Early Plant Molecular Genetic Research

Molecular Marker (RFLP) Order Conserved in Grasses

- Early genetic research discovered conserved marker order (1993)
  - Rice vs. Corn
  - Corn has a duplication
  - Liguless gene
    - Map to corresponding genetic positions

![Diagram showing conserved linkage between rice chromosome 4 and maize chromosomes 2 and 10.](image)

**Fig. 3.** Conserved linkage between rice chromosome 4 and maize chromosomes 2 and 10. Loci connected by a line are detected by the same clone in both genomes. Maize chromosome 10 is shown in reversed order to clarify the relationship of it with other chromosomes. Approximate positions of centromeres are indicated by solid bars left of chromosomes. Note that the majority of rice chromosome 4 corresponds to a single chromosome arm of both maize chromosomes 2 and 10. Three loci in the middle of rice chromosome 4 (RZ53, RZ467, and RZ86) are not located on either maize chromosome 2 or 10 but, instead, are found on maize chromosomes 4 and 5. The rearrangement(s) leading to this difference between rice and maize likely occurred before polyploidization of maize. See legends of Figs. 1 and 2 for information about locus names and map construction.
Recombination Rates Among Species

- Rates similar between rice and corn (t-test result)
  - Even though genome sizes are very different

Table 1. Comparison of map distance in selected intervals of conserved regions

<table>
<thead>
<tr>
<th>Interval</th>
<th>Map distance, cM</th>
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<tbody>
<tr>
<td></td>
<td>Rice</td>
</tr>
<tr>
<td>CDO455 – CDO920</td>
<td>21.5</td>
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<tr>
<td>CDO718 – RZ166</td>
<td>21.8</td>
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<tr>
<td>CDO395 – CDO400</td>
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<tr>
<td>CDO20 – CDO1081</td>
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<tr>
<td>BCD450 – RZ630</td>
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<td>RZ67 – CDO312</td>
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<td>CDO346 – CDO202</td>
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<td>RZ395 – CDO405</td>
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<tr>
<td>CDO99 – RZ28</td>
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<td>RZ588 – RZ2</td>
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<td>RZ682 – CDO78</td>
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<td>BCD386 – CDO98</td>
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<td>CDO87 – BNL8.29</td>
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<td>RZ569 – BCD135</td>
<td>2.9</td>
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<tr>
<td><strong>Total</strong></td>
<td><strong>137.2</strong></td>
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RZ, rice leaf DNA; CDO, oat leaf cDNA; BCD, barley leaf cDNA; BNL8.29, clone BNL8.29.
Plant Species Loci and QTL are Colinear
Paterson et al. (1995) Science 269:1714
Domestication Loci Common Among Grass Species
Why Should We Study Plant Genome Evolution???

Genomic Synteny

- Large sequence **BLOCKS** Shared in the Same Order between species
  - Same **GENES MAY** control the Same PHENOTYPES
- Warm season legumes have shared gene **MACROSYNTENY**

**Figure 3.** Synteny view between cowpea (**Vu**; *Vigna unguiculata*) and other closely related diploid species. These include: (a) adzuki bean (**Va**; *Vigna angularis*); (b) mung bean (**Vr**; *Vigna radiata*); and (c) common bean (**Pv**; *Phaseolus vulgaris*) using the revised cowpea chromosome numbering system.
Micro-synteny in Grasses

- **Q gene**
  - Major wheat domestication gene
  - Confers free thrashing of naked grain


**Figure 1.** Micro-colinearity between the Q locus of *Triticum monococcum*, *Brachypodium sylvaticum*, and rice. Genes are shown as colored boxes along the physical maps of each species, and transcriptional orientations are indicated by arrows above the boxes. A kilobase (kb) scale is shown above each physical map. The black- and blue-hatched box on the *B. sylvaticum* map indicates a degenerate gene. Orthologous genes are connected by dotted lines. The *T. aestivum* genetic map of the chromosome 5A Q region derived from CS × CS-DIC 5A (Faris et al. 2003) is shown at the top and was used to determine the genetic locations of the EST-based markers XBE406609 and XBG263210, which are orthologous to *BsMIIP/OsMIIP* and *BsPHD/OsPHD*, respectively.
Viridiplantae Phylogeny

Fig. 2: Phylogenetic inferences of major clades. Phylogenetic inferences were based on ASTRAL analysis of 410 single-copy nuclear gene families extracted from genome and transcriptome data from 1,153 species, including 1,090 green plant (Viridiplantae) species (Supplementary Table 1). a, Phylogram showing internal branch lengths proportional to coalescent units ($2N_e$ generations) between branching events, as estimated by ASTRAL-II15 v.5.0.3. b, Relationships among major clades with red box outlining flowering plant clade. Species numbers are shown for each lineage. Most inferred relationships were robust across data types and analyses (Supplementary Figs. 1–3) with some exceptions (Supplementary Fig. 6). Data and analysis scripts are available at https://doi.org/10.5281/zenodo.3255100.
Phylogenetic Relationships and Divergence Times of Land Plants Orders


Angiosperms 238 MYA

Seed Plants 338 MYA

Mesangiosperms 179 MYA

Gymnosperms 446 MYA

Landplants 493 MYA

Tracheophytes 446 MYA

Euphyllophytes

Plant Genome Evolution

How have plants evolved over time to express their extensive biological, cellular, and molecular diversity?

From: Bowles et al (2020) Current Biology 30:530

- Compared gene sets from 208 sequenced genomes across the photosynthetic organisms
  - Viridiplantae = photosynthetic organisms
    - 500,000 species
      - Chlorophyta – Green algae
      - Streptophyta – other algae and land plants
  - Two functions added over time BEFORE the appearance of land plants
    - Multicellularity
    - Terrestrialization
Consistent with other research

- Evolved basic functions of plants included
  - Embryogenesis
  - Plant hormones
  - Symbiotic interactions with:
    - Arbuscular mycorrhizae
    - Rhizobacteria
Data Analysis of Bowles et al. 2020

208 sequenced eukaryotic plant genomes analyzed

- 9 million proteins
  - Clustered into ~650,000 Homology Groups
  - Identified five evolutionarily distinct classifications of HG
    - **Ancestral**
      - present in the Last Common Ancestor (LCA) of a clade **BUT** not necessarily in all members of the clade
    - **Ancestral core**
      - present in **EVERY** representative species within a clade (or absent only in one genome)
    - **Novel**
      - present in the LCA of a clade **AND** absent in all outgroup taxa
    - **Novel core**
      - present in **EVERY** representative species within a clade (or absent only once **AND** absent in all outgroup taxa
    - **Lost**
      - lost in the Last Common Ancestor of a clade

Observations on Core Genes

- Significant core genes added to
  - Streptophyta: n=50
  - Embryophyta: n=103
- Only a few core genes added since these events
  - Only a few core genes added since Tracheophyta and more recent clades
### Number of Homolog Groups Across the Plant Phylogeny

<table>
<thead>
<tr>
<th>Age (MYA)</th>
<th>Group</th>
<th>Homology Group Types</th>
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<tr>
<td></td>
<td>Ancestral</td>
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<tr>
<td>1271</td>
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<td>1184</td>
<td>Viridiplantae</td>
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<td>493</td>
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<td>446</td>
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<td>338</td>
<td>Tracheophyta</td>
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<td>238</td>
<td>Spermatophyta</td>
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<td>Angiosperms</td>
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<td>160</td>
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<td>165</td>
<td>Monocot</td>
<td>9,380</td>
</tr>
<tr>
<td></td>
<td>Eudicot</td>
<td>10,105</td>
</tr>
</tbody>
</table>

**Novel Core Homology Group**

- Represent new functions
  - Each taxonomic level gains new functionality
    - New functionality associated with evolution

**Streptophyta: n = 50 HGs**

- Protein Functions
  - Gene regulation
    - Transcription factors
  - Cell structure, movement, and division
    - Cytoskeletal proteins
- Biological Functions
  - Multicellularity
    - Roots
    - Lateral organ development
Embyrophyta: n = 103 HGs

- **Protein Functions**
  - Protein modification
    - Transferase, oxidoreductase, and ligase
  - Protein transport
    - Transporter proteins and membrane traffic proteins

- **Biological/Molecular Functions**
  - Terrestrialization
    - Lignin biosynthesis
    - UV protection
  - Cell signaling
    - Growth hormone system
  - Transcription factors
    - Beta-helix-loop-helix TFs
Evolution of Plant Genomes

Introduction

Modern plant genomes are quite variable
- ~150 megabase (Mb) Arabidopsis thaliana genome.
- 18,000 Mb hexaploidy wheat genome.

Why understanding the evolutionary history of genomes?
- Applied genetics perspective
- Application of comparative genomics for gene discovery.
  - Arabidopsis terminal flower 1 (tfl1)
    - Encodes a transcription factor
    - It controls indeterminacy/determinacy phenotype
    - Arabidopsis tfl1 as a reference gene
  - Homolog of this gene also controls the phenotype in other
    - Dicot species
      - Snapdragon (Antirrhinum)
      - Pea (Pisum sativum)
    - Monocot
      - Rice (Oryza sativum)
  - Mutations all results in a determinate phenotype
The relevant question
- To what degree are functional genes in one plant species conserved in another species?
  - Important to trace
    - Evolutionary events
    - Related to current organization of plant genomes
Polyploidy and the Construction of Plant Genomes

Whole genome duplication (WGD)
- Common event in the evolution of plant species
  - Entire genome doubles in size
  - Duplicates the same genome
- Two related diploid species merge
  - During mitosis
    - Chromatids migrate to separate daughter cells
  - If they migrate to only one cell
    - The cell will be a tetraploid
- If the 2x duplicate cell is involved in reproduction
  - Resulting gamete
    - 2x the normal number of cells
  - If 2x gamete unites
    - Offspring will be tetraploid

Polyploidy
- An organism that contains extra sets of chromosomes.
  - Tetraploids
    - Cultivated potato
    - Alfalfa
- For a success of any polyploidy
  - It must generate balanced gametes.
    - The same number of chromosomes as other gametes
- Embryos from gametes with the same number of gametes
  - Successfully survive

GENOME DUPLICATIONS
a common EVOLUTIONARY EVENT in all plant species

AUTOPOLYPLOIDS
the species duplicates its OWN genome
Other Polyploids

- Allopolyploids
  - Two species with very similar chromosomal structure and number intermate.
  - After chromosomal doubling, genome will have
    - Number of chromosomes equal to the sum of the number of chromosomes from each of the parent species.

- Examples of allopolyploid species
  - Tetraploid durum wheat (x=14)
  - Hexaploid bread wheat (x=21).

- Durum wheat arose from
  - Union of two diploid species (x=7) species

- Bread wheat arose from
  - Diploid wheat species with the tetraploid wheat species
Constructing the *A. thaliana* genome as a model for eudicot genome evolution

- With the whole genome sequence
  - Study the duplication history of the *A. thaliana* genome.
  - Ancestral duplication signatures could be inferred
    - Blastp analysis
      - Protein vs. protein comparison
      - Identifies gene pairs
        - E-value < -10 used in Fig. 1
      - **Suggests genes are ancestrally related**
    - Duplicates are mapped relative position in the genome
      - Displayed using a dot blot
        - Blocks observed
          - Linear arrayed dots
          - Form a diagonal in the dot blot,
            - Signatures of a duplication event
**Figure 1.** Top right, α duplications. Both x and y axes represent 26,028 genes in their chromosomal order. The best-matching gene pairs are plotted, colour-coded to indicate same (red) or opposite (green) transcriptional orientations. For further analysis, 57 adjacent duplicated regions with opposite orientation and order explicable by localized inversions were combined into 26 ‘large’ duplications (α01–α26) that each included ≥1% (260) of the genes. Eight shorter duplications were pooled (α27). Lower left, β and γ duplications. Both x and y axes represent 21,749 genes, in an inferred ancestral order that accounts for the composition of the 26 large α duplications (at left and bottom). Twenty-nine β or γ duplications (see text) are highlighted. Colours show how the four modern *Arabidopsis* chromosome segments contribute to β or γ duplications, distinguishing contributions to the segments at left and bottom respectively from the: (1) lower-numbered chromosomes (red); (2) higher- and lower-numbered chromosomes (light blue); (3) lower- and higher-numbered chromosomes (dark blue); (4) higher-numbered chromosomes (green). Higher-resolution versions of the figure and lists of gene orders are available (see Supplementary Information).
Figure 1A

- Early comparison of the proteins in the *A. thaliana* genome
  - Red and green diagonals in the upper right panel
    - Block α3
      - Chromosome 1 vs. chromosome 1 block
      - Signature of a duplicated block of genes
      - Genes that have the same conserved order
      - At two ends of the *A. thaliana* chromosome 1
    - Block α5
      - Another pairs of duplicated genes on chromosome 1
    - Block α8
      - Shared block on chromosomes 1 and 3
    - Block, α11
    - Largest block
    - Ends of chromosomes 3 and 2
  - Total
    - 27 major duplicated blocks
      - Strong signals
      - *Signals of a recent duplication*

So how does this relate to the mechanism of genome construction?

- *A. thaliana* underwent a WGD
  - Chromosomes were broken
  - Rearranged into new chromosomes
  - New chromosomes developed
    - *Represent blocks of DNA from the progenitor species*
Figure 2. Comparative physical map of *A. thaliana* and the genetic map of *A. lyrata*. (from: Yogeeswaran et al. Genome Research 15:505)

*A. thaliana* chromosome V built from *A. lyrata* chromosome 8 and *A. lyrata* chromosome 7

Figure 2. Colinearity of A. lyrata linkage map with the *A. thaliana* genome. *A. thaliana* chromosomes (At Chr I – V) are represented as patterned bars (drawn to scale, 1 unit = 1 Mbp; gray rectangles, centromeres; gray circles, heterochromatic knobs). *A. lyrata* linkage groups (Aly LG 1 – 8) are shown in black (drawn to scale, 1 unit = 5 cM). Sixteen collinear blocks are highlighted with the same pattern as the At chromosome to which they correspond. Markers defining the ends of each collinear block are shown on the map in black lettering. Markers mapping with LOD score less than 3.0 are featured in parentheses. Italicized markers map to translocated or nontysntetic regions in *A. lyrata*. Translocations T1 and T2 are highlighted by arrows whose positions correspond to the At chromosome where their collinear region lies. Major inversions I1 and I2 and minor inversion I3 are highlighted in light gray. Three chromosomal fusions are denoted as F1-F3.
Progenitor *Arabidopsis* genome

- How it was modified by the duplication event
- Compare to species that is evolutionary close.
  - *A. lyrata*
    - 8 chromosomes
  - *A. thaliana*
    - 5 chromosomes
- Genetic maps developed using shared loci were

**Fig. 2**

- Five *A. thaliana* chromosomes
  - Constructed from ancestral genome with eight chromosomes
- At Chr I
  - Blocks of AlyLG1 + AlyLG2
- At Chr II
  - Blocks of AlyLG3 + AlyLG4.
- Conclusion
  - *Two species with different chromosome numbers consist of the same chromosomal blocks*
Fig. 1B – Early duplication events

- Shows evidence of more ancient duplications
  - 27 $\alpha$ duplications reoriented
    - Notice block $\alpha$5
    - Two duplicates blocks in the same order
    - Two in an opposite orientation
      - Presumed ancestral order derived from these four blocks
    - Same procedure that uncovered the $\alpha$ blocks.
  - Two types of blocks discovered.
    - 22 $\beta$ blocks
      - Another duplication event in the A. thaliana lineage

The 7 $\gamma$ blocks

- Controversial
  - Hypothesis 1
    - Early duplication in the angiosperm lineage
  - Hypothesis 2
    - Duplication after the split of monocots and dicots
- Grapevine genome sequenced
  - Evidence from the genome appears to have resolved this question
    - Grape
      - Ancestor of the rosids
        - Group of species included A. thaliana.
    - Blast and dot blot analysis of grape genome
Figure 3. Dot blot representation of duplicate regions of the grapevine genome. (from: Jaillon et al. 2007. Nature 449:463)

Figure S5. The grape genome originated from a polyploidy event that joined three ancestral genomes. The nineteen chromosomes of grape are represented on both the x and y axis. Dots represent the positions of paralogous pairs of genes. For clarity, intrachromosomal paralogs are not shown. Clusters of paralogs form a succession of dots, that indicate that the gene order of the ancestral genome was locally maintained. These clusters are painted in seven colours. Each colour marks paralogous blocks, that were colinear in the ancestors of the three constituents of the grape genome. Some regions are not painted in triplicate in this grid, either because a whole region is not visible in synteny with two others in the present-day grape genome (too many rearrangements or gene loss), or because one or two syntenic regions lie in supercontigs which are still not anchored.
Figure 3
- Any genes shared with two other regions of the genome
  - Grape genome has a hexaploid history
- How about other species
  - Signal of hexaploidy is detected
    - Figure 4
      - Grape and poplar genomes were compared
      - Only triplicated regions in grape used
        - Triplicated regions
          - Two copies in poplar
        - *Hexaploid ancestry concept is supported*
        - *Poplar under went an additional WGD after its divergence from the grape lineage*

Shared duplications in dicot and monocot analysed
- Grape and rice orthologs analyzed
  - Hypothesis 1
    - Rice shared the hexaploid ancestry
      - 3-to-3 relationship
        - Not observed
  - Hypothesis
    - Rice does not share the same hexaploid ancestry
      - 3-to-1 relationship observed
  - Conclusion
    - *Monocots and dicots do not share the same hexaploid history.*

(Note: See Tang et al. 2008. Genome Research18:1944 for an alternative perspective.)
**Figure 4.** Comparison of the triplicated blocks and the Poplar genome. (from: Jaillon et al. 2007. Nature 449:463)

**Figure S6.** The distribution of 8,604 orthologous genes between *Vitis vinifera* (x axis) and *Populus trichocarpa* (y axis) chromosomes.
Summary of Eudicot Evolution

- Two diploid mate
  - Tetraploid species developed
- Tetraploid species mated to another diploid
  - Produce the ancestral hexaploid
    - All subsequent eudicots derived from this ancestor
      - Signatures of the same duplications
        - Should be observed in their genome history

Monocot genome evolution.
- Monocots also have a duplication history.
  - Figure 5
    - Compared rice and maize.
      - Maize chromosomes (y-axis) as the reference
        - Most rice genes found in two copies
      - Rice chromosomes (x-axis) as the reference
        - Blocks found three or four times in maize.
    - Conclusion
      - WGD event in the history of monocots
      - An additional duplication occurred in the maize lineage.
Figure 5. A comparison of maize and rice duplication events. (from: Wei et al. (2007) PLoS Genetics 3(7):e123, 1254)

Figure 1. Dotplot Analysis of the Integrated Maize Map against Rice Pseudomolecules

Synteny blocks were detected, and background noise was filtered with SyMAP [37]. The interactive dotplot can be viewed at http://www.agcol.arizona.edu/symap. When clicking the related synteny block, the detailed window with contig number will pop up. The viewer can select the preferred area and double click the selection, and then a graphic alignment is displayed.
doi:10.1371/journal.pgen.0030123.g001
**Figure 6.** A unified model of grass genome evolution. (from: Vogel et al. 2010. Nature 463:763.)

**Supplementary Figure 18. Grass chromosome evolution model.** The monocot chromosomes (r1-r12 for rice, t1-t7 for Triticeae, bd1-bd5 for Brachypodium, s1-s10 for sorghum, and m1-m10 for maize) are represented with a five colour code to illustrate the evolution of segments from a common ancestor with five proto-chromosomes and a n=12 intermediate as described in 62, and are named according to the rice nomenclature. The events that have shaped the structure of the 5 different grass genomes including the 7 Brachypodium chromosome nested insertion events during their evolution from the common ancestor are indicated as whole genome duplication, ancestral chromosome translocations and fusions, and lineage-specific nested chromosome insertions.
Unified model of grass evolution – developing the ancestor
• Based on sequences of genome sequences of
  o Rice
  o Sorghum
  o Brachypodium (a model grass species)
  o Maize
• 56-73 MYA
  o Ancestral grass species containing five chromosomes
    ▪ Duplicated
    ▪ Genome with ten chromosomes appeared
  o Then
    ▪ A4 and A6 fractionated
      • Chromosomes A4, A6, and A2 appear
    ▪ A7 and A10 fractionated
      • Chromosomes A7, A10, and A3 appear
  o Paleopolyploid developed
    ▪ 12 chromosomes
      • Progenitor of all of the modern grasses

Unified model of grass evolution – developing the lineages
• Rice genome structure
  o Represents the ancient paleotetraploid.
    ▪ Basic set of chromosomes
      • Building blocks for other genomes
Figure 6

- Breakage/translocation/fusion events
  - Involve chromosomal fragments from the n=12 ancestor.
    - Developed
      - Brachypodium
      - Poideae (representing the wheat lineage)
      - Panicoideae (representing the maize/sorghum lineage)
      - Panicoideae
        - Simplest history
        - Arose from only four breaks
    - Other lineages
      - More complex patterns of evolution
        - Maize genome
          - Underwent additional duplication
          - Additional breakage/translocation/fusion events
          - *Constructed the modern maize chromosomes*
Summary

- **Plant genomes**
  - A long history of genome duplications
    - Unlike animal and fungal genomes.

- **Figure 7**
  - Illustrates the duplication history
    - (The γ event should be moved to the origin of the eudicot lineage.)
  - Significant role of WGD in development of plant species
    - Many duplications appear 55-70 MYA
      - Transition point
        - Cretaceous and Tertiary periods
          - Mass extinction of species
      - Hypothesis
        - Duplications gave plants the needed gene repertoire
          - *To survive this extinction*
          - *Flourish on earth*

(see Fawcett et al. 2009. PNAS USA 106:5737)

- **Figure 8**
  - Additional species were analyzed
  - Extended the analysis to deeper phylogeny
  - Additional duplication events determined
    - Ancestral seed plants
      - ζ at ~330 MYA
    - Ancestral angiosperms
      - ε at ~220 MYA
Fig. 4: The distribution of inferred ancient WGDs across lineages of green plants. **a**, The locations of estimated WGDs are labelled red in the phylogeny of all 1000 Plants (1KP) samples. **b**, The number of inferred ancient polyploidization events within each lineage is shown in the violin plots. The white dot indicates the median, the thick black bars represent the interquartile range, the thin black lines define the 95% confidence interval and the grey shading represents the density of data points. The sample sizes for each lineage are shown within parentheses along with taxon names on the phylogeny. The phylogenetic placement of inferred WGDs is illustrated in Supplementary Fig. 8 and data supporting each WGD inference are provided in Supplementary Table 2.
Figure 7. History of Plant Genome Duplications at the Cretaceous/Tertiary Border

Journal Article Figure 13. Figure 1. The Distribution of Known Whole-Genome Duplication (WGD) Events within the Plant Kingdom. Most events are shown from Van de Peer et al. [91] but have been updated. The length of each bar along the branch indicates the current estimate for its age. Duplication events of unknown origin are shown in navy blue, triplications in red, known autopolyploidy events in yellow, and allopolyploidy events in green. The white bar associated with Caryophyllales represents 26 independent WGD events, some of which are autopolyploidy and some allopolyploidy. Named duplication events are shown alongside their Greek letter. Abbreviations: Camb., Cambrian; Carb., Carboniferous; Ord., Ordovician; Neo., Neogene; Pal., Paleozoic; Sil., Silurian.
Figure 8: Ancestral polyploidy events in seed plants and angiosperms. [Jiao et al (2011) Nature 473:97]

Original figure legend from manuscript. Two ancestral duplications identified by integration of phylogenomic evidence and molecular time clock for land plant evolution. Ovals indicate the generally accepted genome duplications identified in sequenced genomes (see text). The diamond refers to the triplication event probably shared by all core eudicots. Horizontal bars denote confidence regions for ancestral seed plant WGD and ancestral angiosperm WGD, and are drawn to reflect upper and lower bounds of mean estimates from Fig. 2 (more orthogroups) and Supplementary Fig. 5 (more taxa). The photographs provide examples of the reproductive diversity of eudicots (top row, left to right: Arabidopsis thaliana, Aquilegia chrysanth, Cirsium pumilum, Eschscholzia californica), monocots (second row, left to right: Trillium erectum, Bromus kalmii, Arisaema triphyllum, Cypripedium acaule), basal angiosperms (third row, left to right: Amborella trichopoda, Liriodendron tulipifera, Nuphar advena, Aristolochia fimbriata), gymnosperms (fourth row, first and second from left: Zamia vazquezii, Pseudotsuga menziesii) and the outgroups Selaginella moellendorfii (vegetative; fourth row, third from left) and Physcomitrella patens (fourth row, right). See Supplementary Table 4 for photo credits.
a, Paralogous gene pairs in *Eucalyptus* for the identified palaeohexaploidization (bottom) and palaeotetraploidization (top) events. Each line represents a duplicated gene, and colours reflect origin from the seven ancestral chromosomes (A1, A4, A7, A10, A13, A16, A19). b, Number of synonymous substitutions per synonymous site (\(K_s\)) distributions of *Eucalyptus* paralogues (top) and *Eucalyptus–Vitis* orthologues (bottom). Blue bars (top) indicate \(K_s\) values for 378 gene pairs from the palaeotetraploidization WGD event (red dot), and red bars show \(K_s\) values for 274 gene pairs of the palaeohexaploidization event (red star). c, Evolutionary scenario of genome rearrangements from the Eudicot ancestor to *Eucalyptus* and other sequenced plant genomes; palaeohistory modified from ref. 49.
The Gene-based Evolution of Duplicated Genes

If duplications are a major signature of plant genomes

- Copy number of genes should equal the number of rounds of duplication.

Table 1

- Number of genes found within plant species
  - Complete genome sequence
    - If the hexoploidy concept is true for dicots, and
    - Grape only contains this hexaploid event
      - Estimate
        - Ancestral dicot contains ~10,000 genes
          (=30,000/3$\Omega$).
Table 1. The estimated number of genes in sequenced plant genomes.

<table>
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<tr>
<th>Species</th>
<th>Estimated # of Genes (from <a href="http://www.phytozome.net">www.phytozome.net</a>)</th>
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<td><em>Eudicots</em></td>
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<td>Cucumber</td>
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<tr>
<td>Medicago</td>
<td>50,692</td>
</tr>
<tr>
<td>Soybean</td>
<td>66,153</td>
</tr>
<tr>
<td>Arabidopsis</td>
<td>27,343</td>
</tr>
<tr>
<td>Papaya</td>
<td>27,332</td>
</tr>
<tr>
<td>Grape</td>
<td>30,434</td>
</tr>
<tr>
<td>Mimulus</td>
<td>25,530</td>
</tr>
<tr>
<td><em>Monocots</em></td>
<td></td>
</tr>
<tr>
<td>Sorghum</td>
<td>34,496</td>
</tr>
<tr>
<td>Maize</td>
<td>32,540</td>
</tr>
<tr>
<td>Brachypodium</td>
<td>25,532</td>
</tr>
<tr>
<td>Rice</td>
<td>31,500</td>
</tr>
</tbody>
</table>
Similarly
- Poplar underwent an additional duplication,
  - Theoretically # of genes = 60,000 genes
- *A. thaliana* underwent two duplications
  - Theoretically # of genes = 120,000 genes
- *Not observed*

Monocot calculations
- Rice, Brachypodium, and sorghum only contain a duplication event
  - Number of ancestral monocot genes
    - 15,000 (=30,000/2).
  - Maize
    - Additional duplication event
      - But has undergone a reduction to ~30,000 genes
- Conclusion
  - Necessary to reduce the number of genes to ensure the success of the species.
**Diploidization.**

**The polyploid past history of plants**
- Surprising result for Arabidopsis and rice genomes
  - Why??
    - Selected for sequencing because of their small genome sizes

**Consequences of polyploidy?**
- Doubling or tripling of the number of chromosomes
  - Evident for monocots.

**Fate of the additional gene set from the WGD**
- Concept
  - Species cannot maintain the entire set of duplicate chromosomes
  - New genes a problem
    - Generate deleterious mutations
    - Compromises the fitness of a genome
  - Genome must transition back to its original state.
    - Process is called
  - *Diploidization.*
To revert back to the diploid state
  • Many duplicate genes must be eliminated from the gene set
    o But a recently duplicated genome
      ▪ Soybean
        • Withstands the extra copies
        • Genome about 2X the basic set of 30,000 genes of hexoploid ancestral eudicot

Events associated with diploidization
  • Duplicate genome must change its chromosome pairing pattern
    o After the duplications,
      ▪ Four chromosomes pair
      ▪ Form quadravalents
    o Chromosomal structure must be changed so
      ▪ Bivalents must be formed
  • Result
    o Doubling of the chromosome number
      ▪ Seen for the monocot lineage
    • Once bivalents are formed
      o Gene sets can evolve
        ▪ Processes
          • Deletions and chromosomal rearrangements
Duplicate genes can undergo specific changes
- Common fate
  - Gene death of new copies
    - Loses associated with
      - Chromosomal breakage
      - Rearrangements.
  - Result
    - New basic set of chromosomes and genes will have appeared

Duplicate genes fate differs
- Some are retained as multicopy
  - Up to the ploidy level for that species
- Other reduced to only a single copy

“Deletion resistant” genes
- Not reduced to single copy
  - Dosage dependent
    - Mainly encode
      - Transcription factors
    - May lead to
      - Complex morphologies

“Duplication resistant” genes
- Must be maintained as single copy
  - Mainly encode
    - Enzymes or genes of unknown function
Developing new functions

Duplicate set of genes cannot be maintained
- Deleterious mutations can arise
- Duplicate genes are modified
  - Changes will provide
    - New functions
    - Altered altered functions
  - New functions may lead to the evolution of the species
    - Higher level of fitness
    - Evolutionary modifications of duplicate genes

**Neofunctionalization.**
- One duplicate gene maintains its original function
- Second gene evolves a function
  - May increase the adaptability of an individual

**Subfunctionalization**
- Modifies the duplicates
- Basic structure of both copies altered
  - Expression pattern of the gene changes
    - Results in a higher level of the protein production
- Alternately, the function of the original gene is maintained
  - Structure of both copies is significantly changed.
    - New copies retains
      - Part of the original function
    - Two genes work together
      - Function of the original gene maintained
Synteny: The Result of WGD and Reconstructing Plant Genomes

Synteny among plant species.

- Major result of the duplication history
  - Synteny
    - Maintenance of gene order between two species
  - Classic approach to synteny
    - Based on shared markers mapped onto two different species.

- Macrosynteny is detected by
  - Large scale chromosomal blocks shared by two species.

Fig. 9

- Example of macrosynteny
  - Tomato and eggplant
    - Eggplant linkage group 4
      - Evolutionarily related to tomato
    - Linkage groups 10S and 4L.
      - Highly conserved marker order over many centimorgans of the two genomes
Figure 9. Macrosynteny between tomato and eggplant, including a QTL for a shared domestication trait. (from: Doganlar et al. 2002. Genetics 161:1713.)
**Genetic mapping of shared genes**
- First method of comparing species
- Only way to compare species that have not been sequenced
- Many examples of synteny mapping in plants.
- The power of synteny mapping
  - Discovery of shared loci from two species
    - Control the same phenotype
    - Map to the same genetic location.

**Fig. 9 again**
- Major QTL for fruit striping
  - Eggplant linkage 4.
  - Previous work with tomato
    - Major QTL
      - Linkage group 10 of tomato
  - Syntenic marker and QTL observed here
- Hypothesis
  - Multiple loci are shared in the same macrosyntenic order
    - Same ancestral gene is controlling this trait in these two species.
Leveraging knowledge in one species for gene discovery in a second species

- Phenotypic traits mapped extensively in one species
  - Points a researcher working on a second species
  - Likely location of a similar gene in second species.
  - Leverage is
    - Great aid for genetic discovery
    - For species in where the discovery of important genetic factors are limited by a lack of funding