Transposable Elements: Powerful Contributors to Angiosperm Evolution and Diversity

Keith R. Oliver*, Jen A. McComb, and Wayne K. Greene

School of Veterinary and Life Sciences, Murdoch University, Perth, Western Australia, Australia.

*Corresponding author: E-mail: K.Oliver@murdoch.edu.au.

Accepted: September 9, 2013

Abstract

Genome Biology and Evolutic

SMBE

Transposable elements (TEs) are a dominant feature of most flowering plant genomes. Together with other accepted facilitators of evolution, accumulating data indicate that TEs can explain much about their rapid evolution and diversification. Genome size in angiosperms is highly correlated with TE content and the overwhelming bulk (>80%) of large genomes can be composed of TEs. Among retro-TEs, long terminal repeats (LTRs) are abundant, whereas DNA-TEs, which are often less abundant than retro-TEs, are more active. Much adaptive or evolutionary potential in angiosperms is due to the activity of TEs (active TE-Thrust), resulting in an extraordinary array of genetic changes, including gene modifications, duplications, altered expression patterns, and exaptation to create novel genes, with occasional gene disruption. TEs implicated in the earliest origins of the angiosperms include the exapted *Mustang, Sleeper*, and *Fhy3/Far1* gene families. Passive TE-Thrust can create a high degree of adaptive or evolutionary potential by engendering ectopic recombination events resulting in deletions, duplications, and karyotypic changes. TE activity can also alter epigenetic patterning, including that governing endosperm development, thus promoting reproductive isolation. Continuing evolution of long-lived resprouter angiosperms, together with genetic variation in their multiple meristems, indicates that TEs can facilitate somatic evolution in addition to germ line evolution. Critical to their success, angiosperms have a high frequency of polyploidy and hybridization, with resultant increased TE activity and introgression, and beneficial gene duplication. Together with traditional explanations, the enhanced genomic plasticity facilitated by TE-Thrust, suggests a more complete and satisfactory explanation for Darwin's "abominable mystery": the spectacular success of the angiosperms.

Key words: TE-Thrust, hybridization, polyploidy, adaptation, speciation, domestication.

Introduction

The origin and extremely rapid diversification of flowering plants, which Darwin famously referred to as an "abominable mystery," is one of the most extraordinary, and still not yet fully explained, phenomena in evolutionary history. As the dominant plant taxon, angiosperms are estimated to contain at least 350,000 extant species (Soltis et al. 2008), placing them second only to insects in terms of species richness. This contrasts with their ancient woody competitors, the gymnosperms, which are apparently in stasis and comprise less than 1,000 species. Angiosperms are vastly diverse in form, from the hyphal-like strands of the endoparasitic Pilostyles and 1-mm-long single floating leaf of Wolffia to the giant banyan trees (*Ficus*) that may cover over a hectare and the >80 m tall eucalypts. Key characteristics of angiosperms are flowers with ovules in an enclosed ovary, double fertilization to produce both a zygote and a (usually) triploid endosperm to nourish the zygote, and the development of fruit-containing seeds. Such evolved reproductive features have been critical to the success of the angiosperms, which occupy a wide range of ecological niches and include all carnivorous and parasitic plants.

Although once said to be "junk," or "parasitic," DNA (Doolittle and Sapienza 1980; Orgel and Crick 1980), a recent large and rapid accumulation of evidence indicates that transposable elements (TEs) have been a significant factor in the evolution of a wide range of eukaryotic taxa (Bennetzen 2000; Kazazian 2004; Biémont and Vieira 2006; Feschotte and Pritham 2007; Böhne et al. 2008; Hua-Van et al. 2011). We have proposed TEs as powerful facilitators of evolution (Oliver and Greene 2009), formalized this proposal into the TE-Thrust hypothesis (Oliver and Greene 2011), and more recently, expanded and strengthened this hypothesis (Oliver and Greene 2012).

The TE-Thrust hypothesis has great explanatory power with regard to adaptation and evolution and was developed from

[©] The Author(s) 2013. Published by Oxford University Press on behalf of the Society for Molecular Biology and Evolution.

This is an Open Access article distributed under the terms of the Creative Commons Attribution Non-Commercial License (http://creativecommons.org/licenses/by-nc/3.0/), which permits non-commercial re-use, distribution, and reproduction in any medium, provided the original work is properly cited. For commercial re-use, please contact journals.permissions@oup.com

empirical evidence among the metazoans, principally mammals. It has offered an explanation for the great fecundity of some lineages and the paucity of species in other lineages, for stasis, and for "living fossils" (Oliver and Greene 2009, 2011, 2012). Owing to variable TE activity over time, TE-Thrust also suggests strong support for punctuated equilibrium (Eldredge and Gould 1972; Gould 2002). The TE-Thrust hypothesis posits that the genome-modifying effects of TEs can be either active or passive. Active gene/genome modification is due to transposition of TEs, which often occurs in bursts, whereas passive gene/genome modification is due to ectopic recombination between TEs of similar sequence, scattered throughout the genome (Oliver and Greene 2009, 2011, 2012).

Novel TEs/TE insertions can be acquired in germ lines by endogenous de novo synthesis (e.g., SINEs), de novo modification to resident TEs, de novo formation of chimaeras (e.g., SVAs, SINE-Variable number of tandem repeats-Alus; Wang et al. 2005), endogenization of viral sequences (Feschotte and Gilbert 2012), genomic perturbations such as hybridization or polyploidy (Kawakami et al. 2010; Parisod et al. 2010), and by horizontal transposon transfer, often between completely unrelated taxa (Schaack et al. 2010). Acquisitions of new TEs or reactivation of TEs in germ line genomes can result in intermittent bursts of TE activity, and these have been reported in various lineages, including metazoans (Gerasimova et al. 1985; Marques et al. 2005; Ray et al. 2008) and angiosperms (Naito et al. 2009; Lu et al. 2012; Palmer et al. 2012; El Baidouri and Panaud 2013).

A concept central to the TE-Thrust hypothesis is that either fixed or unfixed TEs in a lineage can facilitate adaptation and evolution by means of their various interactions with the genome that can create realizable intragenomic potential. Intragenomic potential is a continuum that ranges from adaptive potential to evolutionary potential. Adaptive potential, also termed capacitance (Pigliucci 2007), can be realized over periods of tens to hundreds of years, whereas evolutionary potential can be realized over thousands or millions of years (Oliver and Greene 2012).

In angiosperms, genome size and structure is largely determined by TEs because they often constitute the major fraction (up to 84%) of their DNA (table 1). Angiosperm genes in large and TE-rich genomes such as maize have been described as islands surrounded by seas of nested TEs (SanMiguel and Bennetzen 1998). Furthermore, the islands themselves are a moonscape of ancient and recent impacts from TEs. Among the retro-TEs, Copia-like and Gypsy-like long terminal repeat (LTR) elements are abundant in angiosperms, whereas Long Interspersed Elements (LINEs) and Short Interspersed Elements (SINEs), that are prominent in mammalian genomes, are less common. An important unanswered question for future investigation is the function and significance of the envelopeclass LTR retro-TEs found in angiosperms (Vicient et al. 2001) and at least one gymnosperm species (Pinus pinaster; Miguel et al. 2008), as these could possibly be the equivalent of the vertebrate endogenous retroviruses, which were prominent in the evolution of mammals (Feschotte and Gilbert 2012; Oliver and Greene, 2012), and are particularly active in the murid rodents (Gibbs et al. 2004; Maksakova et al. 2006). DNA-TEs found in angiosperms belong to several superfamilies and include CACTA, hAT, Harbinger, Mutator, Helitron, and Mariner-like elements (table 1). The generally high TE content of flowering plants, which includes active DNA-TEs and homogeneous LTR retro-TE populations, makes them highly suited for TE-Thrust, both active and passive. Indeed, the relative instability of angiosperm genomes compared with mammals in terms of gene movement and genome rearrangement (Bennetzen 2005; Kejnovsky et al. 2009), implies that TE-Thrust may be especially powerful in flowering plant lineages. We therefore see TE-Thrust as having much explanatory value, in addition to other accepted explanations, for the rapid diversification of the angiosperms, after their mysterious origin, early in the Cretaceous (135-90 Ma) and their rapid rise to dominance among the vascular plants 100-70 Ma.

In putting forward TE-Thrust as an important facilitator of evolution, we do not suggest that it is entirely universal or that other mechanisms of evolution are not significant. In fact, as we have noted previously (Oliver and Greene 2009, 2011, 2012), TE-Thrust, although very important in most extant taxa, is one of many possible facilitators of evolution, which include hybridization (Soltis PS and Soltis DE 2009), polyploidy/whole genome duplication (Van de Peer et al. 2009), recombination (Gaut et al. 2007), and horizontal gene transfer (Keeling and Palmer 2008). In some rare extant species belonging to reasonably fecund genera, TE-Thrust appears to now have little to do with ongoing adaptive potential or evolutionary potential, as such species currently have genomes that are largely devoid of TEs. An example among the angiosperms is the small 80 Mb genome of the recently sequenced bladderwort, Utricularia gibba, which is remarkable for having only 3% TE content, yet belonging to a successful genus comprising more than 200 species (Ibarra-Laclette et al. 2013). Significantly, the evolutionary history of this species has been marked by repeated rounds of whole genome duplication (Ibarra-Laclette et al. 2013), whereas the clade to which U. gibba belongs exhibits extreme mutation rates that are among the highest within the angiosperms (Müller et al. 2004). These, and possibly other factors, may account for its evolution and also for some current adaptive potential and evolutionary potential.

TE-Thrust Acts in Concert with Other Factors Widely Acknowledged as Promoting Angiosperm Diversity and Dominance

Hybridization and Polyploidy

Frequent tolerance of hybridization and polyploidy (with or without hybridization) are widely acknowledged factors

Table 1

TE Composition (%) of Representative Flowering Plant Genomes

			A. Dicc	otyledons				
Family	Rosad	eae	Vitaceae	Brassicaceae	Fab	aceae	Solana	iceae
Species	Malus x domestica	Fragaria vesca	Vitus vinifera	Arabidopsis thaliana	Glycine max	Medicago truncatula	Solanum lycopersicum	Solanum tuberosum
Genome size (Mbp)	742	240	487	125	1,115	375	900	844
Haploid chromosome number ^a	17	7	19	5	20	8	12	12
Type I: Retro-TEs								
LTR/Gypsy	25.2	6.0	14.0	5.2	29.5	1.4	19.7	15.2
LTR/Copia	5.5	4.6	4.8	1.4	12.5	2.4	6.3	3.8
LTR/other	0.4	3.8	_	_	_	9.6	35.8	33.2
LINE	6.5	0.2	0.6	0.9	0.25	3.4	0.4	0.7
SINE	_	0.06	_	_	_	0.1	0.2	0.3
Unclassified	_	_	_	_	_	_	_	
Total Retro-TEs	37.6	14.7	19.4	7.5	42.2	16.9	62.3	53.2
Type II: DNA-TEs								
CACTA	_	2.6	0.2	0.9	10.2	0.1	0.1	0.1
Helitron	_	0.07	_	5.6	0.5	0.2	_	
hAT	0.3	0.6	0.8	0.3	0.04	0.1	0.1	0.2
PIF/Harbinger	_	0.2	_	0.2	0.3	0.2	_	0.1
Tc1/Mariner	_	—	_	0.3	0.03	_	—	
Mutator	_	0.2	0.4	3.1	4.5	0.6	—	
Other		—		0.1	0.09		0.3	0.3
MITE/Tourist	0.6 ^b	1.6 ^b	_	—	0.3	0.1 ^b	—	_
MITE/Stowaway			_	_	0.5		_	
Unclassified	_	_	_	0.5	_	0.2	0.2	0.4
Total DNA-TEs	0.9	5.2	1.4	11.0	16.5	1.4	0.9	1.2
Unknown	3.9	0.9	0.7	—		—	—	
Total TEs	42.4	20.7	21.5	18.5	58.7	18.3	63.2	54.4

B. Monocotyledons

Family				Poaceae				Musaceae
Species	Triticum aestivum	Hordeum vulgare	Zea mays	Sorghum bicolor	Oryza sativa	Brachypodium distachyon	Setaria italica	Musa acuminata
Genome size (Mbp)	17,000	5,100	2,300	730	389	272	423	523
Haploid chromosome number ^a	7	7	10	10	12	5	9	11
Type I: Retro-TEs								
LTR/Gypsy	44.0	18.0	46.4	19.0	12.0	16.0	22.1	11.4
LTR/Copia	17.4	8.5	23.7	5.2	2.5	4.9	7.2	25.6
LTR/other	1.5	25.8	_	30.2	9.0	0.5	0.3	_
LINE	0.8	_	1.0	0.04	0.8	1.9	1.8	5.4
SINE	0.004	_	_	_	0.4	_	0.2	_
Unclassified	_	0.5	4.5	0.05	1.1	_	_	_
Total Retro-TEs	63.7	52.7	75.6	54.5	25.8	23.3	31.6	42.4
Type II: DNA-TEs								
CACTA	12.8	3.9	3.2	4.7	3.4	2.2	4.7	_
Helitron	0.2	0.04	2.2	0.8	0.3	0.2	0.2	_
hAT	0.04	0.02	1.1	0.02	0.5	0.2	0.6	1.2
PIF/Harbinger	0.3	0.2	_	0.02	_	0.4	_	0.01
Tc1/Mariner	1.0	0.06	_	_	0.02	0.07	_	_
Mutator	0.4	0.3	1.0	0.06	1.8	0.6	_	0.01
Other	0.01	0.04	_	_	1.3	_	_	0.03
MITE/Tourist	_	0.5 ^b	1.0	0.9	1.5	0.2	1.4	_
MITE/Stowaway	_		0.1	0.2	1.7	0.9	0.6	_
Unclassified	0.06	0.04	_	0.7	3.1	_	1.9	_
Total DNA-TEs	14.9	5.0	8.6	7.5	13.7	4.8	9.4	1.3
Unknown	1.2	0.7	_	_		_	5.4	
Total TEs	79.8	58.4	84.2	62.0	39.5	28.1	46.4	43.7

^aAll are diploids except *S. tuberosum* (tetraploid), *T. aestivum* (hexaploid), and *M. acuminata* (doubled haploid). ^bIncludes all MITEs.

thought to have promoted angiosperm diversification (Baack and Rieseberg 2007; Soltis PS and Soltis DE 2009; Jiao et al. 2011). The emergence of vigorous hybrids can result in gene and TE introgression between species. Such hybrids can sometimes become stabilized into new species, especially if polyploidy also occurs. Significantly, hybridization and polyploidy are often accompanied by extensive transposition of TEs, leading to new genomic modifications and changes in genome size (Liu and Wendel 2000; Shan et al. 2005; Josefsson et al. 2006; Ungerer et al. 2006; Kawakami et al. 2010; Parisod et al. 2010; Piednoël et al. 2013). Potentially deleterious effects on genomes that might result from such bursts of TE activity may be cushioned through gene duplication in polyploids (Matzke MA and Matzke AJ 1998). A good example of a TE burst following hybridization was documented in three diploid sunflower (Helianthus) hybrids, where massive TE derepression resulted in genomes at least 50% bigger than either diploid parent (Ungerer et al. 2006; Kawakami et al. 2010). Intriguingly, in contrast to their parent species, each of these hybrids is capable of occupying extreme (arid or saline) habitats.

The frequent and recurring production of polyploids, both autopolyploids and allopolyploids in angiosperms (Tate et al. 2005), reflects a high rate of production of unreduced gametes, especially in hybrids (Brownfield and Köhler 2011; Leitch AR and Leitch IJ 2012). Although polyploidy often represents a bottleneck due to difficulties with meiosis, nuclear enlargement, and/or epigenetic instability (Comai 2005), it has the potential to promote longer-term evolutionary success (Mayrose et al. 2011). Polyploidy may lead to speciation, as tetraploids for example, are usually reproductively isolated from their parental diploids, and polyploid populations can frequently occupy habitats not available to their parent species. In polyploids, mutations that lead to the formation of bivalents and the elimination of multivalents will be strongly selected. Therefore, genomes with active or many passive TEs (to promote TE-Thrust) may show faster homolog divergence, diploidization, and return to full gamete fertility. All angiosperms are thought to have had at least one polyploidization event in their evolutionary history usually followed by a diploidization process (Jiao et al. 2011), and individual polyploid taxa typically form multiple times (Tate et al. 2005). The widespread prevalence of this phenomenon is reflected in the recent finding that about one third of extant vascular plants are recent polyploids (Wood et al. 2009). Polyploidy has a major impact on genome size in angiosperms; however, the effect of TE amplification (and removal) is even greater (Bennetzen 2005). As new polyploid populations are small and reproductively isolated, they could result in drift to either fixation or extinction of TE families or superfamilies; an example may be the Gypsy-like Gorge LTR retro-TEs specific to the Gossypium genus (Hawkins et al. 2006).

Polyploidy is implicated in the promotion of TE proliferation in a variety of angiosperm species (Parisod et al. 2010; Piednoël et al. 2013), although its effect on TEs appears to be complex and may involve not only transposition but also TE-associated epigenetic changes and DNA recombination events (Parisod et al. 2010). Such events may lead to major genomic restructuring, producing abundant genetic novelty for adaptive evolution. A good example of a successful allopolyploid is the recently emerged and highly invasive dodecaploid species Spartina anglica involved in widespread colonizations of salt marshes and estuaries (Thompson 1991). Although no transposition burst was detected in S. anglica, major structural and epigenetic changes in the vicinity of TE insertions were observed, supporting a central role for TEs in genome reorganization during allopolyploid speciation (Parisod et al. 2009). Thus, the evolutionary impact of hybridization and/or polyploidy in angiosperms, which are important factors in their own right, would appear to be greatly magnified through the ability to enhance TE-Thrust.

Stress

Cellular TE repression mechanisms are generally sensitive to perturbation. Thus, stress can induce TE activity, which can create intragenomic potential at opportune times to facilitate adaptation in response to environmental challenge (Zeh et al. 2009; Casacuberta and González 2013). In angiosperms, TE mobilization has been reported for a variety of abiotic or biotic stress conditions including high or low temperatures, UV light, wounding, and pathogen attack (Mhiri et al. 1997; Walbot 1999; Grandbastien et al. 2005; Fujino et al. 2011; Matsunaga et al. 2012). Tolerance to one stress factor in particular, fire, has been a major factor in the success of many angiosperms (Keeley et al. 2011), including grasses and resprouting plants that are long lived and rarely reproduce from seed. Bursts of TE activity induced by the heat and damage of fire could result in genetic differences between the multiple apices that regenerate allowing somatic evolution, particularly in very long-lived resprouters and vegetatively reproducing species that rarely reproduce sexually. TEs are known to cause somatic variation in vegetatively propagated plants such as the grapevine, Vitis vinifera (Fernandez et al. 2010; Carrier et al. 2012), indicating that TE-Thrust can create intragenomic potential in the soma as well as in the germ line. This is an additional and hitherto undescribed aspect of the TE-Thrust hypothesis.

Genomic Imprinting in Endosperm

A characteristic of speciation is the emergence of pre- or postzygotic barriers to genetic exchange. Maturation of the angiosperm embryo after either intraspecific or interspecific pollination is dependent on normal development of the (usually) triploid endosperm in most taxa, which in turn is dependent on a proper balance in gene imprinting (Kinoshita 2007), consistent with a matching endosperm balance number (EBN) (Johnston et al. 1980). Thus, epigenetic mismatch/differing EBNs resulting in incorrect gene expression dosage are a frequent cause of failure of crosses and a powerful causal factor of reproductive isolation or incipient speciation. Imprinting in plants is intimately associated with changes to methylation of TEs (Gehring et al. 2009; Wolff et al. 2011), and TE activity is known to alter DNA methylation patterns and gene imprinting in plant genomes (Kashkush et al. 2003; Haun et al. 2009; Parisod et al. 2009). Thus, TEs seemingly have a significant potential to change imprinting patterns in the endosperm, resulting in reproductive isolation, and thereby indirectly promoting speciation and diversity.

Ecological Factors: Horizontal TE Transfers to Angiosperms

Angiosperms have coevolved with pollinators, fruit and seed eaters, browsers, grazers, fungi, prokaryotes, and exogenous and endogenous viruses, and likely with specialized endogenous retroviruses. Specific pollinators, mainly among insects, birds, and bats can seek out and fertilize scattered individuals of a species, allowing high species diversity in populations of angiosperms with biotic compared with wind pollination. Coevolution with metazoans for seed and fruit dispersal is also an important driver of species diversity in many angiosperms. Horizontal transfers of TEs between angiosperm genomes have been documented (Diao et al. 2006; Cheng et al. 2009; Roulin et al. 2009; Woodrow et al. 2012). This is of significance for potentially enabling TEs to prompt genomic variation within new lineages and therefore influence evolutionary trajectories (Schaack et al. 2010). An intriguing possibility worthy of future investigation is the extent to which interactions with metazoans facilitated horizontal transposon transfer to angiosperms, and also possibly, horizontal gene transfer. The same could apply to prokaryotes, fungi, and exogenous viruses.

TE-Thrust and the Evolutionary Success of Angiosperms Compared with Gymnosperms

Among plants, the angiosperms have undergone tremendous evolutionary innovations and radiations when compared with their sister clade, the gymnosperms. Apparent explanations for the lower genomic plasticity, morphological diversity, and rates of speciation in gymnosperms include lack of hybridization, polyploidy, and genetic imprinting, as well as decreased base substitution rate (Ahuja 2005; Buschiazzo et al. 2012). Significantly, despite having an abundance of TEs, TE-Thrust also appears to have been much less effective in gymnosperms. Since the angiosperm divergence, gymnosperms have experienced low TE activity with a very slow and steady accumulation of a diverse set of TEs, mainly LTR retro-TEs (Kovach et al. 2010; Nystedt et al. 2013). Thus, TEs in extant gymnosperms appear to be ancient and nonviable, whereas those in extant angiosperms are much younger and show evidence of repeated bursts of activity within the relatively recent past (Stuart-Rogers and Flavell 2001; Naito et al. 2009; Kovach et al. 2010; Lu et al. 2012; Palmer et al. 2012; El Baidouri and Panaud 2013). Moreover, in contrast to the relatively few TE subfamilies that were expanded in angiosperms (Nystedt et al. 2013), the diversity of TEs in gymnosperms makes their genomes relatively poorly suited for passive TE-Thrust. In keeping with this, gymnosperm TEs appear to be removed less frequently by unequal recombination than those in angiosperm genomes (Nystedt et al. 2013), a key outcome being the development of very large genomes that are characteristic of this lineage (Bennett and Leitch 2005). This one-way road to genomic obesity in gymnosperms may be a compounding factor in their relative lack of evolutionary diversity, as smaller angiosperm genomes offer advantages in terms of rapid seedling establishment, short generation times, and the costs and rates of reproduction (Bennett 1987). However, some angiosperm genomes are very large, as they have a much greater variety in size due to dynamism in terms of their TE amplification and TE-mediated recombination processes (Devos et al. 2002). Thus, ongoing and large amplifications, and removals, of both retro-TEs and DNA-TEs confined to the angiosperms, offer a plausible additional explanation for the lack of evolutionary innovation and speciation in the gymnosperm lineages as compared to the remarkable success of angiosperms.

Mechanisms by Which Plant Genomes Are Modified by TEs

Active TE-Thrust

TEs can powerfully facilitate genetic changes to angiosperm genomes and create intragenomic potential (standing variation), as they do in metazoans, in a large variety of ways, both active and passive. In their active role, TEs can be exapted to create new genes or functional sequences (also referred to as molecular domestication). Although not particularly common, exaptation can nevertheless have enormous impacts, such as the generation of adaptive immune system in jawed vertebrates (Schatz 2004) and of the mammalian placenta (Rawn and Cross 2008; Oliver and Greene 2012). A significant number of genes whose sequences are largely TE derived have now been reported in angiosperms (He et al. 2000; Bundock and Hooykaas 2005; Cowan et al. 2005; Muehlbauer et al. 2006; Lin et al. 2007; Roccaro et al. 2007; Duan et al. 2008; Joly-Lopez et al. 2012; Knip et al. 2012), but not in examined gymnosperms, indicating that TEs have made beneficial contributions specifically to the angiosperm gene repertoire. It is likely that further examples will be identified as more genomes are analyzed. Through exaptation, TEs can blur the distinction between themselves and their host genomes by becoming entwined with normal host cell biology. For instance, most of the matrix attachment regions (MARs) in rice and sorghum were found to colocalize with miniature inverted-repeat TEs (MITEs), suggesting that these DNA-TEs can actually serve as MARs (Avramova et al. 1998). Similarly, TEs have been found to act as source DNA for long tandem arrays at some centromeres in a variety of plant species including the potato (*Solanum tuberosum*) (Macas et al. 2009; Gong et al. 2012).

In addition to donating entire genes, TEs can contribute partially to individual genes, for instance, through the creation of introns, exons, or chimeric genes. These are not rare events, and a substantial proportion of genes in angiosperms harbor TEs, as for example rice, where more than 10% of transcripts are reported to contain TEs (Sakai et al. 2007). Significantly, this includes a contribution to about 2% of rice protein coding regions (Sakai et al. 2007). In the model plant Arabidopsis thaliana, 7.8% of expressed genes were found to contain a region with close similarity to a known TE sequence (Lockton and Gaut 2009). Brassicaceae lineage-specific genes in Arabidopsis showed an even greater percentage (about 10%) that were partly derived from TEs (Donoghue et al. 2011), which lends support for our proposal that TEs can be an important factor in lineage divergence (Oliver and Greene 2009, 2011, 2012).

Changes to gene regulation play a critical role in evolution (Carroll 2008) and a major way that TEs act to functionally modify genomes is by inserting novel regulatory elements adjacent to genes to alter or expand their expression patterns (Rebollo et al. 2012). Indeed, it was the very ability of TEs to affect gene activity in plants (specifically maize) that prompted their discoverer McClintock (1984) to refer to them as "controlling elements." A growing body of evidence now indicates that TE-derived regulatory elements can act conventionally as binding sites for transcription factors. Alternatively, they may cause epigenetic gene silencing by being targets for DNA methylation, as in the case of *Arabidopisis FLC* and *FWA* loci, and maize *B1* locus (Selinger and Chandler 2001; Fujimoto et al. 2008; Zhai et al. 2008).

Beyond their effect on the expression of individual genes, TEs can impact on gene regulation on a genomewide scale by acting as modular carriers of readymade promoters and/or enhancers via their ability to transpose throughout the genome (Britten and Davidson 1969; Feschotte 2008). This enables the widespread dissemination of discrete regulatory elements with those that confer benefit likely to be retained. A striking case is a subset of MITE DNA-TE insertions that have generated regulatory networks in rice that render adjacent gene stress inducible (Naito et al. 2009). Also striking is the presence of GC (guanine/cytosine)-rich Pack-MULE (mutatorlike element) DNA-TEs at the 5'-end of many grass genes, which may act to epigenetically control gene expression (Jiang et al. 2011). Similarly, LTR retro-TEs of the recently amplified Dasheng family have been implicated in the methylation and tissue-specific expression of adjacent rice genes (Kashkush and Khasdan 2007). Bursts of TE activity may thus be crucial for rapidly generating the large-scale genetic diversity required by angiosperms in the face of environmental and ecological challenges. They also provide a plausible mechanism by which entire sets of genes can become coregulated to fashion new cellular pathways or build on existing ones, thus potentially enhancing the extraordinary diversification of angiosperms.

The capacity of TEs to partake in the regulation of host genes is particularly supported by data from the rice genome. One sixth of rice genes are associated with retro-TEs, with insertions either in the gene itself or within putative promoter regions (Krom et al. 2008), whereas 58% are associated with a MITE (Lu et al. 2012). Thus, a large proportion of rice gene promoters appear to contain a TE. Recent evidence also indicates that many exapted plant TE sequences may actually be transcribed to function as microRNAs (miRNAs) that regulate gene expression posttranscriptionally. Now acknowledged as an important class of regulatory genes in eukaryotes, many regulatory miRNA genes found in rice are derived from TEs that have the potential to regulate thousands of genes (Li et al. 2011; Ou-Yang et al. 2013). Moreover, MITEs generate nearly a guarter of all small RNAs identified in rice (Lu et al. 2012). Thus TEs, which provide a mechanism to account for the origin of miRNAs (Buchon and Vaury 2006), appear to fulfill essential functions in plants by serving as master regulators with widespread regulatory influence.

Rather than directly contributing functional sequences to angiosperm genomes, another major way that TEs can actively generate genetic novelty is by using their transpositional (or retrotranspositional) mechanisms to delete, rearrange, or partially or fully duplicate genes or chromosomal segments. Gene duplication, in particular, is a crucial aspect of evolution and constitutes the principal means by which organisms evolve new genes (Ohno 1970). Both DNA-TEs and retro-TEs have a propensity to capture and transpose genes or gene fragments, which can result in gene duplication, exon shuffling, or regulatory element seeding, depending on the nature of the sequence involved. For example, in the rice genome, there are reportedly more than 1,200 retrogenes (Wang et al. 2006). Many of these are conserved, which implies that they have been advantageous. This includes the huge pentatricopeptide repeat gene family that likely expanded in angiosperms as a consequence of one or more waves of retrotransposition by retro-TEs (O'Toole et al. 2008). The rice genome also harbors thousands of Mutator superfamily (Pack-MULE) DNA-TEs containing fragments derived from more than 1,000 genes (Jiang et al. 2004; Juretic et al. 2005). The unparalleled ability of TEs to generate genetic novelties is reflected in the fact that many of the Pack-MULEs contain sequences from multiple chromosomal loci that are fused to form new open reading frames, some of which are expressed as chimeric transcripts. Importantly, many have undergone purifying selection (Hanada et al. 2009), indicating that they have acquired highly beneficial functions. In maize,

there is a similar situation with a very high rate of gene capture and exon shuffling by *Helitron* DNA-TEs that replicate via a rolling circle mechanism (Gupta et al. 2005; Lai et al. 2005; Morgante et al. 2005; Yang and Bennetzen 2009). The number of novel transcripts expressed by *Helitrons* is at least 11,000 or 25% of the total number of genes in the maize genome (Du et al. 2009; Barbaglia et al. 2012). Other DNA-TEs implicated in the capture and integration of gene fragments in angiosperm species include the *CACTA* and *Harbinger* DNA-TE superfamilies (Paterson et al. 2009; Vogel et al. 2010). Thus, TEs seemingly have a vast ability to concoct new coding regions and combinations of coding regions as an indispensable form of realizable intragenomic potential (standing variation) for possible future selection, whether by natural or human means.

Besides duplication, TEs are adept at moving genes, both protein and RNA coding, to new locations within a genome. Such movement has the potential to reprogram gene expression through a change in regulatory elements. An illustrative example can be found in grasses where a substantial number of miRNA genes appear to have been relocated by TEs (Abrouk et al. 2012). TEs can also induce DNA deletions through their transpositional activity, as has been observed by hAT elements in maize (Zhang and Peterson, 2005). Beyond molecular-scale changes, active TE transposition can mediate large-scale chromosomal rearrangements leading to karyotypic variation, which is a factor in the formation of reproductive barriers and speciation (Rieseberg 2001; Levin 2002). This is best documented in maize, where alternative hAT element transposition reactions can cause major changes to chromosomal architecture, including deletions, inversions, and translocations (McClintock 1950; Zhang et al. 2009). Karyotypic variability is common in angiosperms and can even occur within species. Although many TE-mediated karvotypic differences may be incidental to speciation, they represent an important potential contributory mechanism to reproductive isolation, angiosperm diversification, and species radiations.

Passive TE-Thrust

The presence of large numbers of similar TEs in genomes can separately play a passive role in plant evolution by promoting gene or segmental duplications (or deletions) through homology-driven ectopic recombination of DNA (Oliver and Greene 2009, 2011, 2012). Duplication events are particularly important because they create functional redundancy and the potential for gain of function.

TE-induced recombination events are often difficult to detect, especially those from the distant evolutionary past, which may now be untraceable. Thus, compared with active TE-Thrust, the passive effects of TEs have been less well documented. Nevertheless, passive TE-Thrust has been in evidence in *Arabidopsis* where *Copia*-like LTR retro-TEs, and *CA*

CTA and *Mutator* DNA-TEs, apparently generated segmental duplications that occurred after the monocot-dicot divergence and probably after the Rosales and Brassicales divergence (Hughes et al. 2003). On the whole, and as we outline in further detail below, the evidence points to both the active and passive effects of retro-TEs (mainly LTRs) and DNA-TEs as having greatly facilitated and influenced the trajectory of flowering plant evolution.

Evidence for Intragenomic Potential Derived from TEs in Angiosperms: Specific Examples of Traits Generated by TEs

Most data on the genomic impact of TEs are presently derived from mammals and angiosperms. In this context, realizable intragenomic potential due to TE-Thrust previously demonstrated in mammals (Oliver and Greene 2011, 2012) is also demonstrable in angiosperms, with numerous studies reporting genotypic changes due to TEs being correlated with the generation of specific flowering plant phenotypes (tables 2 and 3). Although these examples are biased toward traits of domesticated plant species, they nevertheless provide a good illustration of the power of TEs to uniquely create diverse and elaborate intragenomic potential, which can be realized by selection. Human selection in plant domestication and improvement has foresight and strategy, but is a selective force that, unless using induced mutation, must rely on the same generators of change as blind natural selection.

Tables 2 and 3 list 65 known instances in which TEs have altered or created individual plant genes and thus were directly implicated at a genomic level in the origin of various traits, both domesticated and wild. Notably, DNA-TEs were the major contributors to these traits, accounting for nearly two thirds of the total (fig. 1A). The autonomous hAT and CA CTA elements and nonautonomous MITE DNA-TEs were particularly prevalent contributors, whereas LTR retro-TEs were responsible for the remaining one third of traits. This suggests that DNA-TEs may be particularly effective at facilitating evolution, at least via active TE-Thrust (Oliver and Greene 2011), which accords with findings in disparate lineages, including the vespertilionid bats (Ray et al. 2008; Pagán et al. 2012; Mitra et al. 2013). Traits associated with cultivated plants were most commonly a consequence of gene disruption (50%; fig. 1B and table 2) rather than due to the creative effects of TEs. Although gene disruptions by TEs occur in natural populations, they generally result in a reduction of fitness and were therefore expected to be relatively uncommon. However, gene disruption features prominently in domesticated plant traits due to humans having selected for desirable null phenotypes. By contrast, traits facilitated by TEs that could be of value in wild populations were more diverse in origin and most commonly were the result of regulatory changes to plant

Specific Examples of TEs Implica	ted in Flowering	Plant Domestication and Divers						
TE-Generated or Modified Trait	Gene Affected	Gene Function	TE Responsible	Taxon	Type of Event	Effect	Type of TE-Thrust	Reference
Spring growth habit	Vrn1	Transcriptional regulator	LTR (gypsy-like)	Triticum turgidum	Regulatory	Positive regulation	Active	Chu et al. 2011
Purple coloration	BoMyb2	Transcriptional regulator	Harbinger	Brassica oleracea	Regulatory	Positive regulation	Active	Chiu et al. 2010
Floral branching	Apo1	F-box protein	һАТ	Oryza sativa	Regulatory	Positive regulation	Active	Ikeda-Kawakatsu
								et al. 2009
Fruit cluster morphology	VVTFL1A	Plant development	hAT	Vitis vinifera	Regulatory	Positive regulation	Active	Fernandez et al. 2010
Blood orange	Ruby	Transcriptional regulator	LTR (copia-like)	Citrus sinensis	Regulatory	Stress responsiveness	Active	Butelli et al. 2012
Chinese blood orange (Jingxian)	Ruby	Transcriptional regulator	LTR	Citrus sinensis	Regulatory	Stress responsiveness	Active	Butelli et al. 2012
Apical dominance	Tb1	Transcriptional regulator	LTR (copia-like)	Zea mays	Regulatory	Enhancer	Active	Studer et al. 2011
Plant pigmentation	B1	Transcriptional regulator	LTR	Zea mays	Regulatory	Epigenetic silencing	Active	Selinger and
-			H.	ľ	3			Chandler 2001
waxy kernels	, XVV	Granule-bound starch symmase	nAI	zea mays	I ransposition	Altered protein	ACTIVE	Wessier et al. 1980
Flower color pattern	niv	Anthocyanin pigmentation	hAT	Antirrhinum majus	Transposition	Altered expression	Active	Lister et al. 1993
Orange kernels and cob glume	P-00	Transcriptional regulator	hAT	Zea mays	Transposition	Novel fusion gene	Active	Zhang et al. 2006
Double flowers	DP	Transcriptional regulator	CACTA	Ipomoea nil	Transposition	Gene loss	Active	Nitasaka 2003
Elongated fruit	Sun	Auxin transport	LTR (copia-like)	Solanum lycopersicum	Retrotransposition	Duplicated gene	Active	Xiao et al. 2008
High-latitude cultivation	GmphyA2	Photoperiod sensitivity	LTR (copia-like)	Glycine max	Gene disruption	Gene inactivation	Active	Kanazawa et al. 2009
Bread-making quality	Glu-1	Glutenin seed storage protein	LTR (copia-like)	Triticum aestivum	Gene disruption	Gene inactivation	Active	Harberd et al. 1987
Parthenocarpic fruit production	MdPI	Transcriptional regulator	LTR	Malus domestica	Gene disruption	Gene inactivation	Active	Yao et al. 2001
Golden hull coloration	osCHI	Flavonoid biosynthesis	LTR	Oryza sativa	Gene disruption	Gene inactivation	Active	Hong et al. 2012
Wrinkled seed	Sbel	Starch-branching enzyme	hAT	Pisum sativum	Gene disruption	Gene inactivation	Active	Bhattacharyya
								et al. 1990
White flowers	Dfr-B	Anthocyanin pigmentation	Helitron	Ipomoea tricolor	Gene disruption	Gene inactivation	Active	Choi et al. 2007
White/variegated flowers	Chs-D	Anthocyanin pigmentation	hAT	Ipomoea purpurea	Gene disruption	Gene inactivation	Active	Habu et al. 1998
Pale flowers/lvory seed	bHlh2	Anthocyanin pigmentation	hAT	lpomoea purpurea	Gene disruption	Gene inactivation	Active	Park et al. 2007
Yellow seed	BrTT8	Transcriptional regulator	Helitron	Brassica rapa	Gene disruption	Gene inactivation	Active	Li et al. 2012
High oleate seeds	ahFAD2B	Microsomal oleoyl-phospatidyl	MITE	Arachis hypogaea	Gene disruption	Gene inactivation	Active	Patel et al. 2004
		choline desaturase						
Waxy millet	Gbss1	Granule-bound starch synthase	Multiple	Setaria italica	Gene disruption	Gene inactivation	Active	Kawase et al. 2005
Glutinous rice	WX	Granule-bound starch synthase	LTR	Oryza sativa	Gene disruption	Truncated transcript	Active	Hori et al. 2007
Variegated pigmentation	×	Phlobaphene pigmentation	CACTA	Sorghum bicolor	Gene disruption	Aberrant splicing	Active	Chopra et al. 1999
Pink flowers, lighter color,	dМ	Flavonoid biosynthesis	CACTA	Glycine max	Gene disruption	Aberrant splicing	Active	Zabala and
and higher protein								Vodkin 2005
content of seeds								
White fruit	VvMyba1	Transcriptional regulator	LTR (gypsy-like)	Vitis vinifera	Gene disruption	Low expression	Active	Walker et al. 2007
Waxy kernels	WX	Granule-bound starch synthase	LTR	Zea mays	Gene disruption	Low expression	Active	Varagona et al. 1992
Higher kernel oil content	ZmGE2	Cytochrome P450 enzyme	Mutator	Zea mays	Gene disruption	Low expression	Active	Zhang et al. 2012
Plant pigmentation	S1	Transcriptional regulator	CACTA	Zea mays	Duplication	Novel gene	Passive	Walker et al. 1995
Red fruit	VvMyba1	Transcriptional regulator	LTR (gypsy-like)	Vitis vinifera	Deletion	Regained expression	Passive	Kobayashi et al. 2004
Grain hardness	Pina, Pinb	Lipid-binding proteins	Various	Triticum aestivum	Deletion	Gene loss	Passive	Chantret et al. 2005

Specific Examples of TE	s Implicated in I	Howering Plant Physiology, Dé	evelopment, or stress	Kesistance				
TE-Generated or Modified Trait	Gene Affected	Gene Function	TE Responsible	Taxon	Type of Event	Effect	Type of TF-Thrust	Reference
Growth and flowering	Mustang 1-8	Transcriptional regulator	Mutator	Angiosperms	Domestication	Novel gene	Active	Cowan et al. 2005;
Development	Sleeper	Transcriptional regulator of	hAT	Angiosperms	Domestication	Novel gene	Active	Joly-Lopez et al. 2012 Bundock and Hoovkaas
-		plant development		-				2005; Knip et al. 2012
Light-induced responses	Fhy3	Transcriptional regulator of	Mutator	Angiosperms	Domestication	Novel gene	Active	Lin et al. 2007
		light signaling						
Light-induced responses	Far1	Transcriptional regulator of licht signaling	Mutator	Angiosperms	Domestication	Novel gene	Active	Lin et al. 2007
	Garv	Unknown	hAT	Cereal grasses	Domestication	Novel aene	Active	Muehlbauer et al. 2006
Fungal resistance	Rim2	Unknown	CACTA	Oryza sativa	Domestication	Novel gene	Active	He et al. 2000
	AtCopeg1	Hormone and nutrient stress	LTR (copia-like)	Arabidopsis thaliana	Domestication	Novel gene	Active	Duan et al. 2008
		signaling						
Flower development	TamRSI	Transcriptional regulator	CACTA	Antirrhinum majus	Domestication	Novel gene	Active	Roccaro et al. 2007
					and transposition			
Virus resistance	Z	Disease resistance	MITE	Nicotiana glutinosa	Exonization	Novel isoform	Active	Kuang et al. 2009
	OsRp16-1	Ribosomal protein	Harbinger	Oryza sativa	Exonization	Enhanced expression	Active	Kubo et al. 2008
Plant stress response	ALP-A3	Acireductone dioxygenase-like	CACTA	Triticeae (diploid)	Regulatory	Major promoter	Active	Akhunov et al. 2007
Fungal resistance	Pit	Disease resistance	LTR (copia-like)	Oryza sativa	Regulatory	Positive regulation	Active	Hayashi and Yoshida 2009
	Hsp70	Heat shock protein	MITE	Oryza sativa	Regulatory	Positive regulation	Active	Zhang et al. 2012
Growth and development	Abp1	Auxin-binding protein	MITE (Tourist)	Zea mays	Regulatory	Positive regulation	Active	Elrouby and Bureau 2000
Aluminium resistance	AltSB	Efflux transporter	MITE (Tourist)	Sorghum bicolor	Regulatory	Positive regulation	Active	Magalhaes et al. 2007
Stress response	OsGSTL2	Detoxification enzyme	hAT/MITE (Stowaway)	Oryza sativa	Regulatory	Herbicide/hormone	Active	Hu et al. 2011
						responsiveness		
Light-induced responses	TCS	Ribosome-inactivating protein	MITE	Trichosanthes kirilowii	Regulatory	Light responsiveness	Active	Xu et al. 2007
Flowering behavior	FLC	Transcriptional regulator	hAT	Arabidopsis thaliana	Regulatory	Epigenetic silencing	Active	Zhai et al. 2008
Flowering behavior	FWA	Transcriptional regulator	SINE	Arabidopsis thaliana	Regulatory	Epigenetic silencing	Active	Kinoshita et al. 2007;
								Fujimoto et al. 2008
Dessication tolerance	CDT-1	siRNA	Unknown retro-TE	Craterostigma plantagineum	Regulatory	siRNA silencing	Active	Hilbricht et al. 2008
Stress response	siRNA854	siRNA	LTR (gypsy-like)	Arabidopsis thaliana	Regulatory	siRNA silencing	Active	McCue et al. 2012
	Adh1	Alcohol dehydrogenase	hAT	Zea mays	Transposition	Enhanced expression	Active	Dawe et al. 1993
						in pollen		
Biosynthesis	Cyp72A27	Cytochrome P450	Helitron	Zea mays	Transposition	Novel gene	Active	Jameson et al. 2008
		monooxygenase	:		:			
	ZmCda3	Cytidine deaminase	Helitron	Zea mays	Transposition	Novel gene	Active	Xu and Messing 2006
Stress response	ALP-A3	Acireductone dioxygenase-like	Unknown	Triticeae (diploid)	Transposition	Novel gene	Active	Akhunov et al. 2007
Reproductive development	Bs1	Unknown	LTR (copia-like)	Zea mays	Retrotransposition	Novel gene	Active	Elrouby and Bureau 2010
Sexual reproduction	N17	Unknown	LTR	Paspalum notatum	Retrotransposition	Novel gene	Active	Ochogavía et al. 2011
Sexual reproduction	N22	Unknown	LTR (gypsy-like)	Paspalum notatum	Retrotransposition	Novel gene	Active	Ochogavía et al. 2011
	PPRs	Gene expression	Unknown	Angiosperms	Retrotransposition	Novel genes	Active	O'Toole et al. 2008
Flowering behavior	FLC	Transcriptional regulator	MITE	Arabidopsis thaliana	Gene disruption	Low expression	Active	Michaels et al. 2003
Flowering behavior	FLC	Transcriptional regulator	LTR (copia-like)	Arabidopsis thaliana	Gene disruption	Low expression	Active	Michaels et al. 2003
Seed development	z1C cluster	Seed storage proteins	Unknown	Zea mays	Duplication	Novel genes	Passive	Song et al. 2001

1894 Genome Biol. Evol. 5(10):1886–1901. doi:10.1093/gbe/evt141 Advance Access publication September 23, 2013



Fig. 1.—Summary of the effect of TEs on angiosperm adaptation and evolution. (*A*) Types of TEs implicated in the generation of traits in flowering plants. (*B*) Types of events mediated by TEs underlying flowering plant domestication and diversification. (*C*) Types of events mediated by TEs underlying wild traits in flowering plants. Based on the published data shown in tables 2 and 3.

genes (33%; fig. 1*C* and table 3). As outlined below, TE-generated traits in angiosperms could be classified into one of the four phenotypic groups, which are not necessarily mutually exclusive.

Domestication and Diversification of Crops and Ornamentals

Cultivated plants possess artificially selected characteristics that often greatly distinguish them from their wild progenitors. TEs have substantially contributed to plant domestication, in particular through gene disruption, to generate null alleles and by reprogramming gene expression (fig. 1B and table 2). The domestication of various angiosperm species provides a model to observe recent and ongoing adaptive potential due to TE-Thrust, a prominent example of which is cultivated maize. The morphology of maize, which underwent a very marked transformation from a highly branched wild progenitor (teosinte) to its modern apically dominant form, is explained in large part by the insertion of a Copia-like LTR retro-TE into a regulatory region of the teosinte branched 1 (tb1) gene to create an enhancer element (Studer et al. 2011). The resultant TE-modified (TEm) allele has increased expression of tb1, which encodes a transcriptional regulator that represses branching. The timing of the tb1 retro-TE insertion predates maize domestication by at least 10,000 years (Studer et al. 2011), indicating that human selection realized adaptive potential (standing variation) due to TE-Thrust. This closely parallels the recent realization of adaptive potential due to TE-Thrust observed in Drosophila melanogaster, where preexisting TEm alleles were adaptive for insecticide resistance and colonization of temperate climates (González et al. 2010; Schmidt et al. 2010).

Further to plant domestication per se, there is a clear link between TEs and crop improvement and/or varietal diversification. TE-generated null mutations have been particularly useful in this regard, leading to a range of agronomically useful traits (table 2), as well as Mendel's wrinkled peas (Bhattacharyya et al. 1990). Remarkably, the generation of TE-destroyed (TEd) alleles of the granule-bound starch synthase (GBSS1) gene have been repeatedly observed to underlie low amylose/sticky and waxy traits in a number of grass species including rice, maize, and millet (Varagona et al. 1992; Kawase et al. 2005; Hori et al. 2007). In the case of foxtail millet, multiple low amylose and waxy alleles of GBSS1 have been created via independent insertions of Copia- and Gypsy-like LTR and non-LTR retro-TEs, as well as autonomous (Mutator) and nonautonomous/MITE (Tourist) DNA-TEs (Kawase et al. 2005). These findings seemingly implicate TEs as a major source of new mutations, at least in some angiosperm lineages. They also suggest that the destructive power of TEs may be a significant factor in regressive evolution, a phenomenon where certain species lose features (e.g., floating aquatic plants with no roots). However, with the

occasional exception, gene disruption by TEs would be unlikely to have much value in nature as an adaptation or contribute to the evolution of a lineage.

Resistance to Stress and Disease

Plants are not mobile and must adapt to many adverse stresses such as drought, soil conditions, and temperature. TEs are known to be intimately associated with plant stress responses. both biotic and abiotic, and undergo transposition and transcription in response to stress (Grandbastien 1998). Moreover, recent findings suggest that TE-Thrust has directly made genomic contributions to the molecular and physiological responses that underlie the ability of plants to cope with stresses (table 3). Examples discovered in A. thaliana are the Copia evolved gene 1 (AtCopeg1), which is implicated in hormone and nutrient stress signaling, apparently having been domesticated from a Copia-like LTR retro-TE (Duan et al. 2008), and a Gypsy-like LTR retro-TE, which when epigenetically activated, produces a siRNA (siRNA854) that regulates expression of the UBP1b gene involved in responding to and regulating cellular stress (McCue et al. 2012). TEs have also been found to underlie stress responses in cultivated plants, for example, in sorghum, where the insertion of a MITE (Tourist) element upstream of an organic acid efflux transporter locus (AltSB) is implicated in enhanced root apex expression of the AltSB gene to confer tolerance to aluminum in soil (Magalhaes et al. 2007). Attesting to the ability of TEs to cause genetic change above and beyond traditional mutagens (Oliver and Greene 2012) is the evolution of the ALP-A3 gene (encoding an acireductone dioxygenase-like protein) in some Triticaceae species, including diploid wheat. Remarkably, TEs facilitated both the creation of this gene through DNA transposition and its subsequent expression by virtue of a promoter sequence derived from a CACTA DNA-TE (Akhunov et al. 2007). TEs have also enhanced the ability of plants to defend against disease. The Rim2 gene implicated in defense against fungal infection appears to have been directly exapted from part of a CACTA DNA-TE element (He et al. 2000), whereas an inactive rice blast disease resistance gene, Pit, was refunctionalized by the recruitment of a Copia-like LTR element as a promoter (Hayashi and Yoshida 2009).

Growth and Development

Growth, reproduction, and development are key fitness determining factors that have been influenced by TEs (table 3). Two particularly striking examples of fitness benefits brought about by TEs in flowering plants are the *Mustang* and *Sleeper* gene families, whose sequences derive from exapted transposases from *Mutator*-like DNA-TEs and *hAT* DNA-TEs, respectively (Bundock and Hooykaas 2005; Cowan et al. 2005; Joly-Lopez et al. 2012; Knip et al. 2012). *Mustang* genes are present only in the angiosperm lineage and encode putative transcriptional regulators that play important roles in growth, flower development, and reproduction. They are important for fitness because plants harboring mutated *Mustang* genes show major defects in floral organ development, fecundity, and reproductive timing (Joly-Lopez et al. 2012). Similar findings have been reported for *Sleeper* genes (Bundock and Hooykaas 2005; Knip et al. 2012). Because *Mustang* and *Sleeper* genes are found in all examined angiosperms, they appear to have been important factors in the phyletic differentiation of the angiosperms and seemingly represent key instances of realized evolutionary potential due to TE-Thrust.

Physiological and Metabolic Adaptations

TEs underlie a variety of adaptations associated with plant physiology and metabolism (table 3), including responses to light, which plants not only harness as a source of energy but also monitor constantly in order to grow and respond to seasonal changes. Most processes regulated by light involve alterations in gene expression. TEs can impart light responsiveness on genes via insertion into gene regulatory regions, as in the Chinese cucumber (Trichosanthes kirilowii) TCS gene, which has a MITE DNA-TE in its promoter (Xu et al. 2007). Two genes identified in Arabidopsis associated with light-induced responses, *Fhy3* and *Far1*, represent further prime examples of exaptation. These genes were co-opted from an ancient transposase belonging to a *Mutator*-like DNA-TE (Hudson et al. 2003; Lin et al. 2007) and encode transcriptional regulators that jointly act downstream of the photoreceptor phytochrome A to specifically modulate far-red light-responsive gene expression. This is crucially required for various processes such as chlorophyll biosynthesis, circadian rhythm, shade tolerance, seed germination, and flowering (Nagy et al. 2000; Allen et al. 2006; Tang et al. 2012). Such key light-sensing mechanisms have been suggested to be a critical development in angiosperm evolution, conferring upon this lineage an adaptive advantage as well as promoting their extraordinary diversification (Mathews 2006).

Conclusion

By assessing the available evidence, we conclude that TE-Thrust operates in, and has been crucial to, the evolution of flowering plants. The additional involvement of TEs in the artificial arena of plant domestication provides direct and relatively recent evidence for the importance of TEs in the generation of selectable variation in angiosperms. TE-Thrust is therefore potentially a general phenomenon that may have very widespread significance to many lineages of life on earth. Nevertheless, TE-Thrust is only one of the many facilitators of evolution, and its relative importance may vary from lineage to lineage and from age to age. A comprehension of the full magnitude of the contributions that TEs have made to angiosperm evolution will require complete genome sequencing and detailed trait characterization in a wide range of plant species, including nondomesticated species of angiosperms and species from other plant phyla. However, any measure of TE impact will likely be an underestimate owing to important contributions having been made by ancient TEs that have been lost or are no longer recognizable.

Accepted explanations for angiosperm diversity are valid and persuasive, but still cannot fully account for the extreme diversity of angiosperms. We add to this explanation by proposing that there is good evidence that the TE-Thrust hypothesis, in addition to the accepted explanations, gives a fuller and more complete explanation for the extraordinary angiosperm diversification. The same realizable intragenomic potential due to TE-Thrust shown in metazoans, particularly mammals, is affirmed here in angiosperms. Thus, the remarkable advancement and radiation of the angiosperms appears to have been significantly aided by TE-Thrust powered by the prominent presence of LTR elements in partnership with active DNA-TE families. However, due to a paucity of data regarding the deeper evolutionary history of angiosperms and the short timescale of human selection, adaptive potential, rather than evolutionary potential, is more readily apparent at present. Nonetheless, exceptional examples of evolutionary potential appear to include the TE-derived Mustang and Sleeper genes, which may have underpinned the development of floral organs, a key morphological divergence of the angiosperms. All things considered, current evidence points to TEs being a highly significant facilitator of evolution in the angiosperms, as we have previously proposed them to be in other lineages (Oliver and Greene 2009, 2011, 2012), and this significantly broadens the applicability of, and base of support for, the TE-Thrust hypothesis.

Literature Cited

- Abrouk M, et al. 2012. Grass microRNA gene paleohistory unveils new insights into gene dosage balance in subgenome partitioning after whole-genome duplication. Plant Cell 24:1776–1792.
- Ahuja MR. 2005. Polyploidy in gymnosperms: revisited. Silvae Genet. 54: 59–69.
- Akhunov ED, Akhunova AR, Dvorak J. 2007. Mechanisms and rates of birth and death of dispersed duplicated genes during the evolution of a multigene family in diploid and tetraploid wheats. Mol Biol Evol. 24: 539–550.
- Allen T, et al. 2006. *Arabidopsis* FHY3 specifically gates phytochrome signaling to the circadian clock. Plant Cell 18:2506–2516.
- Avramova Z, Tikhonov A, Chen M, Bennetzen JL. 1998. Matrix attachment regions and structural colinearity in the genomes of two grass species. Nucleic Acids Res. 26:761–767.
- Baack EJ, Rieseberg LH. 2007. A genomic view of introgression and hybrid speciation. Curr Opin Genet Dev. 17:513–518.
- Barbaglia AM, et al. 2012. Gene capture by *Helitron* transposons reshuffles the transcriptome of maize. Genetics 190:965–975.
- Bennett MD. 1987. Variation in genomic form in plants and its ecological implications. New Phytol. 106:177–200.
- Bennett MD, Leitch IJ. 2005. Genome size evolution in plants. In: Gregory TR, editor. The evolution of the genome. San Diego (CA): Elsevier. p. 89–162.
- Bennetzen JL. 2000. Transposable element contributions to plant gene and genome evolution. Plant Mol Biol. 42:251–269.

- Bennetzen JL. 2005. Transposable elements, gene creation and genome rearrangement in flowering plants. Curr Opin Genet Dev. 15: 621–627.
- Bhattacharyya MK, Smith AM, Ellis TH, Hedley C, Martin C. 1990. The wrinkled-seed character of pea described by Mendel is caused by a transposon-like insertion in a gene encoding starch-branching enzyme. Cell 60:115–122.
- Biémont C, Vieira C. 2006. Genetics: junk DNA as an evolutionary force. Nature 443:521–524.
- Böhne A, Brunet F, Galiana-Arnoux D, Schultheis C, Volff JN. 2008. Transposable elements as drivers of genomic and biological diversity in vertebrates. Chromosome Res. 16:203–215.
- Britten RJ, Davidson EH. 1969. Gene regulation for higher cells: a theory. Science 165:349–357.
- Brownfield L, Köhler C. 2011. Unreduced gamete formation in plants: mechanisms and prospects. J Exp Bot. 62:1659–1668.
- Buchon N, Vaury C. 2006. RNAi: a defensive RNA-silencing against viruses and transposable elements. Heredity 96:195–202.
- Bundock P, Hooykaas P. 2005. An *Arabidopsis hAT*-like transposase is essential for plant development. Nature 436:282–284.
- Buschiazzo E, Ritland C, Bohlmann J, Ritland K. 2012. Slow but not low: genomic comparisons reveal slower evolutionary rate and higher dN/dS in conifers compared to angiosperms. BMC Evol Biol. 12:8.
- Butelli E, et al. 2012. Retrotransposons control fruit-specific, cold-dependent accumulation of anthocyanins in blood oranges. Plant Cell 24: 1242–1255.
- Carrier G, et al. 2012. Transposable elements are a major cause of somatic polymorphism in *Vitis vinifera L*. PLoS One 7:e32973.
- Carroll SB. 2008. Evo-devo and an expanding evolutionary synthesis: a genetic theory of morphological evolution. Cell 134:25–36.
- Casacuberta E, González J. 2013. The impact of transposable elements in environmental adaptation. Mol Ecol. 22:1503–1517.
- Chantret N, et al. 2005. Molecular basis of evolutionary events that shaped the *hardness* locus in diploid and polyploid wheat species (*Triticum* and *Aegilops*). Plant Cell 17:1033–1045.
- Cheng X, Zhang D, Cheng Z, Keller B, Ling HQ. 2009. A new family of Ty1copia-like retrotransposons originated in the tomato genome by a recent horizontal transfer event. Genetics 181:1183–1193.
- Chiu LW, et al. 2010. The purple cauliflower arises from activation of a MYB transcription factor. Plant Physiol. 154:1470–1480.
- Choi JD, Hoshino A, Park KI, Park IS, lida S. 2007. Spontaneous mutations caused by a *Helitron* transposon, *Hel-It1*, in morning glory, *Ipomoea tricolor*. Plant J. 49:924–934.
- Chopra S, Brendel V, Zhang J, Axtell JD, Peterson T. 1999. Molecular characterization of a mutable pigmentation phenotype and isolation of the first active transposable element from *Sorghum bicolor*. Proc Natl Acad Sci U S A. 96:15330–15335.
- Chu CG, et al. 2011. A novel retrotransposon inserted in the dominant *Vm-B1* allele confers spring growth habit in tetraploid wheat (*Triticum turgidum* L.). G3 1:637–645.
- Comai L. 2005. The advantages and disadvantages of being polyploid. Nat Rev Genet. 6:836–846.
- Cowan RK, Hoen DR, Schoen DJ, Bureau TE. 2005. *MUSTANG* is a novel family of domesticated transposase genes found in diverse angio-sperms. Mol Biol Evol. 22:2084–2089.
- Dawe RK, Lachmansingh AR, Freeling M. 1993. Transposon-mediated mutations in the untranslated leader of maize *Adh1* that increase and decrease pollen-specific gene expression. Plant Cell 5:311–319.
- Devos KM, Brown JK, Bennetzen JL. 2002. Genome size reduction through illegitimate recombination counteracts genome expansion in *Arabidopsis*. Genome Res. 12:1075–1079.
- Diao X, Freeling M, Lisch D. 2006. Horizontal transfer of a plant transposon. PLoS Biol. 4:e5.

- Donoghue MT, Keshavaiah C, Swamidatta SH, Spillane C. 2011. Evolutionary origins of Brassicaceae specific genes in *Arabidopsis thaliana*. BMC Evol Biol. 11:47.
- Doolittle WF, Sapienza C. 1980. Selfish genes, the phenotype paradigm and genome evolution. Nature 284:601–603.
- Du C, Fefelova N, Caronna J, He L, Dooner HK. 2009. The polychromatic *Helitron* landscape of the maize genome. Proc Natl Acad Sci U S A. 106:19916–19921.
- Duan K, et al. 2008. *AtCopeg1*, the unique gene originated from AtCopia95 retrotransposon family, is sensitive to external hormones and abiotic stresses. Plant Cell Rep. 27:1065–1073.
- El Baidouri M, Panaud O. 2013. Comparative genomic paleontology across plant kingdom reveals the dynamics of TE-driven genome evolution. Genome Biol Evol. 5:954–965.
- Eldredge N, Gould SJ. 1972. Punctuated equilibria: an alternative to phyletic gradualism. In: Schopf TJM, editor. Models in paleobiology. San Francisco (CA): Freeman Cooper. p. 82–115.
- Elrouby N, Bureau TE. 2000. Molecular characterization of the *Abp1* 5'flanking region in maize and the teosintes. Plant Physiol. 124: 369–377.
- Elrouby N, Bureau TE. 2010. *Bs1*, a new chimeric gene formed by retrotransposon-mediated exon shuffling in maize. Plant Physiol. 153: 1413–1424.
- Fernandez L, Torregrosa L, Segura V, Bouquet A, Martinez-Zapater JM. 2010. Transposon-induced gene activation as a mechanism generating cluster shape somatic variation in grapevine. Plant J. 61: 545–557.
- Feschotte C. 2008. Transposable elements and the evolution of regulatory networks. Nat Rev Genet. 9:397–405.
- Feschotte C, Gilbert C. 2012. Endogenous viruses: insights into viral evolution and impact on host biology. Nat Rev Genet. 13:283–296.
- Feschotte C, Pritham EJ. 2007. DNA transposons and the evolution of eukaryotic genomes. Annu Rev Genet. 41:331–368.
- Fujimoto R, et al. 2008. Evolution and control of imprinted FWA genes in the genus Arabidopsis. PLoS Genet. 4:e1000048.
- Fujino K, et al. 2011. Temperature controls nuclear import of Tam3 transposase in *Antirrhinum*. Plant J. 65:146–155.
- Gaut BS, Wright SI, Rizzon C, Dvorak J, Anderson LK. 2007. Recombination: an underappreciated factor in the evolution of plant genomes. Nat Rev Genet. 8:77–84.
- Gehring M, Bubb KL, Henikoff S. 2009. Extensive demethylation of repetitive elements during seed development underlies gene imprinting. Science 324:1447–1451.
- Gerasimova TI, Matjunina LV, Mizrokhi LJ, Georgiev GP. 1985. Successive transposition explosions in *Drosophila melanogaster* and reverse transpositions of mobile dispersed genetic elements. EMBO J. 4: 3773–3779.
- Gibbs RA, et al. 2004. Genome sequence of the Brown Norway rat yields insights into mammalian evolution. Nature 428:493–521.
- Gong Z, et al. 2012. Repeatless and repeat-based centromeres in potato: implications for centromere evolution. Plant Cell 24:3559–3574.
- González J, Karasov TL, Messer PW, Petrov DA. 2010. Genome-wide patterns of adaptation to temperate environments associated with transposable elements in *Drosophila*. PLoS Genet. 6:e1000905.
- Gould SJ. 2002. The structure of evolutionary theory. Cambridge: The Belknap Press of Harvard University Press.
- Grandbastien MA. 1998. Activation of plant retrotransposons under stress conditions. Trends Plant Sci. 3:181–187.
- Grandbastien MA, et al. 2005. Stress activation and genomic impact of *Tnt1* retrotransposons in Solanaceae. Cytogenet Genome Res. 110: 229–241.
- Gupta S, Gallavotti A, Stryker GA, Schmidt RJ, Lal SK. 2005. A novel class of *Helitron*-related transposable elements in maize contain portions of multiple pseudogenes. Plant Mol Biol. 57:115–127.

- Habu Y, Hisatomi Y, Iida S. 1998. Molecular characterization of the mutable *flaked* allele for flower variegation in the common morning glory. Plant J. 16:371–376.
- Hanada K, et al. 2009. The functional role of Pack-MULEs in rice inferred from purifying selection and expression profile. Plant Cell 21:25–38.
- Harberd NP, Flavell RB, Thompson RD. 1987. Identification of a transposon-like insertion in a *Glu-1* allele of wheat. Mol Gen Genet. 209: 326–332.
- Haun WJ, Danilevskaya ON, Meeley RB, Springer NM. 2009. Disruption of imprinting by *Mutator* transposon insertions in the 5' proximal regions of the *Zea mays Mez1* locus. Genetics 181:1229–1237.
- Hawkins JS, Kim H, Nason JD, Wing RA, Wendel JF. 2006. Differential lineage-specific amplification of transposable elements is responsible for genome size variation in *Gossypium*. Genome Res. 16: 1252–1261.
- Hayashi K, Yoshida H. 2009. Refunctionalization of the ancient rice blast disease resistance gene *Pit* by the recruitment of a retrotransposon as a promoter. Plant J. 57:413–425.
- He ZH, Dong HT, Dong JX, Li DB, Ronald PC. 2000. The rice *Rim2* transcript accumulates in response to *Magnaporthe grisea* and its predicted protein product shares similarity with TNP2-like proteins encoded by *CACT A* transposons. Mol Gen Genet. 264:2–10.
- Hilbricht T, et al. 2008. Retrotransposons and siRNA have a role in the evolution of desiccation tolerance leading to resurrection of the plant *Craterostigma plantagineum*. New Phytol. 179:877–887.
- Hong L, et al. 2012. A mutation in the rice chalcone isomerase gene causes the *golden hull and internode 1* phenotype. Planta 236:141–151.
- Hori Y, Fujimoto R, Sato Y, Nishio T. 2007. A novel *wx* mutation caused by insertion of a retrotransposon-like sequence in a glutinous cultivar of rice (*Oryza sativa*). Theor Appl Genet. 115:217–224.
- Hu T, et al. 2011. Isolation and characterization of a rice glutathione *S*-transferase gene promoter regulated by herbicides and hormones. Plant Cell Rep. 30:539–549.
- Hua-Van A, Le Rouzic A, Boutin TS, Filée J, Capy P. 2011. The struggle for life of the genome's selfish architects. Biol Direct. 6:19.
- Hudson ME, Lisch DR, Quail PH. 2003. The *FHY3* and *FAR1* genes encode transposase-related proteins involved in regulation of gene expression by the phytochrome A-signaling pathway. Plant J. 34:453–471.
- Hughes AL, Friedman R, Ekollu V, Rose JR. 2003. Non-random association of transposable elements with duplicated genomic blocks in *Arabidopsis thaliana*. Mol Phylogenet Evol. 29:410–416.
- Ibarra-Laclette E, et al. 2013. Architecture and evolution of a minute plant genome. Nature 498:94–98.
- Ikeda-Kawakatsu K, et al. 2009. Expression level of ABERRANT PANICLE ORGANIZATION1 determines rice inflorescence form through control of cell proliferation in the meristem. Plant Physiol. 150:736–747.
- Jameson N, et al. 2008. *Helitron* mediated amplification of cytochrome P450 monooxygenase gene in maize. Plant Mol Biol. 67:295–304.
- Jiang N, Bao Z, Zhang X, Eddy SR, Wessler SR. 2004. Pack-MULE transposable elements mediate gene evolution in plants. Nature 431: 569–573.
- Jiang N, Ferguson AA, Slotkin RK, Lisch D. 2011. Pack-*Mutator*-like transposable elements (Pack-MULEs) induce directional modification of genes through biased insertion and DNA acquisition. Proc Natl Acad Sci U S A. 108:1537–1542.
- Jiao Y, et al. 2011. Ancestral polyploidy in seed plants and angiosperms. Nature 473:97–100.
- Johnston SA, den Nijs TPM, Peloquin SJ, Hanneman RE Jr. 1980. The significance of genic balance to endosperm development in interspecific crosses. Theor Appl Genet. 57:5–9.
- Joly-Lopez Z, Forczek E, Hoen DR, Juretic N, Bureau TE. 2012. A gene family derived from transposable elements during early angiosperm evolution has reproductive fitness benefits in *Arabidopsis thaliana*. PLoS Genet. 8:e1002931.

- Josefsson C, Dilkes B, Comai L. 2006. Parent-dependent loss of gene silencing during interspecies hybridization. Curr Biol. 16: 1322–1328.
- Juretic N, Hoen DR, Huynh ML, Harrison PM, Bureau TE. 2005. The evolutionary fate of MULE-mediated duplications of host gene fragments in rice. Genome Res. 15:1292–1297.
- Kanazawa A, Liu B, Kong F, Arase S, Abe J. 2009. Adaptive evolution involving gene duplication and insertion of a novel *Ty1/copia*-like retrotransposon in soybean. J Mol Evol. 69:164–175.
- Kashkush K, Feldman M, Levy AA. 2003. Transcriptional activation of retrotransposons alters the expression of adjacent genes in wheat. Nat Genet. 33:102–106.
- Kashkush K, Khasdan V. 2007. Large-scale survey of cytosine methylation of retrotransposons and the impact of readout transcription from long terminal repeats on expression of adjacent rice genes. Genetics 177: 1975–1985.
- Kawakami T, Strakosh SC, Zhen Y, Ungerer MC. 2010. Different scales of *Ty1/copia*-like retrotransposon proliferation in the genomes of three diploid hybrid sunflower species. Heredity 104:341–350.
- Kawase M, Fukunaga K, Kato K. 2005. Diverse origins of waxy foxtail millet crops in East and Southeast Asia mediated by multiple transposable element insertions. Mol Genet Genomics. 274:131–140.
- Kazazian HH Jr. 2004. Mobile elements: drivers of genome evolution. Science 303:1626–1632.
- Keeley JE, Pausas JG, Rundel PW, Bond WJ, Bradstock RA. 2011. Fire as an evolutionary pressure shaping plant traits. Trends Plant Sci. 16: 406–411.
- Keeling PJ, Palmer JD. 2008. Horizontal gene transfer in eukaryotic evolution. Nat Rev Genet. 9:605–618.
- Kejnovsky E, Leitch IJ, Leitch AR. 2009. Contrasting evolutionary dynamics between angiosperm and mammalian genomes. Trends Ecol Evol. 24: 572–582.
- Kinoshita T. 2007. Reproductive barrier and genomic imprinting in the endosperm of flowering plants. Genes Genet Syst. 82:177–186.
- Kinoshita Y, et al. 2007. Control of FWA gene silencing in Arabidopsis thaliana by SINE-related direct repeats. Plant J. 49:38–45.
- Knip M, de Pater S, Hooykaas PJ. 2012. The SLEEPER genes: a transposasederived angiosperm-specific gene family. BMC Plant Biol. 12:192.
- Kobayashi S, Goto-Yamamoto N, Hirochika H. 2004. Retrotransposon-induced mutations in grape skin color. Science 304:982.
- Kovach A, et al. 2010. The *Pinus taeda* genome is characterized by diverse and highly diverged repetitive sequences. BMC Genomics 11:420.
- Krom N, Recla J, Ramakrishna W. 2008. Analysis of genes associated with retrotransposons in the rice genome. Genetica 134:297–310.
- Kuang H, et al. 2009. Identification of miniature inverted-repeat transposable elements (MITEs) and biogenesis of their siRNAs in the Solanaceae: new functional implications for MITEs. Genome Res. 19: 42–56.
- Kubo N, Fujimoto M, Arimura S, Hirai M, Tsutsumi N. 2008. Transfer of rice mitochondrial ribosomal protein L6 gene to the nucleus: acquisition of the 5'-untranslated region via a transposable element. BMC Evol Biol. 8:314.
- Lai J, Li Y, Messing J, Dooner HK. 2005. Gene movement by *Helitron* transposons contributes to the haplotype variability of maize. Proc Natl Acad Sci U S A. 102:9068–9073.
- Leitch AR, Leitch JJ. 2012. Ecological and genetic factors linked to contrasting genome dynamics in seed plants. New Phytol. 194:629–646.
- Levin DA. 2002. The role of chromosomal change in plant evolution. Oxford: Oxford University Press.
- Li X, et al. 2012. A large insertion in bHLH transcription factor *BrTT8* resulting in yellow seed coat in *Brassica rapa*. PLoS One 7:e44145.
- Li Y, Li C, Xia J, Jin Y. 2011. Domestication of transposable elements into MicroRNA genes in plants. PLoS One 6:e19212.

- Lin R, et al. 2007. Transposase-derived transcription factors regulate light signaling in *Arabidopsis*. Science 318:1302–1305.
- Lister C, Jackson D, Martin C. 1993. Transposon-induced inversion in *Antirrhinum* modifies *nivea* gene expression to give a novel flower color pattern under the control of *Cycloidea radialis*. Plant Cell 5: 1541–1553.
- Liu B, Wendel JF. 2000. Retrotransposon activation followed by rapid repression in introgressed rice plants. Genome 43:874–880.
- Lockton S, Gaut BS. 2009. The contribution of transposable elements to expressed coding sequence in *Arabidopsis thaliana*. J Mol Evol. 68: 80–89.
- Lu C, et al. 2012. Miniature inverted-repeat transposable elements (MITEs) have been accumulated through amplification bursts and play important roles in gene expression and species diversity in *Oryza sativa*. Mol Biol Evol. 29:1005–1017.
- Macas J, Koblízková A, Navrátilová A, Neumann P. 2009. Hypervariable 3' UTR region of plant LTR-retrotransposons as a source of novel satellite repeats. Gene 448:198–206.
- Magalhaes JV, et al. 2007. A gene in the multidrug and toxic compound extrusion (MATE) family confers aluminum tolerance in sorghum. Nat Genet. 39:1156–1161.
- Maksakova IA, et al. 2006. Retroviral elements and their hosts: insertional mutagenesis in the mouse germ line. PLoS Genet. 2:e2.
- Marques AC, Dupanloup I, Vinckenbosch N, Reymond A, Kaessmann H. 2005. Emergence of young human genes after a burst of retroposition in primates. PLoS Biol. 3:e357.
- Mathews S. 2006. Phytochrome-mediated development in land plants: red light sensing evolves to meet the challenges of changing light environments. Mol Ecol. 15:3483–3503.
- Matsunaga W, Kobayashi A, Kato A, Ito H. 2012. The effects of heat induction and the siRNA biogenesis pathway on the transgenerational transposition of *ONSEN*, a *copia*-like retrotransposon in *Arabidopsis thaliana*. Plant Cell Physiol. 53:824–833.
- Matzke MA, Matzke AJ. 1998. Gene silencing in plants: relevance for genome evolution and the acquisition of genomic methylation patterns. Novartis Found Symp. 214:168–180.
- Mayrose I, et al. 2011. Recently formed polyploid plants diversify at lower rates. Science 333:1257.
- McClintock B. 1950. The origin and behavior of mutable loci in maize. Proc Natl Acad Sci U S A. 36:344–355.
- McClintock B. 1984. The significance of responses of the genome to challenge. Science 226:792–801.
- McCue AD, Nuthikattu S, Reeder SH, Slotkin RK. 2012. Gene expression and stress response mediated by the epigenetic regulation of a transposable element small RNA. PLoS Genet. 8: e1002474.
- Mhiri C, et al. 1997. The promoter of the tobacco Tnt1 retrotransposon is induced by wounding and by abiotic stress. Plant Mol Biol. 33: 257–266.
- Michaels SD, He Y, Scortecci KC, Amasino RM. 2003. Attenuation of *FLOWERING LOCUS C* activity as a mechanism for the evolution of summer-annual flowering behavior in *Arabidopsis*. Proc Natl Acad Sci U S A. 100:10102–10107.
- Miguel C, Simões M, Oliveira MM, Rocheta M. 2008. Envelope-like retrotransposons in the plant kingdom: evidence of their presence in gymnosperms (*Pinus pinaster*). J Mol Evol. 67:517–525.
- Mitra R, et al. 2013. Functional characterization of *piggyBat* from the bat *Myotis lucifugus* unveils an active mammalian DNA transposon. Proc Natl Acad Sci U S A. 110:234–239.
- Morgante M, et al. 2005. Gene duplication and exon shuffling by helitronlike transposons generate intraspecies diversity in maize. Nat Genet. 37:997–1002.
- Muehlbauer GJ, et al. 2006. A *hAT* superfamily transposase recruited by the cereal grass genome. Mol Genet Genomics. 275:553–563.

- Müller K, et al. 2004. Evolution of carnivory in Lentibulariaceae and the Lamiales. Plant Biol. 6:477–490.
- Naito K, et al. 2009. Unexpected consequences of a sudden and massive transposon amplification on rice gene expression. Nature 461: 1130–1134.
- Nagy F, Kircher S, Schäfer E. 2000. Nucleo-cytoplasmic partitioning of the plant photoreceptors phytochromes. Semin Cell Dev Biol. 11: 505–510.
- Nitasaka E. 2003. Insertion of an *En/Spm*-related transposable element into a floral homeotic gene *DUPLICATED* causes a double flower phenotype in the Japanese morning glory. Plant J. 36: 522–531.
- Nystedt B, et al. 2013. The Norway spruce genome sequence and conifer genome evolution. Nature 497:579–584.
- Ochogavía AC, et al. 2011. Characterization of retrotransposon sequences expressed in inflorescences of apomictic and sexual *Paspalum notatum* plants. Sex Plant Reprod. 24:231–246.
- Ohno S. 1970. Evolution by gene duplication. New York: Springer-Verlag.
- Oliver KR, Greene WK. 2009. Transposable elements: powerful facilitators of evolution. BioEssavs 31:703–714.
- Oliver KR, Greene WK. 2011. Mobile DNA and the TE-thrust hypothesis: supporting evidence from the primates. Mobile DNA. 2:8.
- Oliver KR, Greene WK. 2012. Transposable elements and viruses as factors in adaptation and evolution: an expansion and strengthening of the TE-Thrust hypothesis. Ecol Evol. 2:2912–2933.
- Orgel LE, Crick FH. 1980. Selfish DNA: the ultimate parasite. Nature 284: 604–607.
- O'Toole N, et al. 2008. On the expansion of the pentatricopeptide repeat gene family in plants. Mol Biol Evol. 25:1120–1128.
- Ou-Yang F, et al. 2013. Transposable element-associated microRNA hairpins produce 21-nt sRNAs integrated into typical microRNA pathways in rice. Funct Integr Genomics. 13:207–216.
- Pagán HJ, et al. 2012. Survey sequencing reveals elevated DNA transposon activity, novel elements, and variation in repetitive landscapes among vesper bats. Genome Biol Evol. 4:575–585.
- Palmer SA, et al. 2012. Archaeogenomic evidence of punctuated genome evolution in *Gossypium*. Mol Biol Evol. 29:2031–2038.
- Parisod C, et al. 2009. Rapid structural and epigenetic reorganization near transposable elements in hybrid and allopolyploid genomes in *Spartina*. New Phytol. 184:1003–1015.
- Parisod C, et al. 2010. Impact of transposable elements on the organization and function of allopolyploid genomes. New Phytol. 186:37–45.
- Park KI, et al. 2007. A *bHLH* regulatory gene in the common morning glory, *Ipomoea purpurea*, controls anthocyanin biosynthesis in flowers, proanthocyanidin and phytomelanin pigmentation in seeds, and seed trichome formation. Plant J. 49:641–654.
- Patel M, et al. 2004. High-oleate peanut mutants result from a MITE insertion into the *FAD2* gene. Theor Appl Genet. 108:1492–1502.
- Paterson AH, et al. 2009. The *Sorghum bicolor* genome and the diversification of grasses. Nature 457:551–556.
- Piednoël M, Carrete-Vega G, Renner SS. 2013. Characterization of the LTR retrotransposon repertoire of a plant clade of six diploid and one tetraploid species. Plant J. 75:699–709.
- Pigliucci M. 2007. Do we need an extended evolutionary synthesis? Evolution 61:2743–2749.
- Rawn SM, Cross JC. 2008. The evolution, regulation, and function of placenta-specific genes. Annu Rev Cell Dev Biol. 24:159–181.
- Ray DA, et al. 2008. Multiple waves of recent DNA transposon activity in the bat, *Myotis lucifugus*. Genome Res. 18:717–728.
- Rebollo R, Romanish MT, Mager DL. 2012. Transposable elements: an abundant and natural source of regulatory sequences for host genes. Annu Rev Genet. 46:21–42.
- Rieseberg LH. 2001. Chromosomal rearrangements and speciation. Trends Ecol Evol. 16:351–358.

- Roccaro M, Li Y, Sommer H, Saedler H. 2007. *ROSINA (RSI)* is part of a *CAC TA* transposable element, *TamRSI*, and links flower development to transposon activity. Mol Genet Genomics. 278:243–254.
- Roulin A, et al. 2009. Whole genome surveys of rice, maize and sorghum reveal multiple horizontal transfers of the LTR-retrotransposon *Route66* in Poaceae. BMC Evol Biol. 9:58.
- Sakai H, Tanaka T, Itoh T. 2007. Birth and death of genes promoted by transposable elements in *Oryza sativa*. Gene 392:59–63.
- SanMiguel P, Bennetzen JL. 1998. Evidence that a recent increase in maize genome size was caused by the massive amplification of intergene retrotransposons. Ann Bot. 82:37–44.
- Schaack S, Gilbert C, Feschotte C. 2010. Promiscuous DNA: horizontal transfer of transposable elements and why it matters for eukaryotic evolution. Trends Ecol Evol. 25:537–546.
- Schatz DG. 2004. Antigen receptor genes and the evolution of a recombinase. Semin Immunol. 16:245–256.
- Schmidt JM, et al. 2010. Copy number variation and transposable elements feature in recent, ongoing adaptation at the *Cyp6g1* locus. PLoS Genet. 6:e1000998.
- Selinger DA, Chandler VL. 2001. *B-Bolivia*, an allele of the maize *b1* gene with variable expression, contains a high copy retrotransposon-related sequence immediately upstream. Plant Physiol. 125:1363–1379.
- Shan X, et al. 2005. Mobilization of the active MITE transposons *mPing* and *Pong* in rice by introgression from wild rice (*Zizania latifolia* Griseb.). Mol Biol Evol. 22:976–990.
- Soltis DE, Bell CD, Kim S, Soltis PS. 2008. Origin and early evolution of angiosperms. Ann NY Acad Sci. 1133:3–25.
- Soltis PS, Soltis DE. 2009. The role of hybridization in plant speciation. Annu Rev Plant Biol. 60:561–588.
- Song R, Llaca V, Linton E, Messing J. 2001. Sequence, regulation, and evolution of the maize 22-kD alpha zein gene family. Genome Res. 11: 1817–1825.
- Stuart-Rogers C, Flavell AJ. 2001. The evolution of Ty1-copia group retrotransposons in gymnosperms. Mol Biol Evol. 18:155–163.
- Studer A, Zhao Q, Ross-Ibarra J, Doebley J. 2011. Identification of a functional transposon insertion in the maize domestication gene *tb1*. Nat Genet. 43:1160–1163.
- Tang W, et al. 2012. Transposase-derived proteins FHY3/FAR1 interact with PHYTOCHROME-INTERACTING FACTOR1 to regulate chlorophyll biosynthesis by modulating *HEMB1* during deetiolation in *Arabidopsis*. Plant Cell 24:1984–2000.
- Tate JA, Soltis DE, Soltis PS. 2005. Polyploidy in plants. In: Gregory TR, editor. The evolution of the genome. San Diego (CA): Elsevier. p. 371–426.
- Thompson JD. 1991. The biology of an invasive plant. Bioscience 41: 393–401.
- Ungerer MC, Strakosh SC, Zhen Y. 2006. Genome expansion in three hybrid sunflower species is associated with retrotransposon proliferation. Curr Biol. 16:R872–R873.
- Van de Peer Y, Maere S, Meyer A. 2009. The evolutionary significance of ancient genome duplications. Nat Rev Genet. 10:725–732.
- Varagona MJ, Purugganan M, Wessler SR. 1992. Alternative splicing induced by insertion of retrotransposons into the maize waxy gene. Plant Cell 4:811–820.
- Vicient CM, Kalendar R, Schulman AH. 2001. Envelope-class retrovirus-like elements are widespread, transcribed and spliced, and insertionally polymorphic in plants. Genome Res. 11:2041–2049.
- Vogel JP, et al. 2010. Genome sequencing and analysis of the model grass *Brachypodium distachyon*. Nature 463:763–768.
- Walbot V. 1999. UV-B damage amplified by transposons in maize. Nature 397:398–399.
- Walker AR, Lee E, Bogs J, McDavid DA, Thomas MR, Robinson SP. 2007. White grapes arose through the mutation of two similar and adjacent regulatory genes. Plant J. 49:772–785.

- Walker EL, Robbins TP, Bureau TE, Kermicle J, Dellaporta SL. 1995. Transposon-mediated chromosomal rearrangements and gene duplications in the formation of the maize *R-r* complex. EMBO J. 14: 2350–2363.
- Wang H, et al. 2005. SVA elements: a hominid-specific retroposon family. J Mol Biol. 354:994–1007.
- Wang W, et al. 2006. High rate of chimeric gene origination by retroposition in plant genomes. Plant Cell 18:1791–1802.
- Wessler SR, Baran G, Varagona M, Dellaporta SL. 1986. Excision of *Ds* produces waxy proteins with a range of enzymatic activities. EMBO J. 5:2427–2432.
- Wolff P, et al. 2011. High-resolution analysis of parent-of-origin allelic expression in the *Arabidopsis* endosperm. PLoS Genet. 7: e1002126.
- Wood TE, et al. 2009. The frequency of polyploid speciation in vascular plants. Proc Natl Acad Sci U S A. 106:13875–13879.
- Woodrow P, et al. 2012. Ty1-*copia* group retrotransposons and the evolution of retroelements in several angiosperm plants: evidence of horizontal transmission. Bioinformation 8:267–271.
- Xiao H, Jiang N, Schaffner E, Stockinger EJ, van der Knaap E. 2008. A retrotransposon-mediated gene duplication underlies morphological variation of tomato fruit. Science 319:1527–1530.
- Xu JH, Messing J. 2006. Maize haplotype with a *helitron*-amplified cytidine deaminase gene copy. BMC Genet. 7:52.
- Xu L, et al. 2007. *Triton*, a novel family of miniature inverted-repeat transposable elements (MITEs) in *Trichosanthes kirilowii* Maximowicz and its effect on gene regulation. Biochem Biophys Res Commun. 364: 668–674.

- Yang L, Bennetzen JL. 2009. Distribution, diversity, evolution, and survival of *Helitrons* in the maize genome. Proc Natl Acad Sci U S A. 106: 19922–19927.
- Yao J, Dong Y, Morris BA. 2001. Parthenocarpic apple fruit production conferred by transposon insertion mutations in a MADS-box transcription factor. Proc Natl Acad Sci U S A. 98:1306–1311.
- Zabala G, Vodkin LO. 2005. The *wp* mutation of *Glycine max* carries a gene-fragment-rich transposon of the *CACTA* superfamily. Plant Cell 17:2619–2632.
- Zeh DW, Zeh JA, Ishida Y. 2009. Transposable elements and an epigenetic basis for punctuated equilibria. Bioessays 31:715–726.
- Zhai J, et al. 2008. Small RNA-directed epigenetic natural variation in *Arabidopsis thaliana*. PLoS Genet. 4:e1000056.
- Zhang J, Peterson T. 2005. A segmental deletion series generated by sisterchromatid transposition of *Ac* transposable elements in maize. Genetics 171:333–344.
- Zhang J, et al. 2009. Alternative *Ac/Ds* transposition induces major chromosomal rearrangements in maize. Genes Dev. 23:755–765.
- Zhang J, Zhang F, Peterson T. 2006. Transposition of reversed *Ac* element ends generates novel chimeric genes in maize. PLoS Genet. 2:e164.
- Zhang P, et al. 2012. A transposable element insertion within *ZmGE2* gene is associated with increase in embryo to endosperm ratio in maize. Theor Appl Genet. 125:1463–1471.
- Zhang YM, et al. 2012. Functional analysis of the *HS185* regulatory element in the rice *HSP70* promoter. Mol Biol Rep. 39:1649–1657.

Associate editor: Ellen Pritham