone. Mol Biol Evol 2005, 22:1948-1963.
51. Magallon S. Using fossils to break long branches in molecular dating:
a comparison of relaxed clocks applied to the origin of angiosperms. Syst
Biol 2010, 59:384-399.
52. Smith SA, Beaulieu JM, Donoghue MJ: An uncorrelated relaxed-clock
model of angiosperm evolution. Proc Natl Acad Sci USA 2006, 103:14959-14964.
53. Kurtz S, Narechania A, Stein JC, Ware D: A new method to compute K-
mer frequencies and its application to annotate large repetitive plant
genomes. BMC Genomics 2008, 9:517.
54. KEW C-Value Database. (http://data.kew.org/values/).
55. Bowers JE, Arias NA, Asher R, Avise JA, Ball RT, Brewer GA, Buss RW,
Chen AH, Edwards TM, Estill JC, Exum HE, Goff VH, Herrick KL, Steel CE,
Karanakara S, Lafayette GK, Lemke C, Marler BS, Masters SL, McMallon JM,
Nelson UK, Newcombe GA, Nwakam IA, Oden RN, Philips CA, Raick EA,
Roberts CJ, Ryan SP, Slaughter K, Soderlund C, et al. Comparative
phylogenetic and molecular dating of angiosperms. Proc Natl Acad Sci USA 2005,
102:13206-13211.
56. Paterson AH, Bowers JE, Bruggmann R, Dubchak I, Grimwood J,
gundlach H, Haberer G, Hellsten U, Mitros T, Poliaikov A, Schmutz J,
Spannagl M, Tang H, Wang X, Wicker T, Bharti AK, Chapman J, Feltus FA,
Glowik G, Grigoriev IV, Lyons E, Maher CA, Marts M, Narechania A, Ottill RP,
Penning BH, Salanov AA, Wang Y, Zhang L, Carpita NC, et al. The
Sorghum bicolor genome and the diversification of grasses. Nature 2009,
457:551-556.
57. Schnable PS, Ware D, Fulton RS, Stein JC, Wei F, Fesstern S, Liang C,
Zhang J, Fulton L, Graves TA, Minx P, Reily AD, Courtem Y, Krakowski SS,
Tomlinson C, Strong C, Delehaunty K, Friesen C, Courtemey B, Rock SM,
Bellet E, Du F, Kim K, Tuch SM, Cotman M, Levy A, Marchetto P, Ochoa K,
Jackson SM, Gillam R, et al. The B73 maize genome: complexity, diversity,
and dynamics. Science 2009, 326:1112-1115.
58. Korf I: Gene finding in novel genomes. BMC Bioinformatics 2004, 5:59.
59. Blanco E, Abril JF. Computational gene annotation in new genome
assemblies using GeneID. Methods Mol Biol 2009, 537:243-261.
60. Burke C, Kardon S. Prediction of complete gene structures in human
genomic DNA. J Mol Biol 1997, 268:78-94.
61. Wu TD, Watanabe CM, GMAP: a mapping and alignment program for mRNA
and EST sequences. Bioinformatics 2005, 21:1859-1875.
62. Haas BJ, Delcher AL, Mount SM, Wortman JR, Smith RK JR, Hancock U,
Mati R, Ronning CM, Rusch DB, Town CD, Salzberg SL, White O: Improving
the Arabidopsis genome annotation using maximal transcript alignment
events. nucAcidS Res 2003, 31:3654-3666.
63. Swierdowicz D, Wilks C, Lamesch P, Ferrer E, Putnam N, Ralph S,
GVerson H, Li D, Meyer T, Muller R, Ploetz L, Radenbaugh A, Singh S,
Sitting V, Tissier C, Zhang P, Huala E: The Arabidopsis Information Resource
(TAIR): gene structure and function annotation. nucAcidS Res 2008, 36:
D1009-1014.
64. Cannon SB, Stryker L, Rembautz S, Sato S, Cheung F, Gouay J, Wang X,
Mudge J, Vaidervan J, Schier J, Tschap P, Maganison E, Nicholson C,
Humphray SJ, School H, Mayer KF, Rogers J, Gueter FD, Oldroyd GD,
Debelle F, Cook DR, Retzel EF, Roe BA, Town CD, Tabata S, Van de Peer Y,
Young ND. Legume genome evolution viewed through the Medicago
truncatula and Lotus japonicus genomes. Proc Natl Acad Sci USA 2006,
103:14959-14964.
Zucollo et al. Genome Biology 2011, 12:R48
http://genomebiology.com/2011/12/5/R48
Page 13 of 14
72. Itch T, Tanaka T, Barrero RA, Yamasaki C, Fujiy Y, Hiltun PB, Antonio BA, Aono H, Apweiler R, Bruskiewich R, Burea T, Buer F, Costa de Oliveira A, Fuku G, Habara T, Haberer G, Hain B, Harda E, Hiraki AT, Hirochika H, Hoen D, Hokai H, Hosokawa S, Hsing Yi, Iwasa H, Iwao K, Inamishi T, Ito Y, Jaiswal P, Kanno M, et al: Curated genome annotation of Oryza sativa ssp. japonica and comparative genome analysis with Arabidopsis thaliana. Genome Res 2007, 17:175-183.

73. Project IRGS: The map-based sequence of the rice genome. Nature 2005, 436:793-800.

74. Harris RS: Improved pairwise alignment of genomic DNA. PhD Thesis Pennsylvania State University, Biology Department, 2004.

75. Miller Lab Software. [http://www.bx.psu.edu/miller_lab/].

76. Li L, Stoeckert CJ Jr, Roos DS: OrthoMCL: identification of orthologous groups for eukaryotic genomes. Genome Res 2003, 13:2178-2189.

77. Zuccolo A, Sebastian A, Talag J, Yu Y, Kim H, Collura K, Kudrna D, Wing RA: PlantGDB: a resource for comparative plant genomics. Nucl Acids Res 2008, 36:D959-965.

78. PlantGDB. [http://www.plantgdb.org/].

79. Edgar RC: MUSCLE: a multiple sequence alignment method with reduced time and space complexity. BMC Bioinformatics 2004, 5:113.

80. Stamatakis A: RAxML-VI-HPC: maximum likelihood-based phylogenetic analyses with thousands of taxa and mixed models. Bioinformatics 2006, 22:2688-2690.

81. Edw H, Su A, Muppirala U, Sabhanval M, Willerson MD, Lawrence CJ, Lushbough C, Brendel V: PlantGDB: a resource for comparative plant genomics. Nucl Acids Res 2008, 36:D959-965.

82. Luo M, Wing RA: An improved method for plant BAC library construction. Methods Mol Biol 2003, 236:3-20.

83. Kim H, San Miguel P, Nelson W, Collura K, Wissotzki M, Walling JG, Kim JP, Jackson SA, Soderlund C, Wing RA: Comparative physical mapping between Oryza sativa (AA genome type) and O. punctata (BB genome type). Genetics 2007, 176:379-390.

84. Ammiraju JS, Luo M, Goicoechea JL, Wang W, Kudrna D, Mueller C, Talag J, Kim H, Sinieron NB, Blackmon B, Fang E, Tomkins JB, Bar D, MacKIl D, McCouch , Kurata N, Lambert G, Galbraith DW, Arumuganathan K, K Rao, Walling SJ, Gill N, Yu Y, SanMiguel P, Soderlund C, Jackson S, Wing RA: The Oryza bacterial artificial chromosome library resource: construction and analysis of 12 deep-coverage large-insert BAC libraries that represent the 10 genome types of the genus Oryza. Genome Res 2006, 16:140-147.

85. Ewing B, Hillier L, Wendl MC, Green P: Base-calling of automated sequencer traces using phred. I. Accuracy assessment. Genome Res 1998, 8:175-185.

86. Chou HH, Holmes MH: DNA sequence quality trimming and vector removal. Bioinformatics 2001, 17:1039-1104.

87. Zuccolo A, Sebastian A, Talag J, Yu Y, Kim H, Collura K, Kudrna D, Wing RA: Transposable element distribution, abundance and role in genome size variation in the genus Oryza. BMC Evol Biol 2007, 7:152.

88. SeqClean. [http://sourceforge.net/projects/seqclean/].

89. Chevreux B, Pfisterer T, Drescher B, Dresfel AJ, Muller WE, Wetter T, Suhai S: Using the miraEST assembler for reliable and automated mRNA transcript assembly and SNP detection in sequenced ESTs. Genome Res 2004, 14:1147-1159.

90. Sonnhammer EL, Durbin R: A dot-matrix program with dynamic threshold control suited for genomic DNA and protein sequence analysis. Gene 1995, 167:GC1-10.

91. Sputnik. [http://espressosoftware.com/sputnik/index.html]

92. Morganie M, Hanafey MJ, Powell W: Microsatellites are preferentially associated with nonrepetitive DNA in plant genomes. Nat Genet 2002, 30:194-200.

93. Xiong Z, Kim JS, Pires JC: Integration of genetic, physical, and cytogenetic maps for Brassica rapa chromosome A7. Cytogenet Genome Res 2010, 129:199-198.

94. Lee E, Harris N, Gibson M, Chetty R, Lewis S: Apollo: a community resource for genome annotation editing. Bioinformatics 2009, 25:1836-1837.

95. Stein LD, Mungall C, Shu S, Caudy M, Mangone M, Day A, Nickerson E, Stajich JE, Harris TW, Arva A, Lewis S: The generic genome browser: a building block for a model organism system database. Genome Res 2002, 12:1999-1610.

96. Rice Annotation Project. [http://rapdb.dna.affrc.go.jp/].

97. Grape Genome Browser. [http://www.genoscope.cns.fr/externe/GenomeBrowser/Vitis/].

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