



Is post-polyploidization diploidization the key to the evolutionary success of angiosperms?

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Advances in recent years have revolutionized our understanding of both the context and occurrence of polyploidy in plants. Molecular phylogenetics has vastly improved our understanding of plant relationships, enabling us to better understand trait and character evolution, including chromosome number changes. This, in turn, has allowed us to appreciate better the frequent occurrence and extent of polyploidy throughout the history of angiosperms, despite the occurrence of low chromosome numbers in some groups, such as in *Arabidopsis* (*A. thaliana* was the first plant genome to be sequenced and assembled). In tandem with an enhanced appreciation of phylogenetic relationships, the accumulation of genomic data has led to the conclusion that all angiosperms are palaeopolyploids, together with better estimates of the frequency and type of polyploidy in different angiosperm lineages. The focus therefore becomes when a lineage last underwent polyploidization, rather than simply whether a plant is ‘diploid’ or ‘polyploid’. This legacy of past polyploidization in plants is masked by large-scale genome reorganization involving repetitive DNA loss, chromosome rearrangements (including fusions and fissions) and complex patterns of gene loss, a set of processes that are collectively termed ‘diploidization’. We argue here that it is the diploidization process that is responsible for the ‘lag phase’ between polyploidization events and lineage diversification. If so, diploidization is important in determining chromosome structure and gene content, and has therefore made a significant contribution to the evolutionary success of flowering plants. © 2015 The Linnean Society of London, *Botanical Journal of the Linnean Society*, 2016, **180**, 1–5.

ADDITIONAL KEYWORDS: chromosome number – flowering plants – genome downsizing – genome rearrangement – genomics – lag phase – polyploidy – WGD radiation lag-time model.

POLYPLOIDY AND DIPLOIDIZATION

Polyploidy, or whole genome duplication (WGD), is a frequent phenomenon in plants, especially in flowering plants. It has been estimated that *c.* 15% of angiosperm speciation events involve a change in ploidy (neopolyploidy; Wood *et al.*, 2009) and that all flowering plants have experienced at least one WGD episode in their evolutionary history (palaeopolyploidy; Bowers *et al.*, 2003; Blanc & Wolfe, 2004; Van de Peer, Maere & Meyer, 2009; Jiao *et al.*, 2011). Ferns contain an even greater number of speciation events involving polyploidy (~31%; Wood *et al.*, 2009)

and they are also the group of plants with the highest reported chromosome number ($2n = c. 1440$; Abraham & Ninan, 1954). In ferns, multiple rounds of polyploidy occur apparently without the same diploidization processes that mask ancestral polyploidy in angiosperms (Leitch & Leitch, 2012). What role, if any, diploidization plays in the story of fern evolution is currently unknown. Ferns are relatively homogeneous in terms of developmental flexibility, morphological diversity and ecological specialization, at least when compared with angiosperms. In contrast, angiosperms exhibit a plethora of floral and vegetative forms that are often thought to account for their diversification and abundance relative to that of gymnosperms, ferns, lycopods and bryophytes. In

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particular, annual life histories are almost unknown outside the angiosperms (there are a few ferns that have managed this). Might diploidization following polyploidy, particularly following allopolyploidy (hybridization involving polyploidy), be a crucial factor in expanding evolutionary innovation versus relative evolutionary stasis?

EVOLUTIONARY DYNAMICS OF POLYPOIDS

As a result of their prevalence in plants, polyploids have been speculated to hold a selective advantage over diploids through the evolution of novel genetic (and indeed genomic) variation (Soltis & Soltis, 2000; Leitch & Leitch, 2008; Flagel & Wendel, 2009). In theory, duplicated genes provide the substrate for mutation-driven evolution of new copies, as a result of freedom from selective constraints. With multiple copies comes the potential for subfunctionalization and/or neofunctionalization, the two often being difficult to distinguish. The extent of neofunctionalization is currently unknown and is difficult to document empirically. However, subfunctionalization is a relatively common phenomenon in angiosperms. The origin of flowers, for instance, ostensibly requires the concerted function of various MADS box transcription factor complexes, and the evolution of such transcription factors has been attributed to ancient (i.e. as a result of palaeopolyploidy) and recent gene-specific duplications, with subsequent subfunctionalization of paralogous gene copies. It has become apparent in orchids, the most species rich and perhaps most florally diverse family of angiosperms, that subfunctionalization of duplicated B genes has led to the development of three petal-like organs: three outer tepals, two inner tepals and a highly modified lip (Mondragón-Palomino & Theißen, 2011; Mondragón-Palomino, 2013). Polyploids also have increased fixed heterozygosity, leading to increased heterosis and a higher tolerance of selfing (perhaps even promoting the evolution of self-compatibility), which leads to a tolerance of habitat fragmentation and population disturbance. In some cases, polyploids may also occupy new ecological niches or a broader range of niches compared with their diploid relatives. Collectively, these factors may contribute to the success of polyploids as invasive species (Pandit, Pocock & Kunin, 2011).

In contrast with the above advantages, polyploidy can also create a barrier to selection, as new mutations are masked by existing alleles, thereby ‘diluting the effects of new mutations’ (Stebbins, 1971). This depends on the dominance of new beneficial mutations versus the pre-existing allele; therefore, if it is at least partially recessive it will be masked, resulting in

inefficient selection (Stebbins, 1971; Otto, 2007), and leading to the idea that polyploids are ‘evolutionary dead-ends’ (Stebbins, 1950). The loss of some duplicate copies following polyploidy can cause a dosage imbalance and disruptions to gene networks – the gene balance hypothesis (Birchler & Veitia, 2007). This is particularly significant for genes that contribute to macromolecular complexes, which are often retained post-polyploidy in order to maintain a dosage-sensitive relationship (Conant, Birchler & Pires, 2014). Furthermore, recent polyploids typically have reduced fertility as a result of pairing problems at meiosis (Chester *et al.*, 2012; Yant *et al.*, 2013). They can also, on formation, experience ‘genomic shock’ from the combination of two disparate subgenomes in one nucleus, resulting in an elevated frequency of (retro)transposition (McClintock, 1984; Petit *et al.*, 2010) and chromosomal rearrangements that reduce fitness, potentially leading to extinction (Leitch & Leitch, 2008; Mayrose *et al.*, 2011). Newly formed polyploids are at low frequencies in populations, and there is strong selection pressure for self-compatibility to evolve – minority cytotype exclusion (Levin, 1975; Husband, 2000). When they first form, allopolyploids are typically, for many characters and traits, intermediate between their two parents, and they are in instant competition if they occur sympatrically with their parents. They may also lack an ecological niche and/or experience low rates of pollination as a result of no specific adaptations to a pollinator. It is a combination of these problems that often causes neopolyploids to go extinct, but, as soon as a polyploid population forms, there will be selection for particular better adapted genotypes that direct the trajectory of subsequent genome evolution. This includes selection for genotypes with increased fertility, genomic stability and better-balanced gene copies. These and other (see below) directional changes are among the most important pressures that lead to diploidization of the neopolyploid genome.

A comparison of diversification rates across angiosperms has led to the suggestion that (neo)polyploids are more likely to go extinct and less likely to speciate than diploids (Mayrose *et al.*, 2011), although there are sampling and analytical issues that make this a topic of much recent debate (see Soltis *et al.*, 2014). There are also arguments for polyploids not contributing to adaptive radiations *per se*, but rather polyploids simply arising through their immediate reproductive isolation from parental lineages (purely as a result of differences in chromosome number). Where recurrent polyploidization occurs between the same or different parental species, (non-adaptive) radiations can result (Gorelick & Olson, 2013). Even if neopolyploids do have higher extinction rates and make a lower contribution to recent species diversification, as has been

argued, all angiosperm species nonetheless have (often multiple rounds of) polyploidy in their ancestry; the ramifications of this are significant and an important focus of research.

THE ECOLOGICAL COST OF POLYPLOIDY

With a larger genome comes the ecological burden of needing more macronutrients to build nucleic acids, particularly nitrogen and phosphorus, the latter limiting in many natural environments (Vitousek *et al.*, 2010; Šmarda *et al.*, 2013). In addition, it has been shown that interactions between plant genome size and macronutrient availability influence plant distribution in semi-natural field experiments (Šmarda *et al.*, 2013; M. S. Guignard *et al.*, unpubl. data). Given the general trend towards genome downsizing following polyploidy (Leitch & Bennett, 2004) and the strong skew towards small genome sizes in angiosperms despite recurrent polyploidy and in comparison with other land plant lineages (Leitch & Leitch, 2012), it is probable that there is selection favouring smaller genomes in angiosperms (but see also Oliver *et al.*, 2007 for a neutral theory to explain the skew), thereby negating the effects of genome enlargement generated by polyploidy. However, the extent to which nitrogen and phosphorus availability influences genome size in the natural environment is a topic of debate, and further work is needed (Leitch & Leitch, 2008; Greilhuber & Leitch, 2013; Neiman *et al.*, 2013).

DIPLOIDIZATION IS NECESSARY FOR EVOLUTIONARY PERSISTENCE AND DIVERSIFICATION

Diploidization of the genome post-polyploidization is associated with neofunctionalization, subfunctionalization and genome downsizing. Removal of extra DNA (often repetitive DNA) and extraneous gene copies occurs through recombination-based deletion and other mechanisms, whilst retaining duplicated genes, some of which may have new or altered functions. Selection can then act on individuals with varying genome sizes, and those with smaller genome sizes may be favoured, particularly perhaps in nutrient-poor environments. In addition, diploidization has been associated with chromosome number reduction, potentially involving complex chromosomal rearrangements (Franzke *et al.*, 2011; Mándaková *et al.*, 2012). Chromosome reorganization can be so extensive such that, in some taxa, e.g. *Arabidopsis thaliana* (L.) Heynh. (Brassicaceae), which has had multiple polyploidy events in its ancestry (potentially the species is 48- or 96-ploid), the chromosome number has been reduced to a mere $n = 5$ pairs. Recent work in *Veronica* L. (Plantaginaceae) has suggested a link between increased

diversification and genome downsizing following polyploidy (Meudt *et al.*, 2015).

The WGD radiation lag-time model (Schranz, Mohammadin & Edger, 2012) postulates that species diversification often follows WGD events, but only after a 'lag phase' that can last up to several million years. This model explains the often observed pattern of a depauperate clade sister to a highly diverse one, with an observable time lag between the formation of polyploids and their subsequent diversification. Significant statistical support for this model has been garnered by Tank *et al.* (2015), who analysed nine well-documented ancient WGD events and demonstrated a non-random association between WGDs and a delayed increase in rates of diversification. It is likely that similar approaches will reveal more recent examples of a lag phase below family level (Tank *et al.*, 2015). Schranz *et al.* (2012) tied the context for the lag to 'later migration events, changing environmental conditions and/or differential extinction rates'. Tank *et al.* (2015) suggested it could represent unsampled extinct lineages or the evolution of complex key traits/innovations; however, they emphasized that there is a real need to study the causes and nature of the lag phase in greater detail, in terms of both genomics and ecology. Our hypothesis here is that this lag phase is the time required for diploidization to take effect and provide a polyploid clade with the potential to radiate.

Polyploidy is important for the generation of genetic and genomic novelty, but it also requires extensive genome reorganization in order for this evolutionary potential to be fully realized (i.e. 'diploidization'). Over intermediate timescales, up to tens of millions of years, selection may favour smaller genomes that have an ecological advantage, at the same time favouring genotypes that retain advantageous alleles in enlarged gene families. Genomic rearrangements that occur after polyploidy may also enable novel *cis*-acting gene responses and the accumulation of locally adaptive genes in linkage groups (Yeaman, 2013). In addition, De Smet *et al.* (2013) documented consistent patterns of gene deletion in neopolyploid genomes, indicating that genes controlling expression and those in balanced macromolecular complexes were preferentially retained. This process of turnover takes time, and almost certainly leads to novel patterns of expression during the removal of extraneous gene copies.

CONCLUSIONS

It is clear from global analyses of chromosome number and genome size in a phylogenetic context that, despite the current frequency (and the important legacy) of polyploidization in angiosperms, there is also an irrefutable role for diploidization after polyploidization

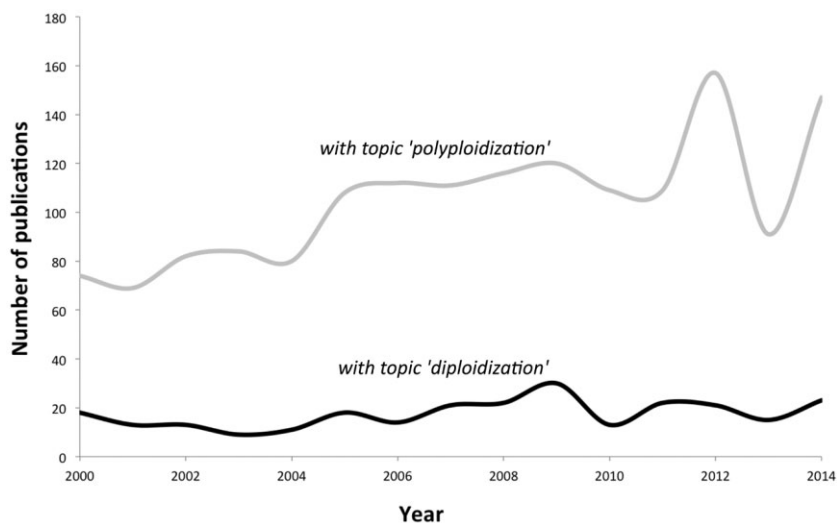


Figure 1. The number of publications with diploidization (black line) or polyploidization (grey line) in the title or keywords over the last 14 years. Source data: Web of Knowledge (Thomson Reuters).

has occurred. Our argument is that diploidization negates the disadvantages of polyploidy, rearranges genomes in novel ways and generates a higher level of genomic and transcriptomic variation upon which selection can act. Many genes return to their original copy number, thereby negating the effects of inefficient selection and the idea that polyploids are ‘evolutionary dead-ends’ (Stebbins, 1950). More sophisticated fine-tuning of expression and subfunctionalization can then enable novel phenotypic changes. The combination of high-throughput sequencing, cytogenetics and evolutionary developmental genetics with our best estimates of phylogenetic relationships will undoubtedly start to uncover the processes that have led to both ecological persistence and diversification of diploidized angiosperms. To test these hypotheses, we suggest physiological (stress) experiments on polyploids of different ages to examine potential ‘genomic plasticity’ enabled by the retention of increased numbers of transcription factors and genes controlling expression. Diploidization subsequent to polyploidization is an under-studied topic (see Fig. 1 for the number of publications on ‘diploidization’ versus ‘polyploidization’ in the last decade). Although the importance of diploidization is often acknowledged, botanists have never been in a better position to begin to answer exactly how post-polyploidization diploidization has contributed to the evolutionary success of the angiosperms.

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