

Useful parasites: the evolutionary biology and biotechnology applications of transposable elements

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Abstract

Transposable elements usually comprise the most abundant nongenic fraction of eukaryotic genomes. Because of their capacity to selfreplicate and to induce a wide range of mutations, transposable elements have long been considered as ‘parasitic’ or ‘selfish’. Today, we recognize that the findings about genomic changes affected by transposable elements have considerably altered our view of the ways in which genomes evolve and work. Numerous studies have provided evidences that mobile elements have the potential to act as agents of evolution by increasing, rearranging and diversifying the genetic repertoire of their hosts. With large-scale sequencing becoming increasingly available, more and more scientists come across transposable element sequences in their data. I will provide examples that transposable elements, although having signatures of ‘selfish’ DNA, play a significant biological role in the maintainance of genome integrity and providing novel regulatory networks. These features, along with the transpositional and mutagenic capacity to produce a raw genetic diversity, make the genome mobile fraction, a key player in species adaptation and microevolution. The last but not least, transposable elements stand as informative DNA markers that may complement other conventional DNA markers. Altogether, transposable elements represent a promising, but still largely unexplored research niche and deserve to be included into the agenda of molecular ecologists, evolutionary geneticists, conservation biologists and plant breeders.

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Introduction

In the late 1940s, Barbara McClintock challenged the existing concepts of genome organization and functioning when she discovered genes prone to mobility (McClintock 1950), which were later called ‘transposable elements’ (TEs). Although, the existence of TEs was accepted relatively soon after by the scientific community, the biology and applications of mobile genetic elements took decades to be widely recognized. With the discovery that many of these sequences are able to selfreproduce and to induce mutations, the selfish or parasitic DNA hypothesis was born. It said that these sequences served no function in the host organism, but were simply maintained by their ability to replicate and spread copies of themselves within and even between genomes (Doolittle and Sapienza 1980; Orgel and Crick 1980). In this ‘selfish’ way, TEs introduce genomic conflict trying to maximize their own fitness at the expense of the host’s genes (Burt and Trivers 2006; Werren 2011). Although, the TEs are

primarily selfish and having deleterious effects, their activities may occasionally and stochastically confer a fitness advantage to their hosts. Nowadays, with the improvement of molecular tools for genome analysis including next generation sequencing technologies, the majority of scientists recognize that mobile elements, even behaving selfishly, play a significant biological role in the maintainance of genome integrity and diversification of the genetic repertoire of their hosts. Nevertheless, there is still an underestimation and/or lack of comprehension among scientists about the opportunity of studying TEs for resolving important research issues. The aim of this review was to highlight the significance of TEs as enhancers of genome dynamics and evolution, and to further disseminate this research issue to the biological community. First, I will provide a short overview of TEs, their distribution among eukaryotes and relation to genome size variation. Then, I will emphasize the evolutionary consequences of TEs for genome functioning and integrity through some examples in the plant and animal kingdoms. Finally, I will focus on the practical applications and perspectives of TEs for genome analysis and manipulation.

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Structure and abundance of TEs

TEs are DNA sequences that can change their positions within the genome through the mechanism called transposition. There are two major types of TEs depending on their mechanism of transposition (Wicker *et al.* 2007). Class II elements (or DNA transposons) move through ‘cut and paste’ mechanism which comprises excision of a TE copy from one place in the genome followed by its reinsertion into another place. DNA transposons are present in low–moderate copy numbers in almost all eukaryotes. Class I retrotransposons transcribe RNA intermediates from genomic copies that are reverse-transcribed into a double-stranded DNA and integrated into a new position. Several daughter copies can be produced from a mother copy and insert throughout the genome, this proliferation has made retrotransposons a major fraction of large genomes. Eukaryote genomes are populated with large fractions of TEs. In plants, the repeat sequences range from 3% of the small genome of *Utricularia gibba* (Ibarra-Laclette *et al.* 2013) to more than 85% of the large genomes of cereals and maize (Li *et al.* 2004; Schnable *et al.* 2009). The human genome consists of about 65% of TEs (de Koning *et al.* 2011).

C-value paradox, transposable elements and the concept of ‘junk’ DNA

With the availability of many sequenced genomes (Michael and Jackson 2013) and the recently updated information of whole-genome sequences, it became evident that there is a striking variation in genome size in angiosperms which is often poorly correlated with the number of genes that code for proteins or the presumed evolutionary complexity of species (although we still do not understand how to meaningfully measure an organism’s complexity). This observation is the core of the C-value paradox raised more than 30 years ago where ‘C’ stands for the haploid genome size (Gall 1981). Genome sizes substantially vary between taxas: more than 7000-fold among animals (Gregory and DeSalle 2005) and 2400-fold across land plants as evidenced in the recent update of the plant genome size database containing data for 8510 species (Bennett and Leitch 2012; Garcia *et al.* 2014). A similar trend is observed between close species at a same ploidy level and apparently comparable complexity. A typical example is the plant genus, *Eleocharis*, which comprises more than 250 species, in which *E. acicularis* ($2n = 20$, $C = 0.25$ pg) is 20 times smaller than *E. palustris* ($2n = 16$, $C = 5.5$ pg) (Zedek *et al.* 2010). This genome size variation results from a substantial fraction of extra DNA other than for protein-coding genes and regulatory sequences, and is comprised of introns, pseudogenes, satellite sequences and TEs. This noncoding DNA has been considered a ‘junk DNA’ that has no biological function and conveys little or no selective advantage to the organisms (Orgel and Crick 1980).

The recent discovery that some regions of the noncoding DNA play important functions in genome and cell integrity has challenged the concept of useless ‘junk’ DNA. The

noncoding DNA was found to be essential for maintaining the chromosome structure, the function of centromeres, binding of transcription factors and coding for RNA involved in gene silencing. The ENCODE project represents a significant breakthrough study into the understanding the proportion of human’s genome that is functional. A function was assigned for 80% (dominated by RNA transcription, which alone covered 62%) of the genome particularly outside of protein-coding genes (ENCODE Project Consortium 2012). ENCODE and other studies have indicated that 5–20% of the human genome is under detectable selective pressure. Interestingly, the genome fraction directly involved in gene regulation was found significantly higher (up to 8-fold) than that ascribed to protein-coding exons (1%). Soon after, the ‘function’ by any meaningful sense of the word of the ENCODE-defined functional elements has been criticized and questioned (Eddy 2012; Doolittle 2013; Graur *et al.* 2013; Niu and Jiang 2013). These authors argued that DNA is still ‘junk’ despite the fact that it may bind transcription factors and contain regions of modified chromatin. ENCODE project represents the first genomewide functional annotation of the human genome. However, it does not directly address the ‘junk’ DNA concept which still remains viable.

TEs, particularly retrotransposons, comprise a major fraction of the ‘junk’ DNA and to a larger extent contribute to genome size expansion and variation among species, even as gene numbers remain relatively constant (table 1). The *Arabidopsis* genome, e.g. contains about 27,000 genes and 20 to 25 Mb of retrotransposons, whereas the maize genome contains about 40,000 but more than 1800 Mb of retrotransposon sequences (Liu and Bennetzen 2008; Baucom *et al.* 2009; Schnable *et al.* 2009). The ‘selfish’ nature of TEs acting as molecular parasites and functioning for themselves rather than having an adaptive function for their host genome was postulated by Doolittle and Sapienza (1980) and Orgel and Crick (1980). Since then, TEs have been a subject of tremendous interest because of their abundance, functionality and role in genome evolution. Do TEs embrace the concept of useless and nonfunctional ‘junk’ DNA? Definitely not for the whole genome fraction of TEs. First of all, many TEs are functional. Their DNA is biochemically active, encode proteins, bind proteins, synthesize regulatory RNA, thus meets the ENCODE criterion of functional elements. Second, many TEs are not ‘junk’ as there is a plenty of ways (see below) through which they provide a benefit to their host genomes. There is, however, a substantial fraction of decaying dead TEs generated from active TEs in the evolutionary past. It is suggested that different loads of such TE relics mostly explains the C-value paradox with larger genomes having a larger fraction of them. One should accept that there is a portion of DNA that seems to serve little useful purpose for the organism. However, I would like to point out on the inappropriateness of referring to DNA sequence as ‘junk’ unless we do not completely understand or have characterized it. DNA that appears useless today may provide a reservoir of sequences from which potentially advantageous new genes

Table 1. TE content (%) in representative flowering plant genomes.

Organism	Genome size (Mbp)	TE content (%)	Retro-TEs (%)	DNA-TEs (%)	Unknown (%)
Dicotyledons					
<i>Arabidopsis thaliana</i>	125	18.5	7.5	11.0	–
<i>Fragaria vesca</i>	240	20.7	14.7	5.2	–
<i>Medicago truncatula</i>	375	18.3	16.9	1.4	–
<i>Vitis vinifera</i>	487	21.5	19.4	1.4	0.7
<i>Malus × domestica</i>	742	42.4	37.6	0.9	3.9
<i>Solanum lycopersicum</i>	900	63.2	62.3	0.9	–
<i>Glycine max</i>	1115	58.7	42.2	16.5	–
Monocotyledons					
<i>Brachypodium distachyon</i>	272	28.1	23.3	4.8	0.9
<i>Oryza sativa</i>	389	39.5	25.8	13.7	–
<i>Setaria italica</i>	423	46.4	31.6	9.4	5.4
<i>Musa acuminata</i>	523	43.7	42.4	1.3	–
<i>Sorghum bicolor</i>	730	62.0	54.5	7.5	–
<i>Zea mays</i>	2300	84.2	75.6	8.6	–
<i>Hordeum vulgare</i>	5100	58.4	52.7	5.0	0.7
<i>Triticum aestivum</i>	17000	79.8	63.7	14.9	1.2

The content of this table is from Oliver K. R., McComb J. A. and Greene W. K. 2013 Transposable elements: powerful contributors to angiosperm evolution and diversity. *Genome Biol. Evol.* **5**, 1886–1901.

can emerge in future (Burt and Trivers 2006). In this way, it may be an important genetic basis for evolutionary innovation. After all, there might be a small amount of DNA that is a true junk. This DNA may just act as a protective buffer against the accumulation of harmful mutations.

In parallel to the interplay between genome size variation and TE accumulation, a recent study explored the link between TE diversity (types and number of TEs) and genome expansion (Elliot and Gregory 2015). The authors showed that this correlation is straightforward only to a certain point of genome size (specifically, around 500 Mbp), and then manifested by either a lack of relationship in animals or a negative correlation in plants. The likely common scenario may be that TE diversity and abundance increase as genomes expand up to a moderate size, whereas further genomic growth beyond this point is driven by a proliferation and divergence of a small subset of TE superfamilies. For example, 50% of the barley genome is made up of just 14 families, 12 of which long-terminal repeats (LTR) retrotransposons (Wicker *et al.* 2009). Also, retrotransposons *BARE-1*, *Wis* and *Angela* account for more than 10% of the *Triticeae* genomes. Consistently, differences in *BARE-1* abundance primarily explain genome size variation among *Hordeum* species (Vicent *et al.* 1999). One point that should be emphasized, however, is that the establishment of a more or less stable equilibrium between genome size and TE proliferation within and among taxa is influenced by several other selection factors like population size, mating system, polyploidization events and ecogeographical distribution. The complexity of the interacting factors means that many comparative studies need to be done before patterns of TE abundance and diversity versus taxon-specificity and genome size can be described and understood.

Epigenetic silencing drives the ups and downs of transposable elements

Why TEs, apparently useless and potentially damaging, are widely spread in higher organisms? Generally, the abundance and accumulation rates of TEs result from a balance between two main forces: TE transposition leading to an increase in copy number and, from the other side, the elimination and inactivation of TEs mediated by mutations in their sequences or through the process of ectopic recombination between TE copies at nonhomologous loci (Charlesworth and Charlesworth 1983). Recombination events between TEs at nonhomologous chromosome locations lead to the generation of truncated inactive elements thus reducing their functional activity and accumulation rate. This balance is widely achieved by epigenetic silencing of TEs by siRNA-directed DNA methylation (Ito 2012). The mechanism consists of the synthesis of small TE-specific noncoding RNAs that guide DNA methylation and silencing of homologous DNA sequences at posttranscriptional level (Lister *et al.* 2008). Differential expression of TEs influenced by siRNA-directed DNA methylation was observed between different plant genomes (Alzohairy *et al.* 2014). For instance, differences in TE accumulation and genome size are largely influenced by the extent of TE-silencing as shown in a comparison study between *Arabidopsis lyrata* and *A. thaliana* (Hollister *et al.* 2011). Fedoroff (2012) argued that actually the evolution of epigenetic mechanisms, that control homology-dependent recombination, is driving the accumulation rate of TEs in a long-term aspect. The methylation in prokaryotes represents a nuclear defense system to limit the destructive potential of ‘parasitic sequences’ including TEs (Yoder *et al.* 1997). Consistently, the epigenetic control is not so efficient in

prokaryotes and lower eukaryotes, and ectopic recombination among dispersed TEs would rapidly eliminate them either directly by deletions or indirectly by creating non-viable chromosomes. As a result, these processes keep the genomes of prokaryotes and many lower eukaryotes small. In contrast, higher eukaryotes have larger genomes due to more stringent epigenetic control and lower recombination rates that allow the accumulation of TEs.

Transposable elements are capacitors of genome dynamics

It is obvious that the high frequency of TEs, their capacity to change the location within the host genome and to induce chromosome aberrations (e.g. deletions, duplications and insertions) would confer a potentially negative impact of TEs on genome integrity. Transposable elements can affect genome dynamics with possible effects on phenotypes through multiple mechanisms, depending on the TE itself and its insertion site. Some examples of the effects of TE insertions on gene structure and function are shown in figure 1.

Transposable elements-mediated alterations in the structure and function of genes

The most obvious TE-induced change is gene disruption leading to observable loss of function. A classical example

are mutations in the gene coding for the colour of the maize kernels provoked by the Ac element detected by Barbara McClintock (figure 1c). Similarly, the wrinkled seed phenotype studied by Mendel in pea was found to result from an insertion of a TE into a locus encoding a starch-branching enzyme (Bhattacharyya *et al.* 1990). Beside plant genomes, ~0.3% of all human mutations are caused by TE insertions or rearrangements (Cordaux and Batzer 2009). The currently active nonLTR retrotransposons, *L1* (LINE 1), SVA and *Alu* (SINE), are reported to be the causative factors of many genetic disorders, such as haemophilia A, Apert syndrome, familial hypercholesterolaemia, colon and breast cancer, muscular dystrophy etc. (for review see O'Donnell and Burns 2010; Ayarpadikannan and Kim 2014). *Alu* elements (named for the enzyme used to identify it) are short sequences (~300 bp) that occur almost a million times in the human genome and comprise up to 3.5% of the total DNA. Transposition of *Alu* elements to sites in and near genes, or *Alu*-mediated ectopic recombination events can have occasional deleterious effects on genes. Indeed, many cloned genes were shown to harbour *Alu* elements in their sequences. On the other hand, *Alu* sequences likely have positive regulatory functions as mutations within them have been associated with cancer. Transposable elements can also disrupt existing regulatory motifs (repressors or enhancers) or to provide new regulatory information thereby influencing the gene expression and causing mutant phenotypes.

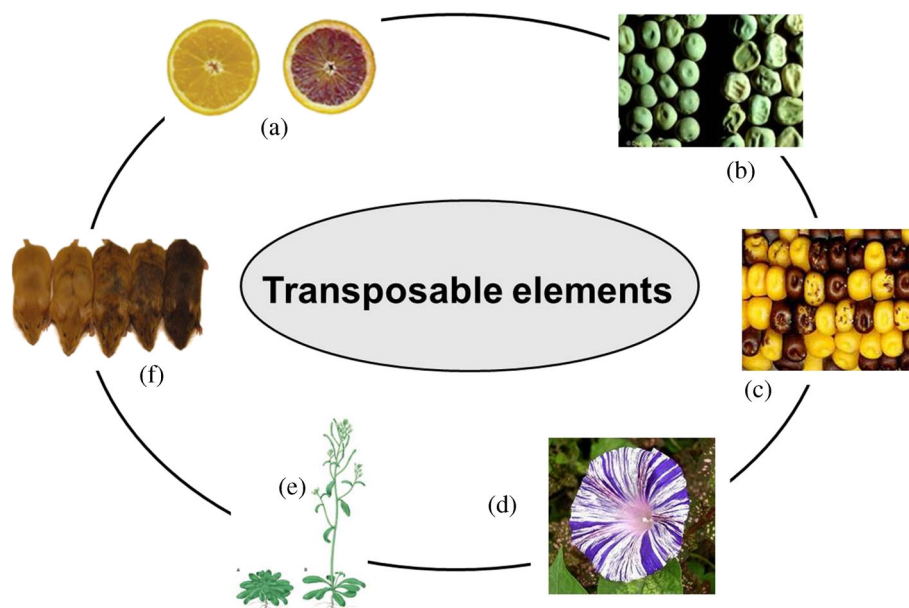


Figure 1. Effects of TE insertions on gene structure and function. (a) Insertion of LTR retrotransposon upstream of the *Ruby* gene in the blood orange provides a new promoter controlling the expression of a gene for flesh colouration of the fruit (Butelli *et al.* 2012). (b) Insertion of TE into the *rugosa* locus encoding a starch-branching enzyme in pea results in a wrinkled seed (Bhattacharyya *et al.* 1990). Insertion of TE into a gene that produces anthocyanin pigments leads to its inactivation and changes in the colour: yellow seeds in maize. (c) White sectors in *Petunia hybrida* (d) Kroon *et al.* (1994). Revertant sectors (dark-spotted seeds or blue stripes) appear when the transposon spontaneously excises from that gene in certain cells and the pigment production is restored. (e) siRNA-controlled methylation of a TE insertion influence the expression of the *FLC* locus (a gene delaying flowering) and leads to earlier flowering in *A. thaliana* (Liu *et al.* 2004). (f) Alterations in the transcription levels of genes by antisense transcription from adjacent TE insertions as observed for the *agouti* colour gene in mice (Morgan *et al.* 1999).

Several examples about such effects of TEs are reported in *Drosophila* (Lerman *et al.* 2003; Lerman and Feder 2005; McCart and Ffrench-Constant 2008) and plants (Salvi *et al.* 2007; Studer *et al.* 2011; Butelli *et al.* 2012).

Transposable elements affect the epigenetic regulation of genes

A substantial part of TEs are targeted by siRNA-directed DNA methylation that is repressing their activity. This epigenetic silencing can spread to genes located in the vicinity of TE insertions and may generate stable epialleles of potential evolutionary relevance (Slotkin and Martienssen 2007; Wang *et al.* 2013). For instance, an insertion of a LTR retrotransposon near the *agouti* gene in mice alters the chromatin state and DNA methylation at this locus (Morgan *et al.* 1999). Variations in the epigenetic status of the retrotransposon ultimately influence the gene transcript level and the colour of the mice coat (figure 1f). Similarly, early flowering of the *Ler* ecotype of *A. thaliana* is controlled by a DNA transposon insertion in the *FLC* gene responsible for a delayed flowering. This insertion is targeted by TE-derived siRNAs which results in the epigenetic silencing of the gene (Liu *et al.* 2004).

Transposable elements can mediate genome restructuring

Recombination events between TE copies at nonhomologous sites in the genome (ectopic recombination) either on one or different chromosomes is important mechanism by which TE mediate genome restructuring (deletions, insertions, inversions, translocations and duplications), thus promoting chromosomal instability (Gray 2000). In addition, TEs can enhance genome reshuffling by capturing and transferring genes within genomes (Jiang *et al.* 2004; Morgante *et al.* 2005; Schaack *et al.* 2010).

The number and the variety of mutations induced by TEs is extraordinary and can be hardly embraced. One can still argue that TEs have predominantly negative impact and increase the genetic load and scientists ask the question: Do TEs, through the induction of mutations or regulatory functions, provide some benefit to the organism itself or just stand as pure 'selfish' and detrimental DNA?

TEs have important biological functions

During the last two decades, a major focus has been on the positive contribution of TEs to the evolution of gene regulation and diversification of host genes. The major break-through in this area was achieved with the advent of high-throughput sequence technologies and software platforms for the annotation of TEs. We have tremendously enriched our knowledge about the biology of TEs and their interaction with other components of the host genome. One way in which TEs contribute to evolution is that their sequences (e.g. genes, binding sites, and terminal repeats) can be coopted to perform functions beneficial for the proper functioning of the host

genome (Sinzelle *et al.* 2009). This genomic coopting of a molecular parasite is often referred to as molecular domestication or exaptation (Gould and Vrba 1982; Miller *et al.* 1999). Here I present examples of the different ways that TEs have evolved from strictly parasitic elements to mutualistic sequences that benefit their host genomes (see also table 2).

Transposable elements as a source of novel regulatory networks

The human genome provides the majority of examples about TEs involved in domestication which has helped to spur remarkable evolutionary innovations. The initial analysis of the human genome has revealed that ~25% of human promoter regions and ~4% of human exons contain sequences derived from TEs (Lander *et al.* 2001; Nekrutenko and Li 2001; Jordan *et al.* 2003; Kapitonov and Jurka 2005; Jurka *et al.* 2007).

V(D)J recombination is a unique mechanism of genetic recombination that occurs only in developing lymphocytes during the early stages of T and B cell maturation. The process results in a highly diverse repertoire of antigen receptors in these cells and is a distinguished feature of the adaptive immune system in vertebrates. The V(D)J recombination is mediated by genes *RAG1* and *RAG2* that are evolutionarily derived from ancient insertions of *Transib* DNA transposons (Zhou *et al.* 2004; Ramsden *et al.* 2010). As expected, based on our knowledge about class II TEs transposition, this process involves the generation of double-strand DNA breaks in a way that is mechanistically similar to the 'cut' component of 'cut and paste' transposition.

Repeat-induced gene silencing involving L1 retroelements has been hypothesized for X-chromosome inactivation, which is necessary to maintain the proper gene dosage in females. Inactivation is initiated at the X-chromosome inactivation centre (XIC) from which the silencing signal spreads along the chromosome. According to one hypothesis, LINE retrotransposons might trigger the heterochromatization in XIC centre and boost the efficient spread of the silencing away this centre (reviewed in Lyon 2006). In support to this idea, the X chromosomes of many mammals, including humans, are rich in LINE elements, except in regions that are prone to escaping X inactivation (Ross *et al.* 2005). It is still unknown whether LINEs function in the spread of heterochromatin on the X chromosome or their enrichment may simply be a consequence of the heterochromatic nature of the inactive X. However, Cohen *et al.* (2007) have shown that short tandem repeats homologous to retrotransposons regulate X-chromosome inactivation by producing bidirectional transcripts in differentiating mouse cells, thus providing indirect evidence that TEs function in both the initiation and spread of X inactivation.

Transposable elements are also functionally implicated in the proper functioning of the mammalian embryo at earliest stages of its development (Macfarlan *et al.* 2012; Tomkins 2013). In this line, it was recently reported that a family of TEs in mammals provide enhancer sequences that modulate

Table 2. Examples of adaptive mutations and exaptations provided by transposable elements.

Affected gene/s	Phenotype	Host species	TE	Reference
<i>Mustang</i> and <i>Sleeper</i> gene families	Growth, flower development, reproduction	Angiosperms	<i>Mutator</i> -like, <i>hAT</i>	Joly-Lopez <i>et al.</i> (2012) Knip <i>et al.</i> (2012)
<i>GmphyA2</i> (phytochrom A)	Adaptation to high latitudes	Soybean	<i>SORE1</i>	Kanazawa <i>et al.</i> (2009)
<i>FAR1</i> and <i>FHY3</i>	Adaptation to light intensity	<i>Arabidopsis</i>	<i>MuDR</i>	Lin <i>et al.</i> (2007)
Teosinte branched 1 (<i>tb1</i>)	Branch outgrowth	Maize	<i>Hopscotch</i>	Studer <i>et al.</i> (2011)
+ <i>ZmCCT</i>	Attenuation of photoperiod sensitivity	Maize	<i>CACTA</i> -like	Yang <i>et al.</i> (2013)
Flowering locus C (<i>FLC</i>)	Flowering time	<i>Arabidopsis</i>	<i>Mutator</i> -like	Liu <i>et al.</i> (2004)
Blast resistance gene <i>Pit</i>	Resistance to fungal infection	Maize	<i>Renovator</i>	Hayashi and Yoshida (2009)
<i>Rim2</i>	Resistance to fungal infection	Rice	<i>CACTA</i> -like	He <i>et al.</i> (2000)
<i>RPP7</i>	Resistance to pathogen	<i>Arabidopsis</i>	<i>COPIA-R7</i>	Tsuchiya and Eulgem (2013)
Juvenile hormone epoxide hydrolase 2 (<i>Jheh2</i>)	Reduced viability	<i>D. melanogaster</i>	<i>Tc1</i> -like	Gonzalez <i>et al.</i> (2009)
<i>CHKov1</i>	Resistance to viral infection and insecticides	<i>D. melanogaster</i>	<i>Doc</i> element	Aminetzach <i>et al.</i> (2005) Darboux <i>et al.</i> (2007) Magwire <i>et al.</i> (2011)
<i>Cyp6g1</i>	Resistance to pesticides	<i>D. melanogaster</i>	<i>Accord</i>	Schmidt <i>et al.</i> (2010)
	Maintainance of telomeres	<i>D. melanogaster</i>	<i>HeT-A</i> , <i>TART</i> and <i>TAHRE</i>	Abad <i>et al.</i> (2004) Shpiz <i>et al.</i> (2007)
<i>CENP-B</i> <i>RAG1/2</i>	Formation of centromeres V(D)J recombination	Human Vertebrates	<i>Pogo</i> <i>hAT</i> , <i>Transib</i>	Casola <i>et al.</i> (2008) Ramsden <i>et al.</i> (2010) Zhou <i>et al.</i> (2004) Kapitonov and Jurka (2005)
	X-chromosome inactivation	Human	<i>L1</i> retroelements	Bailey <i>et al.</i> (2000) Lyon (2006)

the gene expression in placental cells thus regulating the interaction between the mother and the offspring (Chuong *et al.* 2013). In humans, the highly conserved centromere-binding protein CENBP facilitates centromere formation and is derived from transposases of the *pogo* DNA transposon family (Casola *et al.* 2008).

Several examples about TE exaptation are available in the plant kingdom as well. In *Arabidopsis*, *FHY3* and *FAR3* are transcription factors, related to the *MuDR* family of transposases, that bind to promoter regions and activate several genes involved in far-red light and circadian clock signalling (Hudson *et al.* 2003; Lin *et al.* 2007).

A recent study in *Arabidopsis* has shown that the *COPIA-R7* transposon, inserted into the plant disease resistance gene *RPP7*, enhances its host's immunity to a pathogenic microorganism from a large group of fungus-like parasites that cause a number of plant diseases (Tsuchiya and Eulgem 2013). The *Rim2* gene implicated in defence against fungal infection appears to have been directly exapted from part of a *CACTA* DNA TE element (He *et al.* 2000). The rice blast disease resistance gene *Pit* was refunctionalized by the recruitment of a copia-like LTR element as a promoter (Hayashi and Yoshida 2009).

Relatively few data are available about the direct role of TEs in the processes of species domestication. An insertion of the TE *Hopscotch* in the regulatory region of the maize domestication gene, teosinte branched1 (*tb1*), acts as an enhancer of its expression and confers the increased apical dominance in maize compared to its progenitor, teosinte (Studer *et al.* 2011). Insertion of a *CACTA*-like transposon into the promoter of the gene *ZmCCT*, which modulate photoperiod sensitivity, can suppress its expression thus enhancing the spread of maize to long-day temperate regions (Yang *et al.* 2013). *Mustang* and *Sleeper* gene families present in flowering plants have sequences derived from exapted transposases from *Mutator*-like and *hAT* DNA elements, respectively (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.

Transposable elements maintain chromosome stability and functioning

The coexistence of TEs and the host genome has resulted in several regulatory pathways, including a combination

of various epigenetic mechanisms, i.e. DNA methylation, small RNAs and histone posttranslational modifications. In *A. thaliana*, methylated TEs promote the epigenetic silencing and the formation of heterochromatin (Fagegaltier *et al.* 2009). TEs are abundant in heterochromatin-rich regions, centromeres (Casola *et al.* 2008; Mateo and González 2014) and telomeres (Levin and Moran 2011; Pardue and DeBaryshe 2011). Such phenomena might be the consequence of: (i) insertional preferences of TEs into heterochromatin; (ii) positive selection of TE maintenance into heterochromatin for genomic stability; or (iii) an induction of heterochromatin by TE sequences. An example of a host using the movement of a retrotransposon to its advantage was found in the telomere maintenance of *Drosophila*. Two retrotransposons, *HeTA* and *TART*, are present in multiple copies on the telomeres and maintain their length in replicating cells (Abad *et al.* 2004; Shpiz *et al.* 2007). A correlation between TEs activity and the maintenance of heterochromatin integrity was also demonstrated by the observation that the increase in TE expression and transposition leads to an age-related breakdown in heterochromatin structure and subsequent cellular dysfunction (Wood and Helfand 2013). St Laurent *et al.* (2010) argued that the stress-induced activation of LINE/L1 elements, particularly prevalent in mammals, leads to mutagenic insertions and DNA damage that accumulate with age and can cause genomic instability even in the absence of a successful transposition event. This fact shows that LINE elements may play an important role in mammalian ageing and evolution. In support to these data a study in budding yeast, using a chronological ageing model, observed that the yeast retrotransposon *Ty1* showed increased mobility with age, and this was correlated with chromosome rearrangements and other hallmarks of genomic instability (Maxwell *et al.* 2011).

Matrix attachment regions (MARs) are DNA sequences that bind to the nuclear matrix forming functionally independent chromatin domains. Colocalization of TEs with MARs located in introns and 5' flanking regions of genes was observed which suggests the putative role of mobile elements and/or their products (transposases) in the formation of these structures (Avramova *et al.* 1998; Tikhonov *et al.* 2000; Pathak *et al.* 2014). For instance, the insulator protein BEAF-32 in *Drosophila* is entirely derived from a hAT transposase and is involved in chromatin functioning through interactions with the nuclear matrix (Pathak *et al.* 2007). A cross talk between the distribution and genome organization of BEAF-32 contributes to new gene-expression profiles and distinct phenotypes with a putative role in the evolution. Nevertheless, the functional link between MAR and retrotransposons remains to be comprehensively investigated.

Although, several domesticated TEs have been reported so far, their total number may be much higher than currently reported and many more domesticated TEs may await discovery in the near future. Traditional genetic methods may be insufficient to find such TEs and to assess their evolutionary and functional impact, e.g. due to functional

redundancy. Systematic searches that exploit genomic signatures of natural selection have been employed to identify potential domesticated genes, but their predictions have yet to be experimentally verified.

TEs as capacitors of species adaptation to changing environments

Most TEs are usually in a transpositionally silent state but can be occasionally activated in response to different environmental stimuli (Grandbastien 1998), a phenomenon what Barbara McClintock called 'genome shock' (McClintock 1984). The major question is whether TE activity is just an undesired side effect of stress exposure or TE-induced genetic diversity accounts for microevolutionary processes such as rapid adaptive evolution and speciation in natural populations. A prevalent view among evolutionary biologists is that the vast majority of TE insertions are nearly neutral and unlikely to have a strong evolutionary impact. Some host forces do indeed select against TE insertions (due to the deleterious impact of insertions or of their effects through ectopic recombination) and to efficiently purge deleterious insertions. However, the selection on TEs accumulation at population level is affected by a complex of factors and the outcome is often difficult to be predicted. Nevertheless, several reports provide convincing evidence about a turnover of TEs closely matching ecogeographical distribution of gene pools. For instance, increase of full length copies of the retrotransposon *BARE-1* was observed in barley populations in dry environments compared to those grown a few dozen metre apart in less stressful habitats (Kalendar *et al.* 2000). A good example of the capacity of TEs to elicit mutational consequences potentially helpful in adapting to new environments is provided by three diploid sunflower species, *Helianthus anomalus*, *H. Deserticola* and *H. paradoxus*. The three hybrid taxa, independently derived through hybridization events between the two parental taxa, *H. annuus* and *H. petiolaris*, encountered a rapid, retrotransposon-mediated genome expansion and all of them occupy habitats considered abiotically extreme relative to either parental species (Ungerer *et al.* 2006; Kawakami *et al.* 2010). Another example of a TEs involved in the adaptation is the early flowering of the *Ler* ecotype of *A. thaliana* controlled by a DNA transposon insertion (Liu *et al.* 2004). In addition, Gonzalez *et al.* (2010) have shown that at least 32% of the putatively adaptive insertions screened in natural population of *D. melanogaster* have a distribution consistent with selection by contrasted ecogeographical conditions. Most of these TE insertions, belonging to multiple TE families, were linked to functional genes with distinct phenotypic changes.

Insights offered by such reverse population genomic approaches pinpoint the importance of TEs as a source of adaptive variation. However, such surveys should be undertaken in a wider range of species to reliably estimate the impact of TEs on their evolutionary ecology.

Transposable elements are reliable DNA molecular markers

The high copy number, chromosome coverage and variable arrangement pattern even among closely related species give TEs advantage as informative markers to assess natural and stress-induced genetic diversity, and to enhance marker assisted selection (MAS) in plant breeding programmes. Both DNA transposons and retrotransposons can be utilized for the generation of markers, the latter ones being much more efficient. Retrotransposons have been found to comprise the most common class of TEs in eukaryotes, and represent up to 90% in plant genomes (Feschotte *et al.* 2002). They constitute for >50% of the maize and cereal genomes (Meyers *et al.* 2001; Brenchley *et al.* 2012) and 14% of the *Arabidopsis* genome (The *Arabidopsis* Genome Initiative 2000). Moreover, the presence of conserved domains at both ends (LTR) can be easily exploited for the design of PCR primers (Kalendar *et al.* 2011).

Major retrotransposon-based markers and type of inheritance

Retrotransposon-based molecular analysis relies on PCR amplification of DNA sequences (markers) using a primer corresponding to a retrotransposon and a primer matching a section of the neighbouring genome (e.g. microsatellite, restriction site or another TE copy). Among the wide diversity of retransposon-based techniques, the three most frequently used, as tools for diversity studies are sequence-specific amplified polymorphism (SSAP) (Waugh *et al.* 1997), interretrotransposon amplified polymorphism (IRAP) (Kalendar *et al.* 1999), retrotransposon microsatellite amplified polymorphism (REMAP) (Kalendar *et al.* 1999) and retrotransposon-based amplified polymorphism (RBIP) (Flavell *et al.* 1998). A schematic representation of the main marker methods is presented in figure 2. Several molecular marker systems based on the information available for the transposable elements sequences were developed to measure diversity, similarity and cladistic relationships in plants: barley (Kalendar *et al.* 2000; Vicient *et al.* 2001), citrus (Breto *et al.* 2001), genus *Malus* (Antonius-Klemola *et al.* 2006), rice (Branco *et al.* 2007), flax (Melnikova *et al.* 2014), pea (Ellis *et al.* 1998; Pearce *et al.* 2000; Smýkal 2006), wheat (Queen *et al.* 2004; Melnikova *et al.* 2011), pear (Kim *et al.* 2012) and others. Transposable elements find substantial application as genetic markers in animal kingdom as well. For example, the method transposon display has been recently used to study the effect of interspecies hybridization on TEs dynamics in *Drosophila* (Vela *et al.* 2014). In addition, mobile elements are found to be active and to provide polymorphism in human populations (Mills *et al.* 2007). Consistently, SINE (*Alu*) and LINE (*L1*) elements have been used to trace human roots to Africa (Witherspoon *et al.* 2006). Also, *L1* insertion polymorphisms have potential use in forensic analyses (Ray *et al.* 2007).

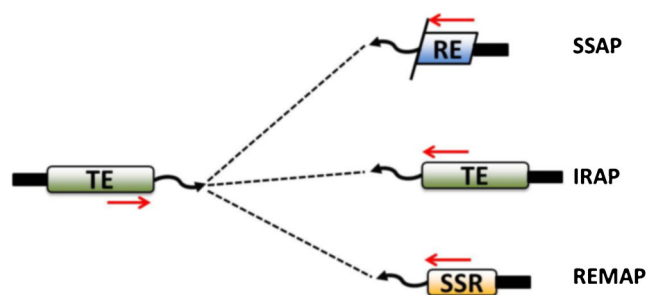


Figure 2. Main transposon-based marker methods tracking polymorphism in the insertion pattern of TEs. The techniques rely on simultaneous PCR amplification of sequences between copies of a candidate TE and adjacent genomic regions which can be other TE copies interretrotransposon amplified polymorphism (IRAP); microsatellite loci, retrotransposon-microsatellite amplified polymorphism (REMAP) or restriction sites, sequence-specific amplified polymorphism (SSAP). Red arrowheads indicate the primers for PCR amplification.

RBIP is the only retrotransposon-based method directing at polymorphism in the integration of an element at a single-copy locus and can distinguish between its heterozygous or homozygous state, thus having a codominant inheritance. Other marker techniques generally behave as dominant (i.e. presence/absence of a TE insertion) and do not allow one to discriminate the allelic state of the locus. Even though, it may be possible to map two polymorphisms to the same TE integration site but this is very tedious in practice and can not be determined directly. The difficulty comes from the complexity of band profiles in multitarget PCR reactions and the less sufficient resolution provided by the commonly used standard agarose gel and polyacrylamid gel electrophoresis (PAGE) methods of amplicon separation. The development of high-throughput marker method for genetic studies, the fluorescent SSAP system, improved the available amplicon number and the accuracy in their scoring which allow to discriminate alleles and to identify heterozygous loci at a resolution of a single nucleotide (Knox *et al.* 2009). Another way to overcome the dominant nature of the other marker systems is to use genetically homozygous material. For mapping populations, this can be achieved using double-haploid, recombinant inbred lines or haploid tissues like the endosperm of gymnosperms. The efficacy of double-haploid populations for the mapping of retrotransposon markers and genes has been well established (Waugh *et al.* 1997; Manninen *et al.* 2000).

Advantages of retrotransposon-based markers

There are few properties of TEs that give them an advantage over other DNA molecular markers.

Transposable elements are prone to activation by stress:

As mentioned earlier in this review, TEs are prone to activation by different stress factors. Beside the few examples of TEs dynamics at population level, a plethora of studies

have also demonstrated the stress-induced escape from the silent state for many TE families under control experimental conditions. In plants, these activating events include artificial interspecific hybridization and allopolyploidization (for review see Parisod *et al.* 2009; Yaakov and Kashkush 2011), infection (Melayah *et al.* 2001; Grandbastien *et al.* 2005), abiotic stresses like drought (Aprile *et al.* 2009), high and low temperatures (Ivashuta *et al.* 2002; Young *et al.* 2005), protoplast isolation, cell culture, wounding, methyl jasmonate, CuCl₂ and salicylic acid (Hirochika 1993; Moreau-Mhiri *et al.* 1996; Mhiri *et al.* 1997; Takeda *et al.* 1998). The activation of wheat retrotransposons under light and salinity stresses has been also reported (Woodrow *et al.* 2010). Stress response may differ between host genotypes possibly reflecting an adaptive response of ancestral populations to different stimuli. Therefore, these and more studies show without doubt that TEs can be used as reliable DNA markers to assess genome response to stress factors both at individual and population levels.

Although, the exact process of transcription induction remains not completely elucidated, it has been shown that TEs are likely to become activated by mechanisms similar to those employed by host defense genes. Indeed, promoter sequences of both TEs and host defense genes share nucleotide similarities and are likely to bind to similar transcription factors (Casacuberta and Santiago 2003). For instance, *Tnt1* retrotransposons from tobacco present regulatory regions with specific motifs that are commonly observed in genes induced by drought, anaerobic conditions or oxidative stresses (Grandbastien *et al.* 2005). Transposable elements could have captured promoters from normal stress-inducible genes or inversely, that they provided their own stress-inducible promoters to some plant defense genes (Grandbastien *et al.* 1997; Takeda *et al.* 1999).

Transposable elements display high level of insertional polymorphism: DNA marker techniques based on TEs are anonymous, producing fingerprints from multiple sites of retrotransposon insertions in the genome (Schulman *et al.* 2004). The outcome is a high degree of heterogeneity and insertional polymorphism observed both within and between species. One can still narrow the regions upon which one is looking for TE polymorphism. To achieve this goal, one should rely on the biology and insertional pattern of TEs. For instance, high copy number families like the active LTR retrotransposon *BARE-1* tend to form clustered (i.e. adjacent insertions) and nested (i.e. insertions within one another) arrangements in intergenic regions of large genomes. In contrast, insertions of short TEs such as Ac/Ds, MITEs, LINEs, SINEs, but also low copy number LTR retrotransposons, seem overrepresented near or within genes in plants and in humans (i.e. mostly in introns, 5' or 3' UTR as well as in flanking regions; Wright *et al.* 2003; Majewski and Ott 2002). In addition, retrotransposons allows to detect large genome changes and seem to be more informative as a

complement to convenient DNA markers like amplified fragment length polymorphism (AFLP), microsatellites (SSR) and single-nucleotide polymorphism (SNP) (Waugh *et al.* 1997; Ellis *et al.* 1998; Yu and Wise 2000; Porceddu *et al.* 2002; Tam *et al.* 2005) which mainly detect single nucleotide changes.

Retrotransposons are homoplasmy-free phylogenetic markers:

The regions of transposon insertions are thought to be more or less random. Thus, TE copies at exactly the same location in homeologous chromosomes within and among individuals appear unlikely. Retrotransposon integrations are also assumed to be irreversible events unless a chromosomal segment containing the repeat becomes deleted. An insertion at a specific genomic location suggests a derived character state, and species which share an insertion at a particular locus are grouped together on the tree. Lack of an insertion at an orthologous locus is regarded as an ancestral state and the corresponding individuals are considered basal in the phylogenetic tree. Thus, as opposed to reversible changes in DNA sequence composition, retrotransposon insertions have been claimed to generate homoplasmy-free phylogenetic markers that provide an extremely accurate picture of evolutionary relationships and have been proven very successful in elucidating problematic phylogenies. For example, SINE elements were evaluated as reliable cladistic markers to resolve phylogenetic relationships among human and nonhuman primates, clinical diagnostics of diseases and forensic identification (for review see Schmitz *et al.* 2005; Ray 2007; Konkel *et al.* 2010; Ray and Batzer 2011).

Transposable elements as a tool for gene manipulation

Insertional mutagenesis

As discussed earlier, TEs can change their genomic location upon activation. If inserted inside the coding or regulatory sequence of a gene, disruption of the reading frame can lead to a loss of gene function. This phenomenon provides the platform for the development of the technique called transposon mutagenesis which has been used to induce mutations, identify and study the function of the responsive gene. The standard mutagenesis platform consists of crossing genetic lines with inactive or nonautonomous TEs with lines containing an active (autonomous) element. The offspring carrying an autonomous transposon, through its transcribing transposase, can mobilize the nonautonomous transposon. The reactivated transposon can further insert randomly into new genomic sites thus causing mutations in the offspring. Hundreds and thousands of individuals can then be screened for a new mutation of interest. If such a phenotype is found, then it can be assumed that TE insertion has inactivated the gene responsible for this phenotype. Because, the sequence of TE is known, the gene can be easily identified either by sequencing the whole genome and searching for the sequence or

using PCR to amplify specifically that gene. Several examples are available on genes successfully tagged using TEs in species like tobacco (Fitzmaurice *et al.* 1999), maize (Howard *et al.* 2014) and tomato (van der Biezen *et al.* 1996).

Transposable elements as gene delivery vehicles

Virtually any DNA sequence of interest can be placed between the terminal inverted repeats (TIRs) of a TE and mobilized *in trans* by supplementing the transposase gene in the form of an expression plasmid or through mRNA synthesized *in vitro*. This feature makes TEs natural and easily controllable DNA delivery vehicles to transfer genes to a host organisms' chromosome for the purposes of introducing new traits and to discover new genes. The use of TEs as non-viral vehicles for persistent gene delivery has emerged with the discovery of DNA transposon 'sleeping beauty' as a tool for genetic modifications and persistent gene expression was demonstrated in a wide variety of vertebrate cell lines and species (Ammar *et al.* 2012). Several other synthetic DNA transposon system vehicles like the *PiggyBac* (PB) transposon with activity in mammalian cells have been studied and tested (for review see Skipper *et al.* (2013)). Transposon-based gene transfer is more efficient for stable expression of foreign genes in vertebrates compared to classic approaches as the latter ones rely on physical methods of gene constructs delivery: transfection, electroporation, sonoporation, or microinjection. The main drawbacks of these approaches are the low rates of genomic integration and unstable expression of the chromosomally integrated gene construct. This is believed to be associated with the phenomenon of concatemerization of the injected DNA before genomic integration and repeat-induced gene silencing (Henikoff 1998).

The ongoing investigations will certainly prompt new ideas and new designs to be developed in the expanding universe of TE-based technologies for genetic and cell engineering. Innovative aspect of these studies represents the evaluation of TE-based delivery systems as a potential approach for the therapy of human diseases. Despite the various examples of preclinical efficacy for *in vivo* gene therapy, the road to the clinic will wind through additional experimentation and evidence of therapeutic effects in large animal models.

Conclusions

Transposable elements occupy a significant portion of eukaryotic genomes and have been long time considered as noncoding 'junk DNA' with no beneficial functions for the host genome. With the advent of high-throughput sequence technologies, however, scientists are beginning to find that intrinsic relationships exist between TEs and the other part of the genome. Perhaps only a small proportion of these relationships evolve to become mutually beneficial over the long term. Transposable DNAs have been expertly integrated into the incredibly complex function of the genome as important

coordinators in many biological processes such as maintenance of chromosome integrity and creation of novel regulatory networks. The number of evidences for the benefits of TEs is constantly increasing, however, this is perhaps the tip of the iceberg. At the end, TEs are the main provider of genetic diversity which is the raw material to promote eucaryotic genome flexibility and evolution. Transposable elements can therefore be viewed as important genomic symbionts and their study should go 'hand-in-hand' with the other part of the genome. The fraction of TEs is often neglected in genomewide studies. For instance, the common practice of scientists is to mask repeat sequences in next generation data focussing on the 'good' stuff: coding regions. One should appreciate, however, that by throwing out all data on TEs, we are turning our back on important findings on genome functioning and evolution. In short, the selfish, junk and beneficial DNA hypotheses are all relevant but by no means mutually exclusive and single label for these relationships is inappropriate and potentially misleading. One of the key take homes from the numerous sequenced plant and animal genomes is that we still have a lot to learn about the organization of genomes, function of genes to comprehensively recognize the diverse functional importance of the 'junk' DNA.

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