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Phenotypic Variation in Plant Polyploids

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Abstract

Polyploidy, or whole-genome duplication (WGD), is an important genomic feature for all eukaryotes, especially many plants and some animals. The common occurrence of polyploidy suggests an evolutionary advantage of having multiple sets of genetic material for adaptive evolution. However, increased gene and genome dosages in autopolyploids (duplications of a single genome) and allopolyploids (combinations of two or more divergent genomes) often cause genome instabilities, chromosome imbalances, regulatory incompatibilities, and reproductive failures. Therefore, new allopolyploids must establish a compatible relationship between alien cytoplasm and nuclei and between two divergent genomes, leading to rapid changes in genome structure, gene expression, and developmental traits such as fertility, inbreeding, apomixis, flowering time, and hybrid vigor. Although the underlying mechanisms for these changes are poorly understood, some themes are emerging. There is compelling evidence that changes in DNA sequence, *cis*- and *trans*-acting effects, chromatin modifications, RNA-mediated pathways, and regulatory networks modulate differential expression of homoeologous genes and phenotypic variation that may facilitate adaptive evolution in polyploid plants and domestication in crops.

Keywords

polyploidy; nonadditive gene expression; epigenetic regulation; RNA interference; evolution

INTRODUCTION

Polyploids can be classified into allopolyploids and autopolyploids based on the origins and levels of ploidy (25,49,135) (Figure 1). An autopolyploid results from doubling a diploid genome (Figure 1*a*). An allopolyploid is formed by the combination of two or more sets of distinct genomes). Mechanisms include interspecific hybridization followed by chromosome doubling (Figure 1*b*), fertilization of unreduced gametes between two diploid species (Figure 1*c*), or interspecific hybridization between two autotetraploids (Figure 1*d*). The two identical chromosomes (red or blue) within a species are homologous, while the chromosomes derived from different species (red and blue) are orthologous but become homoeologous within an allotetraploid. In an allopolyploid, only bivalents are formed because meiotic pairing occurs between homologous chromosomes (Figure 1*b*,*c*,*d*). If the homoeologous chromosomes have some segments that are homologous (Figure 1*e*), pairing may occur between the homoeologous chromosomes, resulting in the formation of multivalents and segmental allotetraploids (135). Here we consider allopolyploids and amphidiploids or disomic allopolyploids to be synonyms. Strictly speaking, only bivalents are formed in the amphidiploids and disomic allopolyploids.

In addition to polyploidy, some plant and animal species exist as intraspecific and inter-specific hybrids (96,124). Many plants that transmit as diploids are actually paleopolyploids (ancient polyploids), which are derived from at least one event of whole-genome duplication (WGD) followed by massive gene loss and genomic reorganization through a process known as

diploidization (152). *Arabidopsis* (14,17,144), rice (154), and maize (45) are good examples of diploidized paleopolyploids. An estimated 30–70% of plant species are of polyploid origin (93,152). That estimate is as high as 100% if paleopolyploids are included (152).

Polyploidy is a fundamental but relatively underexplored biological process. It is widespread but little is known about how duplicate genes and genomes function in the early stages of hybridization, and how the duplicate genes maintain and diverge functions during plant evolution and crop domestication. Many polyploids are ancient, and their exact progenitors are often unknown. Resynthesized polyploids with known progenitors are excellent materials for dissecting gene expression and genomic changes in early stages and comparisons with older polyploids (28,79,132,149). In addition to the parental phenotypes, polyploids give rise to phenotypes that are intermediate between the two parents and to novel phenotypes that are absent in or exceed features of the contributing parents (77,82,119), suggesting nonadditive gene expression. Some traits, such as increasing levels of drought tolerance, apomixis, pest resistance, flowering-time variation, and organ size, may allow polyploids to enter new niches or improve their fitness. Indeed, polyploids may survive better than their diploid progenitors in harsh environments, such as high altitudes and latitudes and cold climates, whereas both diploids and polyploids often thrive and cohabit in mild conditions (49,135). Moreover, polyploidy is a means of permanent fixation of hybrid vigor and dosage regulation, which may be why many crops (e.g., wheat, cotton, oats, canola, potato, peanuts, sugarcane, coffee, and strawberry) are of polyploid origin (63,93). Thus, polyploidy has been studied in the context of evolution, genetics, breeding, and molecular biology (25,29,50,76,79,86,95,111,132,149).

Interspecific hybridization and allopolyploidization occur frequently in plant taxa including Brassica (133), Gossypium (150), Senecio (1), Spartina (10), Tragopogon (141), and Triticum (40,130). Furthermore, hybrids can be formed between different genera including Triticum (wheat)-Secale (rye) (62), Triticum (wheat)-Hordeum (barley) (106), Zea (maize)-Avena (oat) (123), and Zea (maize)-Tripsacum (gamma grass) (58). Some allopolyploids (e.g., Tragopogon miscellus and T. mirus) were produced in natural conditions as recently as ~80 years ago, and new Tragopogon allotetraploids appear to form every year (141). In contrast, polyploids are rarer in animals than in plants (90,107). Interspecific hybrids occur in vertebrates and mammals (e.g., a mule is a hybrid between a horse and a donkey), but they cannot produce offspring probably because of genomic incompatibility and/or imbalance in imprinting and sex chromosome dosage (107,112). Polyspermy (fertilization of more than one sperm into one ovum) causes human triploids in 1-3% of conceptions, and the triploid fetuses are aborted (100). An isolated case of a tetraploid South American rodent (Tympanoctomy barrerae) is still debatable (44,139). Except for endopolyploidy (a diploid individual with cells containing more than 2 C amount of DNA in their nuclei) in some cell types (37), aneuploid and polyploid cells in animals and humans are often associated with malignant cell proliferation or carcinogenesis (136).

MECHANISMS FOR GENE EXPRESSION DIVERGENCE BETWEEN ALLOPOLYPLOIDS AND THEIR PARENTS

Several mechanisms may affect the fate of orthologous and homoeologous genes in polyploids (25,29,76,79,86,95,111,132,149) (Figure 2). First, the majority of homoeologous genes are coexpressed. Second, some duplicate genes are lost, mutate, or diverge (due to genetic changes). The half-life of an active paralogous gene that becomes mutated or lost is estimated to be 2–7 million years (87). Third, epigenetic changes may reprogram gene expression and developmental patterns of new allopolyploids. The impact of these mechanisms on various polyploids can be very different. For example, sequence elimination is predominately observed in wheat and *Tragopogon* allopolyploids (40,130,141); chromosomal translocations and transposition (insertion of a DNA fragment into homoeologous chromosomes) are common in

Brassica allopolyploids (133); and changes in gene expression appear to be a major consequence in *Arabidopsis* and cotton allopolyploids (2,73,147,148). Moreover, genetic and epigenetic changes may be interrelated (25); reactivating transposons by chromatin modifications or RNA-mediated pathways may lead to chromosomal breakages and rearrangements. Eliminating DNA sequences (regulatory and/or coding regions) may alter dosage-dependent gene regulation and chromatin structure.

Mutations, Sequence Eliminations, and Chromosomal Rearrangements (Genetic Changes)

Elimination of chromosome- or genome-specific sequences may occur during polyploid formation (Figure 2*a*). The stochastic changes of duplicate genes may promote polyploid speciation, which is supported by studies in *Brassica* (133), wheat (40,130), and *Tragopogon* (141). In the resynthesized allopolyploids, loss of parental fragments and/or the appearance of novel fragments are commonly observed. Rapid sequence elimination in the resynthesized allopolyploid wheat may account for a relatively high amount (~14%) of genome- or chromosomal-specific DNA sequences (40,130), suggesting that differential elimination of genome-specific sequences facilitates pairing between homologous chromosomes but not homoeologous chromosomes.

Changes in DNA sequence may contribute to the loss of duplicate gene expression and function. Indeed, many isozyme loci are lost during polyploidization, such as chlorophyll a/b binding protein genes in *Polystichum munitum*, leucine aminopeptidase loci in tetraploid *Chenopodium*, and phosphoglucose isomerase loci in homosporous fern and *Clarkia* (48, 116). Estimates indicate that in the salmonid and cyprinid fish, the loss of duplicate isozyme loci can be as high as 35–65%, suggesting that loss of duplicate gene function is common after polyploidization, which occurred 50 million years ago (Mya) in this lineage (41). In *Tragopogon*, 9 of 10 genes that display expression differences are associated with changes in allelic DNA sequence (141). However, loss of gene function may also suggest an epigenetic cause (see below).

Epigenetic Regulation of Orthologous Loci

Genetic mutations can explain the cause of gene loss over evolutionary time, but many silencing phenomena may be epigenetically controlled, especially in the early stages of polyploid formation. When two different genomes are combined into a single cell, they must respond to the consequences of genome duplication, especially duplicate copies of genes with similar or redundant functions. Increased gene or genome dosage may induce disease syndromes and abnormal development (7,38). Thus, the expression of orthologous genes must be reprogrammed through epigenetic mechanisms (Figure 2a) in the early process of polyploidization. This resembles the "genomic shock" phenomenon proposed by McClintock (98). The genomic shock occurs rapidly in interspecific hybrids and allopolyploids, resulting in demethylation of retroelements (92), relaxation of imprinting genes (20,68,145), and silencing and activation of homoeologous genes (2,26,69,73,91,146-148), including rRNA genes subjected to nucleolar dominance (expression of rRNA genes from only one progenitor in an interspecific hybrid or allopolyploid) (117,122). Epigenetic changes, which are potentially reversible, provide an effective and flexible means for a polyploid cell to respond to polyploidy or genomic shock. Moreover, gene silencing or activation that is initially epigenetic and reversible could be one step toward a genetically fixed and irreversible state. Epigenetic silencing may also accelerate sequence mutation rates of the affected genes, as observed in repeat-induced point mutations in duplicated genes of *Neurospora crassa* (128).

Mechanisms for epigenetic regulation of homoeologous genes in the allopolyploids are reminiscent of those for X-chromosome inactivation (75), gametic imprinting (143), paramutation (21,121,134), and homology-dependent gene silencing (11,67,94). However,

ploidy-dependent gene regulation has some unique features. First, epigenetic interactions are established among four alleles of two homoeologous loci in allotetraploids compared with two alleles of one locus in a diploid. Second, homoeologous genes from different parental origins may be up- or downregulated in a chromosomal domain (73,147), which is different from dosage compensation that often refers to concerted or unidirectional changes in gene expression. Third, at least some epigenetic silencing phenomena in allopolyploids are stochastically established and require multiple generations (24,104,148), probably because of the complex process of sorting out chromosome pairing in the allopolyploids. Fourth, pairing occurs mainly between homologous chromosomes, but occasionally between homoeologous chromosomes in allopolyploids (Figure 1*e*), which may affect gene expression. Finally, because of divergence of regulatory sequences between the progenitors and of heterologous proteins produced in the allopolyploids, *cis-* and *trans-* acting effects on homoeologous genes (146,151) in various biological pathways constitute a major mode of gene regulation in the allopolyploids.

Additive and Nonadditive Expression of Orthologous Loci

Many genes display dosage dependency and are expressed additively in aneuploids and polyploids (53). If the levels of gene expression and phenotypic variation in the progenitors are additive, they would have the midparent values (MPVs) in the polyploids; that is, one plus one is equal to 2 (Figure 2*d*). If the gene expression is nonadditive (different from the MPV), the values would be larger than 2 or smaller than 2. The former would suggest gene activation including overdominance, whereas the latter would suggest repression and/or silencing. One model to explain additive gene regulation is that there are extra control settings or rheostat potentials (four levels) in a tetraploid compared with two levels in a diploid (Figure 2*b*). In *Arabidopsis* and *Brassica*, the alleles in the *FLOWERING LOCUS C (FLC)* loci display additive effects on flowering time (102). In *Arabidopsis* allotetraploids, the expression of *A*. *thaliana* and *A*. *arenosa FLC* loci is additive, giving rise to a late flowering phenotype (146). Furthermore, up to ~90% of the transcriptome is expressed additively in resynthesized *Arabidopsis* allotetraploids (147). Although odd and even dosage effects may vary in a ploidy series (see below), coexpression and coevolution of orthologs and paralogs suggest that a selective advantage is obtained from dosage dependency (15,47).

Similar numbers of genes (5-11%) are differentially expressed between the parents and resynthesized allotetraploids in stable allotetraploids and five selfing generations (148), and a slightly low number of genes (~2.5%) are differentially expressed between a natural allotetraploid, *A. suecica*, and its assumed progenitors (73). Most gene expression changes observed in early generations are maintained in the late generations and natural allotetraploids, suggesting that rapid and stochastic changes in the resynthesized allotetraploids are responsible for adaptive evolution (148).

Furthermore, allopolyploidy may induce regulatory incompatibilities as well as selective advantage by combining heterologous protein products (Figure 2*c*). For instance, there is evidence that some protein heterodimers may not function as well as homodimers or vice versa (115,118). Thus, a silencing strategy could balance regulatory incompatibility and the advantages of having multiple copies of orthologous genes or gene products (e.g., transcriptional factors) spontaneously produced in an allopolyploid cell. Alternatively, novel interactions between heterologous protein products may provide a molecular basis for hybrid vigor and novel adaptation.

INSTANTANEOUS SPECIATION AND A MODEL FOR POLYPLOIDY STUDIES

New species are often gradually formed because of geographical and ecological separations from an ancestral species (49). However, new species are believed to have arisen suddenly via

polyploidization in plants and some animals, including vertebrates such as amphibians and lizards (18,49,93,112). For example, *Arabidopsis suecica* (2n = 4x = 26) is a natural allotetraploid formed 12,000 to 1.5 Mya (64,71,125). The two progenitor species, *A. thaliana* and *Arabidopsis arenosa* (108,123), split ~6 Mya (71), similar to the distance between humans and chimpanzees (~6.3 million years) (114). Despite this distance, *A. thaliana* autotetraploid (2n = 4x = 20) and *A. arenosa* tetraploid (2n = 4x = 32) can hybridize to produce *A. suecica*-like plants (2n = 4x = 26) (Figure 3*a*,*b*,*c*). *A. arenosa* is thought to be an autotetraploid (31), but sequencing analysis suggests that it is not a pure autotetraploid (146) (L. Tian, J. Wang & Z.J. Chen, unpublished). The resynthesized allotetraploids are meiotically stable (30,147) and contain five pairs of *A. thaliana* chromosomes and eight pairs of *A. arenosa* chromosomes (28,30,147) (Figure 3*c*). Compared with resynthesized *Brassica* and wheat allopolyploids that undergo rapid changes in chromosomal structure and DNA sequences (40,133), the frequency of aneuploids and chromosome abnormalities in *Arabidopsis* resynthesized allotetraploids is relatively low (30).

The nascent allotetraploids (F_1 individuals) are genetically identical (Figure 3*a*) and showed subtle phenotypic variation. Some variation among F_1 individuls may derive from heterozygosity of the outcrossing tetraploid *A. arenosa* parent, whereas other variation may result from interactions between *A. arenosa* and the different genotypes of *A. thaliana* used in interspecific hybridizations. The degree of variation depends on parental genotypes used in the interspecific hybridization. For example, the seed set is higher in the nascent allotetraploids (F_1) between *A. arenosa* and *A. thaliana* C24 or Ler ecotype than those between *A. arenosa* and *A. thaliana* C01 provide (20,31), indicating genotypic effects on interspecific hybridization. Hybridization was successful only in the crosses using *A. thaliana* as a maternal parent and *A. arenosa* as a pollen donor (31) (J. Wang, L. Tian & Z.J. Chen, unpublished), probably because *A. arenosa* is outcrossing and self-incompatible. Most F_1 individuals and selfing progeny in late generations resemble the *A. arenosa* parent and *A. suecica* (31,91,147) (Figure 3*c*), although diverse phenotypes are observed in segregating populations (F_2 - F_3) (Figure 3*b*). Therefore, *A. arenosa* appears to be phenotypically dominant over *A. thaliana* in the allotetraploids (28,147).

The allotetraploids obtained from selfing the $F_{1}s$ show stable karyoptes in the fifth generation (Figure 3*c*), but exhibit a wide range of variants, some of which are absent in either parent (transgression) (Figure 4*a*). Moreover, the allotetraploids display hybrid vigor: larger rosettes, more leaves, longer and wider leaves, and taller plants than the parents. The fertility rate of the plants in the selfing progeny varies from one lineage to another (30,31). The overall level of fertility improves after each generation of selfing, suggesting that genome incompatibility and gene expression divergence between the progenitors are gradually overcome (148).

The flower colors varied from pink (like *A. arenosa*) in the early generation (F_1) to a mixture of pink and white flowers in the intermediate generations (S2–4) and white in the late generation (S5). During selfing (S3), there is a low frequency of mixed white and pink flowers in the same flower branch (Figure 4b, bottom), which is transient and mosaic (derived from the same zygote). The appearance of varigation within the same flower branch suggests rapid changes in the expression of genes involved in anthocyanin synthesis pathways probably via epigenetic regulation.

NONADDITIVE GENE REGULATION AND TRANSCRIPTOME DOMINANCE IN ALLOPOLYPLOIDS

In 1928, Navashin coined the term "amphiplasty" to describe chromosomal changes in interspecific hybrids of *Crepis* (110). He defined "differential amphiplasty" as specific changes in a few chromosomes (disappearance of satellites or secondary constrictions) and "general

amphiplasty" as the overall changes in chromosomal morphology (shortening, thickening, or lengthening of chromosomes) from one species in the interspecific hybrids or amphidiploids. Changes in chromosomal morphology might also affect gene expression. Indeed, following the pioneering work of Navishin & McClintock (97,110), several contemporary researchers demonstrated that differential amphiplasty is synonymous to nucleolar dominance (117,122). The disappearance of the secondary constrictions is caused by silencing of rDNA loci in those chromosomes (117). Nucleolar dominance is observed in *Drosophila* interspecific hybrids and *Xenopus*, *Arabidopsis*, *Brassica*, and wheat allopolyploids (117,122). The dominance is reversible and developmentally regulated and is controlled by chromatin modifications involving DNA methylation and histone acetylation (26,27). Blocking histone acetylation or DNA methylation derepresses the silenced rRNA genes subjected to nucleolar dominance. Both DNA methylation and histone hypoacetylation reinforce the formation of the "inactive" chromatin state, resulting in gene silencing (72). The silencing of rDNA chromatin requires at least one histone deacetylase (AtHDA6) that is localized in nucleoli (35).

General amphiplasty may be similar to the effects of genomic shock (98). Combining two genomes in a "new" polyploid cell may generate the genomic shock and release some constraints imposed on unstable elements locked in a junk yard (e.g., transposable elements in heterochromatin). Little is known about the consequences of general amphiplasty or genomic shock on interspecific hybrids or allopolyploids that have balanced pairs of chromosomes.

To begin to test this, Wang et al. (144) studied transcriptome divergence in Arabidopsis allotetraploids and their progenitors. First, they compared gene expression differences between the two progenitors using the spotted oligo-gene microarrays designed from ~26,000 annotated genes that share a high percentage of sequence identities between A. thaliana and A. arenosa. Most of the oligos can cross-hybridize with both A. thaliana and A. arenosa genes (74). More than 15% of the transcriptome is differentially expressed between A. thaliana and A. arenosa that diverged ~6 Mya. Approximately 2,100 genes (8%) are more abundantly expressed in A. thaliana than in A. arenosa, whereas 1,818 genes (7%) are expressed at higher levels in A. arenosa than in A. thaliana. Second, Wang et al. (144) compared mRNA abundance in an allotetraploid with the mid parental value (MPV: an equal mixture of RNAs from two parents). If the genes from two progenitors are additively expressed (Figure 2d), their cumulative expression levels in the allotetraploid are equal to MPV. Nonadditive expression suggests that at least one of the homoeologous genes is up- or downregulated. There may also be instances in which silencing of a locus is compensated by increased expression of its homoeologous locus, which cannot be detected in this comparison. Wang et al. (146) found that 2,011 genes (~8%) are nonadditively expressed in two independently derived allotetraploids using a common variance analysis, and up to ~38% genes using a pergene variance analysis. Interestingly, ~68% of the genes that are nonadditively expressed in the allotetraploids are differentially expressed between the two parents, suggesting that the genes with species-specific expression patrterns are subjected to expression changes in the allopolyploids. Remarkably, among the nonadditively expressed genes, more than 65% of the genes are downregulated in the allotetraploids. Among them, >94% of the genes that are expressed at higher levels in A. thaliana than in A. arenosa are downregulated in the allotetraploids. These data indicate that the genes with A. thaliana expression patterns tend to be repressed, whereas the genes with A. arenosa expression patterns are transcriptionally dominant in the allotetraploids, coincident with the phenotypic and nucleolar dominance of A. arenosa in the allotetraploids (117,147) (Figures 3 and 4).

Interestingly, similar levels of transcriptional changes were observed in maize diploid hybrids (9.8%) (140) and polyploid taxa of *Senecio* (5%) (60), wheat (7.7%) (59), and cotton (5%) (2). The high percentage of gene expression changes in the *Tragopogon* allopolyploids (~17.5%) (141) is partly associated with a high level of polymorphism (~11%) within

populations between the two parents and a moderate amount of variation (>2.5%) among allopolyploid populations. These numbers are also similar to those observed in interspecific hybrids of *Drosophila* (103,151), suggesting that the levels of transcriptional changes induced by hybridization may be fairly consistent even across plant and animal kingdoms. Transcriptome dominance is also observed in an analysis of ~210,000 expressed sequence tags (ESTs) derived from an ovular cDNA library of tetraploid cotton (*Gossypium hirsutum* L.) (153). The upland cotton was formed by ancient interspecific hybridization between AA and DD genome species (150). AA subgenome ESTs of all functional classifications including cell cycle control and transcription factor activity were selectively enriched in *G. hirsutum* L., a result consistent with the production of long lint fibers in AA genome species. Therefore, transcriptome dominance is likely a general consequence of hybridization effects on gene expression in interspecific hybrids and allopolyploids.

The number of genes displaying expression changes in *A. thaliana* autopolyploids is much smaller than that in the allotetraploids (147). In yeast, Galitski et al. (43) found that 10 genes are induced and seven genes are reduced in response to an increase in ploidy levels (haploid, diploid, triploid, and tetraploid). The cell size increases with increasing ploidy levels, which is correlated with repression of G1 cyclins (*Cln1* and *Pcl1*). *FLO11*, a gene important to the invasiveness of the yeast cells, is repressed with increasing ploidy levels. The reduction of *FLO11* expression in cells of higher ploidy is correlated with diminished invasion, suggesting a role of ploidy-dependent gene regulation in adaptive evolution. Collectively, the data indicate that genome doubling has smaller effects on gene expression changes than intergenomic hybridization.

What factors affect transcriptome dominance in the allopolyploids? Is the gene repression controlled by widespread chromatin modifications or a few "key" regulatory genes? Over time, the progenitor species may have evolved to possess species-specific gene expression patterns. Modulation of the species-specific expression of these genes may determine the outcome of transcriptional and posttranscriptional competition between the two parental genomes in their offspring. Changes in chromatin landscape on repressed genes may result from concerted modifications of many genes in one species, perhaps by a mechanism similar to that for nucleolar dominance (117). Alternatively, expression changes in a few regulatory genes such as transcription factors and microRNAs may induce *trans*-acting effects on many downstream pathways (25). For example, a Myb transcription factor gene is responsible for hybrid-induced incompatibilities in *Drosophila* interspecific hybrids (8). Also, a single miRNA can regulate hundreds of genes involved in the transition from one developmental stage to another (39).

The role of chromatin modifications in silencing or activating protein-coding genes in allopolyploids has been demonstrated in several recent studies (70,73,91,104,146,148). Silenced genes can be reactivated by aza-dC (73), a chemical inhibitor of DNA methylation, or by downregulation of the genes encoding DNA methyltransferases using RNA interference (RNAi) (148). Treating allotetraploids with aza-dC generates pleiotropic effects on natural and synthetic allotetraploids including reactivation of mobile elements (91). Reactivation of transposons is also observed in the synthetic allotetraploids (92). The above data suggest that two species may have possessed different levels of chromatin modifications for many genes that display species-specific expression patterns. Perturbation of chromatin structure may have occurred during the formation of interspecific hybrids or allopolyploids, leading to the changes in gene expression.

Factors other than chromatin modifications may also be responsible for genomewide nonadditive gene regulation. Non-additively expressed genes are randomly distributed along the chromosomes (147). Within a small chromosomal region in which *TCP3* and *RFP* genes are located, *A. thaliana TCP3* is expressed, whereas *A. arenosa TCP3* is silenced (73). For

RFP, *A. thaliana RFP* is repressed, whereas *A. arenosa RFP* is expressed. Interestingly, the neighboring genes located between *TCP3* and *RFP* loci are coexpressed. The above data are reminiscent of the silenced rRNA genes that are restricted in the rDNA loci (81). Furthermore, in the *met1*-RNAi *A. suecica* lines, several silenced genes are not reactivated (148). These data argue that widespread chromatin remodeling does not explain nonadditive regulation for all genes, but support the notion that each gene is regulated through interactions among homoeologous loci such as paramutation-like phenomena observed in *A. thaliana* tetraploids (104).

ALLOPOLYPLOIDY AND HETEROSIS

Genome-wide nonadditive gene regulation observed in the allotetraploids correlates with expression divergence between the parents. Thus, hybrids derived from distantly related species may induce a high level of gene expression changes in a nonadditive fashion, providing molecular bases of hybrid vigor (13) and phenotypic variation in the allotetraploid progeny (31). Hybrid vigor refers to the performance of an F₁ hybrid higher than MPV or the best parent. The genetic basis for heterosis is predicted to be associated with dominant complementation of slightly deleterious recessives (dominance model) (19,66) or overdominant gene action in which genes have greater expression in heterozygous conditions (overdominance model) (32,36). According to the dominance model, highest performance should be observed when all dominant favorable genes from both parents are in homozygous conditions. The overdominance model suggests that heterosis should reach its peak at the maximum levels of heterozygosity and dissipate when approaching homozygosity. Moreover, overdominance is accompanied by nonallelic or epistatic interactions, and epistasis is involved in most QTLs associated with inbreeding depression and heterosis in corn (137) and rice (84). Comparing genome-wide gene expression data with phenotypic traits (OTLs) may provide new insights into the role of gene expression changes in various biological pathways that give rise to hybrid vigor.

The gene expression changes observed in maize diploid hybrids (6,140) and genomewide transcriptome dominance in *Arabidopsis* allotetraploids (147,148) support both dominance and overdominance models. Many genes in energy, metabolism, cellular biogenesis, and plant hormonal regulation are upregulated in the allotetraploids (147), which may contribute to the hybrid vigor observed in the allotetraploids. Although the underlying mechanisms are unknown, one possibility is modulation of a few key regulators in the allotetraploids that may control downstream genes in various biological pathways (146,147) such as photosynthesis and metabolism (Z. Ni & Z.J. Chen, unpublished).

Alternatively, *cis*- and *trans*-acting effects involving regulatory sequence changes (see below), chromatin modifications, and RNA-mediated pathways (25) (Figure 2*a*) may explain dominance, overdominance, and epistasis. The interactions between the diverged orthologous protein products may determine repression or activation of progenitors' genes in allopolyploids (Figure 2*c*) of *Arabidopsis* (148), cotton (2), *Senecio* (60), and wheat (59,69), interspecific hybrids (151) in *Drosophila*, intraspecific diploid hybrids in maize (6,54,140), and sexdependent gene regulation in *Drosophila* (46,120). These mechanisms are not mutually exclusive, and the diverged protein-protein and protein-DNA interactions in allopolyploids may trigger repression of the protein-coding genes and rDNA loci derived from one progenitor (e.g., *A. thaliana*) via chromatin modifications (26,73,146,148) or novel expression patterns leading to hybrid vigor (Figure 2*c*).

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CIS- AND TRANS-ACTING EFFECTS ON TRANSCRIPTIONAL REGULATION IN ALLOPOLYPLOIDS

Stable allopolyploids provide an excellent system for testing *cis*- and *trans*-acting effects because a common set of protein factors is present in the same allotetraploid cells. After the unification of the distinct genomes, differences in *cis*- and *trans*-regulation contribute to changes in the expression of orthologs that become homoeologous pairs in the allopolyploid or interspecific nucleus (146,151). *Cis*-regulatory divergence directly acts on single genes or localized chromatin domains such as promoters or enhancers and may result in asymmetric accumulation of homoeologous transcripts in allopolyploids.

There is evidence for *cis*- and *trans*-effects on orthologous or homoeologous genes in the allotetraploids (146) and interspecific hybrids (33,151). Differential expression of progenitors' genes in Arabidopsis allopolyploids (73,148) and interspecific hybrids (33), Drosophila interspecific hybrids (151), and maize diploid hybrids (138) is mainly caused by cis-regulatory changes. Progenitor-specific differences in expression in the same cells are most likely due to allelic or epigenetic differences. In contrast, expression divergence due to alterations in trans-regulatory hierarchies should result in two kinds of expression changes. The first is a difference in the sum of homoeologous mRNAs compared with the mid-value of the two parents or nonadditive gene expression. Indeed, the divergently expressed orthologs comprise ~68% of the genes that were expressed in a nonadditive fashion in two allotetraploids (146, 147), implicating *trans*-acting effects. The second is a change in the ratio of homoeologencoded mRNAs in an allopolyploid compared with the ratio of the two orthologs in a 1:1 mixture of the parental mRNAs (147). Such a difference would demonstrate a regulatory interaction between the parental genomes (for example, the failure of an interspecific heterodimer to activate transcription) (25) (Figure 2c). A regulatory hierarchy (12) model suggests that trans-regulatory differences predominate in allopolyploids (25).

The species-specific expression patterns observed in Arabidopsis allotetraploids (147) may result from sequence divergence at regulatory elements during the ~6 million years that separate the parental species. Cis- and trans-acting regulation and epigenetic modifications of homoeologous genes may change regulatory interactions in a biological pathway (Figure 5a). This has been demonstrated in a subset of genes controlling flowering-time variation. A. arenosa and A. thaliana (Ler) diverged in flowering habits probably because of selective adaptation to cold and warm climates (108,123), respectively. Natural variation of flowering time is largely controlled by two epistatically acting loci, namely, FRIGIDA (FRI) (65) and FLOWERING LOCUS C (FLC) (102,131). FRI upregulates FLC expression that represses flowering in A. thaliana. A. thaliana has a nonfunctional AtFRI (65), whereas A. arenosa FRI (AaFRI) is functional (146). Compared with A. thaliana (AtFLC), A. arenosa FLC (AaFLC) loci possess deletions in the promoter and first intron that are important to cisregulation of FLC expression. In resynthesized allotetraploids, AaFRI complements nonfunctional AtFRI and interacts in trans with AtFLC, making the synthetic allotetraploids winter annual in a dosage-dependent manner. AaFRI acts on AtFLC in trans and on AaFLC in cis because A. thaliana FRI is nonfunctional. The different effects of AaFRI on AtFLC and AaFLC loci are likely dependent on the sequence divergence in their cis-regulatory elements (e.g., deletions in the promoter and first intron). AtFLC and AaFLC upregulation is mediated by H3-K9 acetylation and H3-K4 methylation, suggesting a role of FRI in locus-specific chromatin modifications (146).

Although our model (Figure 5*a*) simplifies the flowering pathway that involves >80 genes (80), it offers one explanation of the fate of orthologous genes involved in biological pathways during allopolyploidization. Many orthologous genes might have diverged in their *cis*-regulatory elements that confer strong or weak, dominant or recessive alleles, tissue-specific

expression, and/or developmental regulation. The regulatory networks may be reset by *cis*- and *trans*-acting effects via chromatin modification immediately after allopolyploidization. Over generations, genetic and epigenetic changes are subject to selection and adaptation, and additional genes (e.g., MAF, a FLC-MAF family member of MADS-box genes, in *A. suecica*) (146) may be activated for allopolyploids to occupy an environmental niche. A similar mechanism may be responsible for the functional diversification of orthologous genes in developmental regulation of gene expression, a phenomenon known as subfunctionalization of duplicate genes (88). Flowering time directly affects plant reproduction and adaptation. Therefore, sequence evolution and epigenetic regulation play interactive and pervasive roles in mediating the regulatory incompatibilities between divergent genomes, leading to natural variation and selective adaptation during allopolyploid evolution.

POST-TRANSCRIPTIONAL REGULATION IN ALLOPOLYPLOIDS

Some gene expression variation observed in polyploids may be controlled at the level of posttranscriptional regulation (70). Silencing a duplicate copy of homoeologous RNA in polyploids may be part of an RNA-mediated pathway similar to cosuppression (67,94) or RNAi (42). Silencing of transgenes is correlated with transgene dosage in *Drosophila* (113) and ploidy levels in *Arabidopsis* (105). Activation of Wis 2–1A retrotransposon in the newly synthesized wheat allotetraploids drives the readout transcripts from adjacent sequences including the antisense or sense strands of known genes, leading to the silencing or activation of respective corresponding genes (70). RNA-mediated silencing of duplicate genes in polyploids is a developmental strategy. Production of progenitor-dependent RNA transcripts may be associated with mRNA accumulation and stability during growth and development (55). For example, a subset of genes involved in mRNA stability displayed expression variation in allotetraploids (E. Kim & Z.J. Chen, unpublished). *CCR4*, a gene involved in RNA stability and degradation in yeast and animals (34), is differentially accumulated in leaves and flower buds, suggesting a role of RNA stability in transcript accumulation in allopolyploids.

Over time, species may have adapted to spatial and temporal regulation of RNA transcripts including mRNAs, small RNAs, and additional noncoding RNA transcripts that could accumulate nonadditively in the allotetraploids (Figure 5b). The small RNAs may serve as negative regulators for the expression of target genes originating from two parents. First, the loci encoding miRNAs and siRNAs may diverge during the evolution of the progenitors as observed in the FRI and FLC loci (146). Sequence divergence in promoter regions and cisacting elements leads to the expression variation when these loci are present in the same cell nuclei. Alternatively, differential expression of trans-acting factors may cause gene expression changes in the allopolyploids, as predicted (25). Second, antiviral RNAi genes involved in the biogenesis of small RNAs such as dicers, Argonaute, and RNA-dependent RNA polymerases (9,155) generally diverge faster than other proteins during evolution (106). Combining two divergent proteins in the allotetraploids may alter enzymatic activity and specificity. As a result, different pools of small RNAs could accumulate in the allopolyploids. Third, natural sense and anti-sense transcripts and other read-through transcripts may participate in defense mechanisms (16) and may be accumulated differently in the progenitors. These transcripts affect the expression of neighboring loci as well as other loci via trans-acting effects. Fourth, mRNA transcript abundance is different in the progenitors. Although the coding sequences are very similar among *Arabidopsis* and its related species, sequences at the noncoding regions (5'- and 3'-ends) diverge relatively rapidly (L. Tian, M. Ha & Z.J. Chen, unpublished), which may affect processing and stability of RNA transcripts (55). Finally, each species is differentiated by the presence or absence of species-specific repetitive DNA sequences, including transposons that may affect the chromatin structure and expression of their neighboring genes. Differences in DNA replication and perturbation of chromatin structures

among different species may induce the release of transposons and aberrant RNA transcripts that cause "genomic shock" and many downstream effects, as previously predicted (25,98).

DEVELOPMENTAL REGULATION OF ORTHOLOGOUS AND HOMOEOLOGOUS GENES

During polyploid evolution, both copies of orthologous genes may remain if dosage effects are advantageous (142), or one copy of the gene duplicate may evolve a novel function via neofunctionalization (89). Alternatively, both copies may diverge their functions or expression patterns in different organs or tissues via subfunctionalization (88). Indeed, silenced rRNA genes in vegetative tissues are reactivated during flower development (27). In a survey of 40 genes in cotton, 10 genes (25%) display unequal expression in allotetraploids and exhibit organ-specific expression patterns (2). For 5 genes, the A-subgenome loci are expressed higher than the D-homoeologous loci, whereas for the other 4 genes, the D-subgenome loci are expressed higher than the A-homoeologous loci. For some homoeologous gene pairs, one locus (e.g., *AdhA*) is silenced in one organ, whereas the other locus is silenced in another organ. This silencing scheme is genotype-independent and occurs in both synthetic and natural cotton allotetraploids (3), suggesting rapid subfunctionalization of duplicate genes and stable maintenance of tissue-specific expression patterns during evolution.

Although the mechanisms for developmental control of the expression of orthologous genes are unclear, developmental regulation of orthologous genes immediately after allopolyploid formation suggests that duplicate genes provide genetic robustness against null mutations (52) and dosage-dependent selective advantage (15,142). Moreover, immediate divergence in the expression of orthologous genes in allopolyploids provides a virtually inexhaustible reservoir for generating genetic variation and phenotypic diversification, which facilitates natural selection and adaptive evolution.

ODD AND EVEN DOSAGE EFFECTS ON GENE REGULATION IN POLYPLOIDS

Dosage-dependent gene regulation shows odd and even effects, which may affect additive and nonadditive gene regulation in polyploids. Using B-A chromosome translocation lines in maize, Birchler and his colleagues (53) generated a series of lines with different doses of A chromosomes that could be used to measure gene expression in response to changes in chromosome dosage. Gene expression levels are generally positively correlated with the dosage of the genes or chromosomes in these lines. However, the expression levels of ~10% genes are either reduced or negatively correlated with odd chromosome dosages (e.g., one, three, and five). One possibility is that dosage-dependent gene regulation is associated with chromosome pairing because one or more copies of chromosomes in odd dosages cannot pair properly.

The odd and even effects on gene regulation are also observed in the study of transgene expression in diploid and triploid hybrids derived from the crosses of diploid or tetraploid plants with a diploid strain containing a single copy of a transgenic resistance gene in an active state (105). The expression of the transgene is reduced in the triploids compared with the diploid hybrids, leading to the loss of the resistant phenotype at various stages of seedling development in some individuals. The reduction of gene expression was reversible under selective tissue culture conditions. This type of suppression was observed for a single-copy insert in the absence of other *trans*-acting copies of the transgene and is therefore different from homology-dependent gene silencing. An increase in ploidy or chromosome dosage can give rise to epigenetic gene silencing, generating stochastic variations in gene expression patterns. Although the expression of the transgene in a haploid or a pentaploid was not studied, odd ploidy may result in a new type of epigenetic repression. The expression of the transgene is

repressed only in the triploids in which one set of chromosomes is likely not paired or improperly paired.

Ploidy-dependent gene regulation suggests a sensing mechanism for gene dosage and DNA content via chromosome pairing. Although somatic pairing has not been documented in plants, such transient pairing has been observed in humans, *Drosophila*, and yeast (101). Homologous pairing has been implicated in transvection, position-effect variegation, and transgene gene silencing (5,61,113), all of which involve alterations in gene expression.

PARAMUTATION-LIKE EFFECTS IN POLYPLOIDS

Paramutation is the result of heritable changes in gene expression that occur upon interaction between alleles (21). The phenomenon was first discovered in plants and later found in many other organisms including mammals (mouse and human) (21,121,134). The paramutagenic allele induces the change in the expression state of the paramutable allele. A paramutation-like phenomenon was also discovered in the tetraploid plants containing active and inactive transgene alleles of hygromycin phosphotransferase (HPT) (104). Active alleles that are trans-inactivated by their silenced counterparts are observed in tetraploid but not in diploid plants, and this occurred only in progeny resulting from self-fertilization of plants heterozygous for the active and inactive HPT allele. The occurrence of transgene paramutation only in tetraploid plants indicates that active and inactive alleles go through meiosis together. This led to the hypothesis of pairing-based *trans*-inactivation. This predication is consistent with observations in tetraploid tomato, where the frequency of paramutation of a specific paramutagenic allele at the *sulfurea* locus is different between diploid, triploid, and tetraploid plants and depends on the ratio of paramutagenic to paramutable alleles (57). This suggests a counting mechanism for polyploidy-dependent paramutation, which may be similar to that for X-chromosome inactivation (75).

A paramutation-like phenomenon occurs in the progeny of genetic crosses between heterozygotes and between heterozygotes and wild-type mice independent of gender combination (121). The phenomenon is speculated to be associated with aberrant RNAs resulting from the paramutagenic allele that are packaged in sperm and cause paramutation upon transmission to the next generation. Indeed, paramutation depends on a RNA-dependent RNA polymerase; the *rdr101* mutation prevents paramutation in maize (4). However, paramutation in *Arabidopsis* tetraploids is probably not associated with RNA because *trans*activation does not occur in the F₁ generation (104). Moreover, crosses of decrease in the DNA methylation (*ddm1*) mutant with a paramutable tetraploid do not change paramutation phenotypes in the F₁ or F₂ but do in the F₃ family, which is consistent with the gradual loss of DNA methylation by *ddm1*. The data suggest that methylation occurs later, and is speculated to occur during physical contact of the epialleles during meiosis and after the silencing is established (104). Alternatively, sorting out pairing between homologous and homoeologous chromosomes in polyploids may require a few more rounds of meiosis.

Many paramutation phenomena are associated with repeated sequences (21,134). Multicopy genes or repetitive intergenic regions are a major trigger for the formation of silenced chromatin. Repeated sequences, whether inverted or tandem, can give rise to the production of dsRNA, an important trigger for RNA silencing as well as heterochromatin formation (85). In addition, repetitive sequences are also able to associate physically with their homologs in nonmeiotic cells (134). It is conceivable that different repeat sequences originating in the progenitors may trigger abnormal siRNA production and heterochromation formation that are responsible for paramutation-like or other epigenetic phenomena in allopolyploids.

FERTILITY, SELF-INCOMPATIBILITY, AND CYTOPLASMIC-NUCLEAR INTERACTIONS IN ALLOPOLYPLOIDS

Incompatibility between alien cytoplasmic and nuclear genomes and between alien nuclear genomes is believed to be a barrier leading to reproductive isolation, speciation, and developmental abnormalities in vertebrates and plants (18,77,78,82,112). Breaking down this barrier is essential in forming a new polyploid species (78). Seed fertility may be controlled by a few genes or many genetic loci. Three imprinted genes, *PHERES1, MEIDOS*, and *MEDEA*, are silenced in allotetraploids in a dosage-dependent manner (68). Disrupting maternal imprinting of *AtPHERES1* and paternal imprinting of MEDEA may reduce seed viability in the allopolyploids. Imbalance of paternally and maternally imprinted genes in the endosperm may also cause reproductive failures (20).

Another factor affecting seed fertility is the breeding system. Self-incompatibility is a mechanism for preventing inbreeding in many plant species (22). The sporophytic self-incompatibility system in the family Brassicaceae has been used as a model system to study mating system evolution in plants (99,108). *A. arenosa* is an outbreeder (self-incompatible), and *A. thaliana* is an inbreeding plant (self-compatible). They diverged from the same ancestor ~6 Mya (67,108). However, the natural allotetraploid *A. suecica* and the resynthesized allopolyploids are self-compatible (Figure 6), suggesting that the mating system switches immediately following polyploidization. The loss of self-incompatibility in the first generation of allotetraploids is not caused by the segregation of S-alleles in the allotetraploids because all possible alleles are present. The data suggest rapid epigenetic changes in the expression of the genes important to self-incompatibility, probably including the well-characterized loci encoding S-locus receptor kinase (SRK) and S-locus cysteine-rich (SCR) proteins in Brassicaceae (109,127).

In many cases, polyploidization converts self-incompatible diploids into self-compatible tetraploids in *Nicotiana* and *Solanum* with a gametophytic system, and some allopolyploids become self-compatible regardless of the mating types of their parents (18,49,82). Selfing in the allopolyploids may have an advantage for adapting new allopolyploids because of increased levels of heterozygosity. Inbreeding depression (1/18 or ~5% of homozygosity in one selfing generation) in allopolyploids is relatively low compared with that in a diploid (50%). An extreme form of reproductive modification is apomixis that is commonly associated with polyploidy (18,49). Resynthesized allopolyploids may be released from reproductive failure if they are capable of vegetative or seed apomixis.

ADAPTIVE EVOLUTION AND EXPRESSION DIVERGENCE BETWEEN DUPLICATE GENES

Theoretical prediction suggests that one copy of a gene duplicate would become lost by accumulation of deleterious mutations over an evolutionary timescale (87). Evidently, many duplicate genes are retained during evolution, and the redundancy conferred by duplicate genes may facilitate species adaptation (107) and genetic robustness (52) against changes in environmental conditions and developmental programs. Gene expression analyses indicate that duplicate genes offer genetic robustness against null mutations in yeast (52) and tend to experience expression divergence during development and to evolve faster between *Drosophila* species and within yeast species than single-copy genes (51,83). Using all duplicate gene pairs derived from a recent WGD (14,17,144) and gene expression microarrays in *A. thaliana* (126), Ha et al. (56) found that expression divergence between gene duplicates is significantly higher in response to external stresses than to internal developmental changes. Rapid divergence between gene duplicates in response to abiotic

and biotic stresses may facilitate subfunctionalization (88), neofunctionalization (89), and the evolution of an adaptive mechanism to environmental changes (98,129). A relatively slow rate of expression divergence between the duplicates may provide dosage-dependent selective advantage (15,142) and enable organisms to fine-tune complex regulatory networks. Orthologous and homoeologous genes in allopolyploids may have similar evolutionary fates.

The interactions between cytoplasm-nuclear and nuclear-nuclear genomes in the allopolyploids may induce genomic shock (98) and general amphiplasty (110) that are manifested by differential accumulation of transcripts originating from divergent species, leading to transcriptome dominance and activation or silencing of one or both homoeologous loci through genetic and/or epigenetic mechanisms. As a consequence, allopolyploids display hybrid vigor, flowering-time variation, inbreeding, apomixis, and selective advantage. Over time, orthologous or homoeologous loci in the allopolyploids may diverge their functions via neofunctionalization and subfunctionalization (88,89), as predicted for the paralogous loci (see Sidebar: Adaptive Evolution and Expression Divergence Between Duplicate Genes).

In polyploid populations of separate origin, one population may lose function from one copy of an orthologous gene, while a second population may lose function from a second copy of this ortholog. This "reciprocal silencing" of duplicated genes in polyploid genomes would ultimately lead to hybrid lethality (8), promoting reproductive isolation and the origin of new species. Following this model, the stochastic silencing and subfunctionalization of orthologous genes in different lineages of allopolyploids (148) may play a major role in the origin of new species. Together, these mutually inclusive mechanisms may contribute significantly to the adaptation potential, domestication, and evolution of polyploid plants.

SUMMARY POINTS

- 1. Allopolyploidy induces a wider range of gene expression changes than autopolyploidy, suggesting hybridization has greater effects on gene expression and phenotypic variation than genome doubling.
- 2. Genetic changes occur immediately after interspecific hybridization or allopolyploidization. These include genome-specific sequence elimination, as observed in *Triticum* and *Tragopogon* allopolyploids and chromosomal rearrangements in *Brassica* allotetraploids. *Arabidopsis* and cotton (*Gossypium*) allotetraploids show very low levels of genomic changes.
- **3.** Approximately 15–43% of the transcriptome diverged in expression between *A. thaliana* and *A. arenosa* (split ~6 Mya). The genes that display species-specific expression patterns are preferentially subject to nonadditive regulation in the resynthesized allotetraploids, indicating modulation of transcriptome divergence between the two progenitors.
- **4.** The genes that are highly expressed in *A. thaliana* are repressed in the resynthesized allotetraploids, which is correlated with the nucleolar dominance and overall suppression of *A. thaliana* phenotypes in resynthesized and natural allotetraploids.
- **5.** Some genes that display nonadditive expression are developmentally regulated, which may lead to subfunctionalization of these homoeologous genes in the allopolyploids.

- **6.** Establishment of gene expression patterns in the resynthesized allotetraploids is usually rapid and stochastic and in some cases may require several rounds of meiosis (by selfing).
- 7. Most nonadditively expressed genes in the resynthesized allotetraploids are present in both parents, suggesting an epigenetic cause and *cis* and *trans*-acting effects on nonadditive gene expression.
- 8. Sequence divergence in regulatory elements between the progenitors may cause genetic and epigenetic effects on expression of the genes involved in a biological pathway. *Trans*-acting differences exert their effects on *cis*-regulatory elements via epigenetic mechanisms, indicating interrelated regulatory interactions between genetic and epigenetic effects.
- **9.** Reprogramming gene expression patterns in allopolyploids via chromatin modifications and RNA-mediated pathways may cause changes in adaptive traits such as flowering-time variation, self-incompatibility, and hybrid vigor.
- **10.** Although the majority of data are documented from studies of resynthesized allotetraploids, similar changes in genomic organization and gene expression may occur in the natural allopolyploids and facilitate polyploid evolution and speciation.

FUTURE ISSUES

Addressing the following questions is essential to illuminate new insights into molecular and evolutionary impact on polyploid formation and speciation.

- 1. How and why are genetic dominance and nonadditive expression established rapidly in resynthesized allopolyploids, and how are these molecular processes altered or maintained in natural allopolyploids?
- 2. How do changes in gene expression contribute to intricate regulatory hierarchies and complex biological pathways leading to hybrid vigor and adaptive traits in allopolyploids?
- **3.** Why are ploidy changes more deleterious in animals than in plants, and why are polyploid cells often carcinogenetic in animals?

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Glossary

Polyploid

an individual or cell that has more than two sets of chromosomes

Allopolyploid

a polyploid that contains two or more sets of genetically distinct chromosomes, usually by hybridization between different species

Autopolyploid

a polyploid created by the multiplication of one basic set of chromosomes

Ploidy

the number of basic chromosome sets

Homologous

genes or structures that share a common evolutionary ancestor

Orthologous

chromosomes or genes in different species that have evolved from the same ancestor

Homoeologous

chromosomes or genes in related species that are derived from the same ancestor and coexist in an allopolyploid

Segmental allopolyploid

An allopolyploid that contains some homologous segments between homoeologous chromosomes, in which meiotic pairing occurs between homologous chromosomes as well as homoeologous chromosomes

Paleopolyploid

an ancient polyploid that undergoes chromosomal rearrangements, gene loss, and mutations and eventually becomes diploidized such that it behaves cytogenetically as a diploid

Nonadditive gene expression

levels of gene expression in an auto- or allopolyploid are not equal to the sum of progenitors, suggesting gene activation including dominance, overdominance, or gene repression

Apomixis

only one parent (usually female) contributes genes to the offspring

Aneuploid

an individual in which the chromosome number is not an exact multiple of the typical haploid set for that species

Paralogous

two or more genes in the same species that share a single ancestral origin

Epigenetic

heritable changes in gene expression without changes in primary DNA sequences

Genomic shock

release of genome-wide chromatin constraints of gene expression including activation of transposons in response to genomic, environmental, and physiological changes

X-chromosome inactivation

during mammalian development, the repression of one of the two X chromosomes in the somatic cells of females as a method of dosage compensation

Gametic imprinting

the expression of a gene depends on its parental origin in the offspring

Paramutation

heritable changes in gene expression due to allelic interactions

Homology-dependent gene silencing

repression of gene expression by homologous DNA sequences such as transgenes or endogenous loci

Amphiplasty

alterations of chromosomal morphology in interspecific hybrids or amphidiploids

Neofunctionalization

gain of a novel function (or expression pattern) from one copy of the gene duplicate

Subfuncationalization

functional (or expression) divergence of gene duplicate from the ancestral gene (e.g., tissue-specific expression of gene duplicate)



Figure 1.

Illustration of auto- and allopolyploids. For simplicity, only one pair of homologous chromosomes (either in red or blue) is shown in a diploid. (*a*) Formation of an autotetraploid by doubling a basic set of chromosomes. A triploid (not shown) can be formed by hybridization between a diploid and an autotetraploid. (*b*) Formation of an allotetraploid by interspecific hybridization followed by chromosome doubling. (*c*) Formation of an allopolyploid by fusion of unreduced gametes in two diploid species. (*d*) Formation of an allotetraploid by interspecific hybridization between two autotetraploid species. (*e*) An allotetraploid (*left*) may become a segmental allotetraploid (*right*) if homoeologous chromosomes contain some homologous chromosomal segments.



Figure 2.

Hypotheses to explain changes in genome structure and function in polyploids. (a) Genetic and epigenetic changes in polyploid genomes. Genetic changes include sequence mutations and chromosomal instabilities (deletion/insertion, translocation, transposition, etc.). Epigenetic changes may occur at transcriptional and post-transcriptional levels. Transcriptional regulation is associated with the formation of heterochromatin (black segments in red chromosomes) and euchromatin (*irregularly-shaped long arms of blue chromosomes*), leading to gene silencing or activation. Chromatin modifications [through multiple mechanisms including RNA interference (RNAi)] may transcriptionally silence or activate mobile elements, protein-coding genes, and rRNA genes that are responsible for novel variation in allopolyploids. Posttranscriptional regulation involves RNAi, RNA processing, and stability. AA: diploid genome; AAAA or BBBB: autotetraploid; AABB: allotetraploid. Blue segments in red chromosomes or red segments in blue chromosomes in an allotetraploid indicate nonreciprocal exchanges between homoeologous chromosomes. White segments in the red chromosomes indicate a deletion. (b) A dosage-dependent (12) or "rheostat" (102) model suggests additive effects of gene expression in a diploid (upper panel) and tetraploid (lower panel). The duplicate loci in the tetraploid provide additional levels of controls for gene expression. The levels of gene expression are shown as "off" (0) and by gray arrows (1, 2, and 3), and the maximum level (2) in diploid and 4 in tetraploid) is shown by green arrows. This model may explain dominant effects of hybrid vigor but does not explain overdominant performance or novel variation in allopolyploids. (c) A regulatory compatibility model suggests that regulatory factors (proteins) produced from orthologous genes generate incompatible heterologous products (heterodimers between red squares and blue circles). Alternatively, heterologous proteins may perform better than homologs (homodimers between red squares or blue circles), which may explain overdominant effects and hybrid vigor. (d) Hypotheses for testing additive and nonadditive gene regulation in Arabidopsis allotetraploids. The null hypothesis (Ho) is that gene expression levels in an allotetraploid (Allo) are equal to the sum of two progenitors, A. thaliana

autotetraploid (At4) and *A. arenosa* (Aa). Typical seedling leaves in At4, Aa, and Allo are shown. Ha: alternative hypothesis.





Stable allotetraploid progeny (2n = 4x = 26)



Figure 3.

Production of *Arabidopsis* allotetraploids. (*a*) Schematic karyotypes of F_1 nascent allotetraploids and their progenitors. Seedlings of the two progenitors, *A. thaliana* autotetraploid (At4) and *A. arenosa* tetraploid (Aa), are shown. (*b*) Phenotypic variation was infrequently observed in the F_1 allotetraploids but was very common in the segregating populations (F_2) (*A*–*D*). *A* and *D*: *A. arenosa*-like, *B*: *A. thaliana*-like, and *C*: intermediate. (*c*) Independent lineages of allotetraploids are produced by selfing multiple F_1 lines. Allopolyploids self-pollinate spontaneously (or can be manually self-pollinated), and progressive inbreeding occurs in advanced generations. Karyotypes in a meiotic cell of a resynthesized allotetraploid (Allo733 in the fifth generation) show 5 pairs of *A. thaliana* centromeres (At Cen, *red*) and 8 pairs of *A. arenosa* centromeres (Aa Cen, *green*) (147) (reproduced with kind copyright permission of the Genetics Society of America). Seedling of

Allo733 is shown. All plants were photographed when they were 4–5 weeks old. The size bars indicate 3 cm.

Chen



Figure 4.

(*a*) Phenotypic variation of the plants after five generations of selfing. The plants include two parents, *A. thaliana* autotetraploid (At4) and *A. arenosa* (Aa), six allotetraploids (S–1 to S–6), and a natural allotetraploid, *A. suecica* (As). Allotetraploids have the same chromosome number (Figure 3c and data not shown) but display phenotypic and flowering-time variation. Some plants (S5–5 and S5–6) resemble *A. suecica*, whereas others (S5–1 to S5–4) show novel phenotypes. Differential expression of parental genes may contribute to the phenotypic variation of *Arabidopsis* allotetraploids. (*b*) Evidence for epigenetic silencing in resynthesized allopolyploids. *A. suecica* (As, *white flower*) is the naturally occurring allotetraploid derived from *A. thaliana* (not shown) and *A. arenosa* (Aa), which have white and pink flowers, respectively. The flower colors in the third generation of resynthesized allotetraploids (S3–1,

2, and 3) segregate from all white (S3-1) to all pink (S3-3). Variegated flower colors in the same inflorescence branch (S3-2) indicate epigenetic regulation of genes controlling flower color pathways. The same-size bars are 50 mm in (*a*), except for At4, and 5 mm in (*b*), except for As.

Figure 5.

Models for transcriptional and post-transcriptional regulation of nonadditive gene expression in allopolyploids. (a) Cis- and trans-acting effects on transcriptional regulation of nonadditive gene expression in allotetraploids. I. During evolution, changes in cis-regulatory elements may confer "strong/dominant" and "weak/recessive" loci. II. Combination of these loci in the same allotetraploid cells induces cis- and trans-acting effects. Because A. thaliana FRI (AtFRI) is nonfunctional, A. arenosa FRI (AaFRI) trans-activates A. thaliana FLC (AtFLC) and cisregulates one of the A. arenosa FLC (AaFLC1) loci, leading to overdominance or very late flowering in resynthesized allotetraploids. The cis-regulation and trans-activation are maintained by histone acetylation (Ac in a small purple circle) and methylation (Me in a small orange circle). Functional AaFRI is in a large red circle, and nonfunctional A. thaliana FRI (*atflc*) is in a white small circle with a slash line. The straight and inclined red arrows indicate cis- and trans-acting effects, respectively, by AaFRI. FLC loci are in oval circles. Blue and green indicates active AtFLC1 and AaFLC1, respectively, and gray indicates silenced AaFLC2. The differences in the expression of three FLC loci may reflect *cis*-regulatory variation (146). III. Cis- and trans-acting interactions and epigenetic regulation of FRI and FLC loci may facilitate selection for flowering-time variation in resynthesized allotetraploids (indicated by a circular wheel). IV. In A. suecica, additional selection forces may enhance the expression of AaFLC1 (color change from dark green in II to yellow green in IV) and related genes such as MAFs (146), which facilitate adaptation to cold climate. This simple model may be generalized to explain a mechanism for regulatory interactions between divergent loci in other pathways in the allotetraploids (see text for details). (b) Nonadditive accumulation of mRNAs and small RNAs in allotetraploids (null hypothesis H_0 : 2 + 1 = 3). There is evidence that 15–40% of the genes are expressed differently between two closely related species of Arabidopsis (147), and a subset of those genes are subjected to nonadditive expression in the allotetraploids. Up- or downregulation of mRNAs may be caused by activation of genes originating in parent A, B, or both. We exclude the situation in which the expression of one

locus is compensated by its corresponding homoeologous locus. If the nonadditively accumulated transcripts are small RNAs, such as microRNAs (miRNAs), heterochromatic and transposon-related siRNAs (ht-siRNAs), *trans*-acting siRNAs (ta-siRNAs), and natural antisense (nat-siRNAs), they typically negatively regulate the expression of their target genes. The red and blue wave lines indicate RNA transcripts or small RNAs produced in species A and B, respectively. We hypothetically assign two RNA molecules for species A and one RNA molecule for species B. Note that the number of RNAs in the nascent allotetraploids (F_1) is not equal to 3 (2 + 1), suggesting nonadditive regulation (dominance and overdominance if larger than 3 or repression if smaller than 3). Possible changes (*dashed arrows*) in RNA composition and accumulation (column X) may occur in self-pollinating progeny, which are indicated by orange and green wave lines, respectively. According to the hypothetical outcome, the origins of small RNA upregulation (*red* in 1–3) and repression (*blue* in 4–6) are shown in the last column.

Figure 6.

Switching of self-incompatibility in resynthesized *Arabidopsis* allotetraploids. (*a*) Flower morphology of self-compatible *A. suecica*. (*b*) Self-incompatible *A. arenosa* and (*c* and *d*) two self-compatible synthetic allotetraploid lines. The altered flower organs (elongated stigma and relatively short stamens) in the synthetic allotetraploids (*c* and *d*) may contribute to a low fertility without manual pollination. Size bars (5 mm) are the same in (*a*) and (*b*) or in (*c*) and (*d*). (*e*) Diagram of cytoplasmic and nuclear interactions in resynthesized allotetraploids (Allos, 2n = 4x = 26). T or A and tt or aa denote cytoplasm and nuclear genomes, respectively. The cytoplasmic-nuclear genotypes for autotetraploid *A. thaliana* (At4) are Ttttt, tetraploid *A. arenosa* (Aa), Aaaaa; and allotetraploids, and either Taatt or Aaatt in reciprocal crosses. Blue and gray indicate *A. thaliana* and *A. arenosa* cytoplasms, whereas green, red, and yellow indicate nuclear genomes of *A. thaliana*, *A. arenosa*, and allotetraploids, respectively. *A. thaliana* is self-compatible (SC), whereas *A. arenosa* is self-incompatible (SI). The resynthesized allotetraploids (Allos) are SC. Fertile seeds are readily obtained from the cross using *A. thaliana* as a maternal parent and *A. arenosa* as a pollen donor, and the reciprocal cross is usually unsuccessful (31).