

Modulating signaling networks by CRISPR/Cas9-mediated transposable element insertion

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Received: 4 September 2017 / Revised: 1 October 2017 / Accepted: 9 October 2017
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Abstract In a recent past, transposable elements (TEs) were referred to as selfish genetic components only capable of copying themselves with the aim of increasing the odds of being inherited. Nonetheless, TEs have been initially proposed as positive control elements acting in synergy with the host. Nowadays, it is well known that TE movement into host genome comprises an important evolutionary mechanism capable of increasing the adaptive fitness. As insights into TE functioning are increasing day to day, the manipulation of transposition has raised an interesting possibility of setting the host functions, although the lack of appropriate genome engineering tools has unpaved it. Fortunately, the emergence of genome editing technologies based on programmable nucleases, and especially the arrival of a multi-purpose RNA-guided Cas9 endonuclease system, has made it possible to reconsider this challenge. For such purpose, a particular type of transposons referred to as miniature inverted-repeat transposable elements (MITEs) has shown a series of interesting characteristics for designing functional drivers. Here, recent insights into MITE elements and versatile RNA-guided CRISPR/Cas9 genome engineering system

are given to understand how to deploy the potential of TEs for control of the host transcriptional activity.

Keywords CRISPR/Cas9 · TE-based drives · Genome editing · Miniature inverted transposable elements · Transcriptional control · Transposable element amplification

Introduction

Over the last decades, diverse genome engineering strategies manipulating the functioning of transposable elements (TE) have been intended to achieve site-specific integration of foreign DNA into host genomes. With this aim, the naturally occurring mechanism of transposition was exploited in animals to develop corresponding vectors that allow efficient gene transference and effective integration into genome (Szabo et al. 2003; Yant et al. 2007; Palazzoli et al. 2010). Moreover, the zinc-finger (ZF) and transcription activator-like effector (TALE) programmable nucleases, which can be efficiently redesigned to target specific genome sequences, have been also employed for the development of tools to efficiently achieve genome integration. ZF- and TALE-based constructs display important applications in genome engineering, reverse genetics, and targeting transgenic integration strategies (Walsh and Hochedlinger 2013; Bortesi and Fischer 2015; Cox et al. 2015). Finally, one RNA-guided system based on prokaryotic Type II clustered regularly interspaced short palindromic repeats (CRISPR) and CRISPR-associated Cas9 endonuclease (CRISPR/Cas9) immune system has recently emerged as more efficient, easy to use, and low-cost genome editing tool able to produce desired genomic modifications (Cong et al. 2013; Ran et al. 2013; Lai et al. 2016) (Fig. 1).

Communicated by M. Kupiec.

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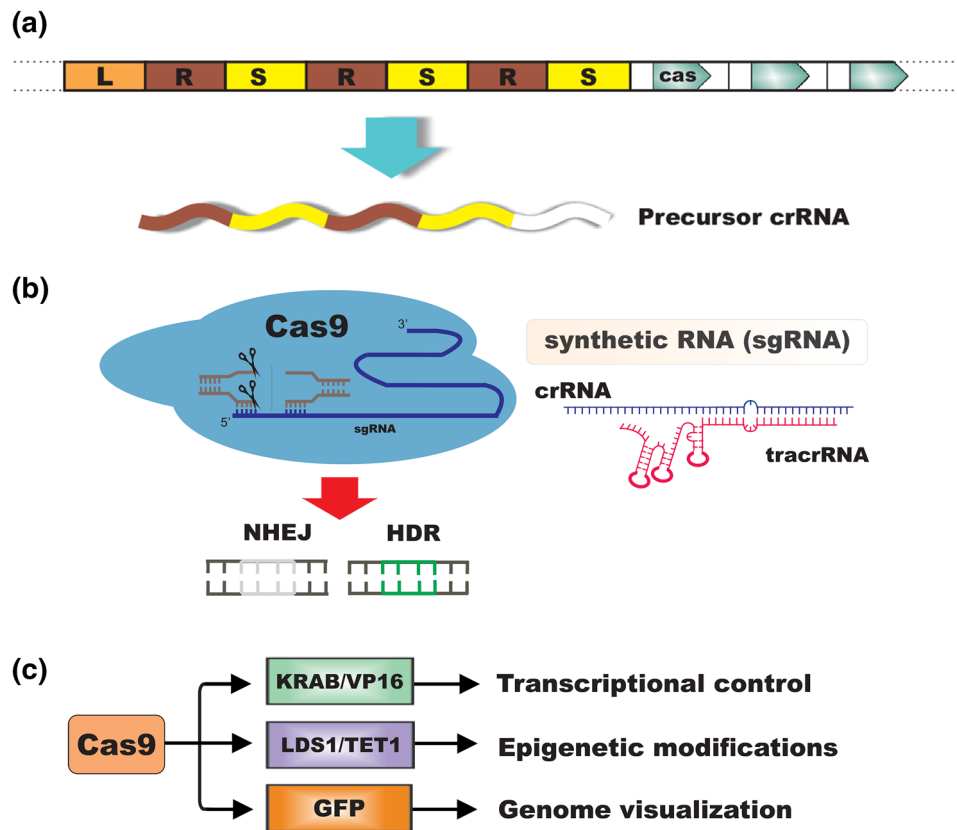


Fig. 1 CRISPR/Cas9 genome editing system. **a** CRISPR locus consists of one leader sequence (L) followed by the CRISPR array containing direct repeats (R) and spacers (S), and CRISPR-associated (Cas) genes. In natural type II CRISPR system, this locus is transcribed as a CRISPR RNA precursor (pre-crRNA) that is processed into mature CRISPR RNAs (crRNAs) molecules which are then bound to transacting CRISPR RNAs (tracrRNAs) to direct targeted Cas9-mediated Double Strand Breaks (DSBs) in complementary crRNA sequences. Thus, the natural CRISPR system is capable of defending the host against infection of matching crRNA pathogenic sequences. **b** In CRISPR/Cas9 genome engineering, a synthetic sin-

gle-guide RNA (sgRNA) that combines the crRNA and tracrRNA is employed to mediate Cas9 DSB cleavage at targeted genomic sites which are subsequently repaired via non-homologous end-joining (NHEJ) or homology-directed (HR) competing cellular DNA repair pathways. By HR-mediated repair, it is possible to achieve targeted integration of desired templates and induce site-specific genome changes. The only requirement of the system is one short Cas9 recognition site known as Protospacer-Adjacent Motif (PAM). **c** Alternative Cas9-based chimeric fusion systems can be also engineered to achieve efficient transcriptional targeting, site-specific epigenetic modifications, cell imaging, etc

The CRISPR/Cas modules are adaptive host defense systems that protect archaea and bacteria against the invasion of viruses and plasmids (Jinek et al. 2012). The CRISPR locus was initially identified in *Escherichia coli* (Ishino et al. 1987) and the CRISPR/Cas modules were further characterized by Jansen et al. (2002), who identified CRISPR and CRISPR-associated (cas) genes in more than 40 species of prokaryotes. CRISPR/Cas systems can be divided into three main types (I, II, and III) and ten subtypes (Makarova et al. 2011). The CRISPR/Cas9 type II genome editing system uses a synthetic single-guide RNA (sgRNA) to produce targeted Cas9 DNA double strand breaks (DSBs) that are subsequently repaired by endogenous host mechanisms (Jinek et al. 2012; Doudna and Charpentier 2014). As a consequence of this, the DSBs generated are rejoined by either non-homologous end-joining (NHEJ) or homologous

recombination (HR) pathways (see Fig. 1b). Depending on the repair mechanism activated, site-specific modifications involving gene disruption, gene replacement, or nucleotide substitution may be effectively generated. NHEJ is prone to produce indel mutations at target sites, while by use of DNA templates harboring desired sequences, it is possible to generate specific genome modifications by activation of HR-mediated repair (Sakuma and Yamamoto 2015). Alternative genome engineering strategies consisting of a nuclease-deactivated Cas9 (dCas9) fused with modular transcriptional domains (activators and/or repressors), chromatin remodelers, and fluorophores also enable efficient transcriptional control, site-specific chromatin modifications, and visualization of loci, respectively (Li et al. 2016; see Fig. 1c). The RNA-guided CRISPR/Cas9 system has dramatically improved our ability to edit the genome of many organisms,

thereby being increasingly employed in biotechnology and therapeutics.

Transposable elements and the CRISPR/Cas9 genome engineering system: control of the host genome

The RNA-guided CRISPR/Cas9 system for design of homing endonuclease gene drives

Diverse endogenous genome elements including, among others, homing endonuclease genes (HEG), and transposable elements (TEs) are capable of exploiting the host machinery to increase the odds of being inherited. Such elements were previously referred to as “gene drives”, or parasitic genome players able to spread in the host genome by copying themselves into target sequences (Burt 2003; Adelman and Tu 2016). Since the beginning, it was believed that these elements might be used to design effective genome engineering strategies that allow coopting endogenous host molecular mechanisms. With this aim, Burt (2003) suggested an interesting, multi-talent, and promising procedure based on engineered HEG drives which would allow addressing the host cell machinery and control gene expression (Fig. 2); however, several technical constraints hindered effective design. Even though Burt’s proposal had not been implemented, his idea predicted the development of future applications. Fortunately, the emergence of the adaptable CRISPR/Cas9 genome editing system has overcome methodological restrictions, thereby Cas9-based drives were already successively engineered to bias inheritance in favor of particular fruit fly (Gantz and Bier 2015) and yeast (DiCarlo et al. 2015; see Fig. 2b) drive constructs which exhibited remarkable high transmission rates (97 and 99%, respectively). Recent genome engineering strategies based on CRISPR/Cas9-mediated gene drives have demonstrated having potential to address the evolution of natural populations (Hammond et al. 2016), whereby displaying the outstanding power that these emerging technologies possess and, besides, underlying an imperative need for public debate before each effective use (Esvelt et al. 2014).

CRISPR spacer acquisition and insertion of transposable elements: themes in common

Several parallels between the CRISPR/Cas9 system and the eukaryotic RNA interference (RNAi) mechanism have been interestingly discovered (Mojica et al. 2005; Bolutin et al. 2005; Barrangou et al. 2015). In nature, CRISPR loci contain arrays of direct repeats (DR) associated with spacers sequences which likely derived from bacteriophages or plasmids (see Fig. 1a). The CRISPR DRs act as

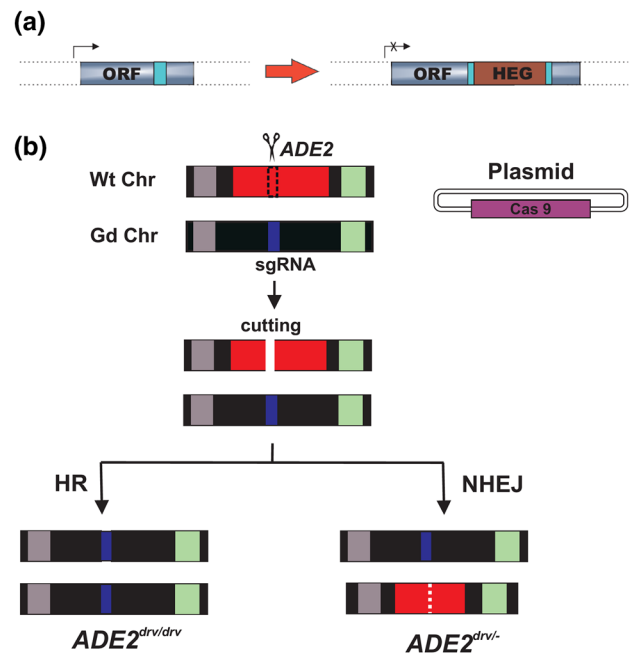


Fig. 2 Endonuclease-based gene drive systems. **a** Burt’s proposal (2003) based on targeted homing endonuclease gene (HEG) drives. Genome engineering HEG drives may spread in the genome host via targeted integration into recognition sites (light blue), thereby activating recombination while simultaneously preventing the cleavage of chromosomes that carry them. Following site-specific HEG insertion, HEG-based gene drives are expected to be able to modify host cellular functions (in this case, causing loss of function by gene disruption). **b** Biased inheritance of a CRISPR/Cas9 gene drive system in *S. cerevisiae* (Di Carlo et al. 2015). Gametes carrying a drive construct targeting the gene encoding the phosphoribosylaminoimidazole carboxylase (ADE2) enzyme mate with wild-type haploid cells and thus generate CRISPR/Cas9-based DSBs which are subsequently repaired by HR (gene replacement) or NHEJ (gene disruption) pathways. CRISPR/Cas9 genome editing can be monitored in the progeny by loss of function of ADE2. Blue box DNA sequence encoding sgRNA (single-guide RNA), *Wt Chr* wild-type chromosome, *Gd Chr* gene drive chromosome, *HR* homologous recombination, *NHEJ* non-homologous end-joining

Cas9-mediated cleavage sites, whereas invasion-acquired spacers mediate immunologic responses to host reinvasion, thereby mimicking the eukaryote post-transcriptional gene silencing (PTGS) pathway mediated by small RNAs (sRNAs). Likewise CRISPR DRs, TEs also are flanked by cleavage sites recognized by particular enzymes (transposases) that mediate their own transposition. Essentially, there are two types of TE flanking sequences referred to as long terminal repeats (LTRs) or terminal inverted repeats (TIRs), belonging to elements which are classified into Class I and Class II, respectively. TEs are categorized according to their transposition intermediates into RNA-mediated (Class I) and DNA-mediated (Class II) elements, which are transposed through “copy-and-paste” and “cut-and-paste” mechanisms, respectively (for review, see Casacuberta

and Santiago 2003). Remarkably, flanking CRISPR DRs derived from insertions of particular Class II TEs known as miniature inverted-repeat transposable elements (MITEs) have been discovered (Mai et al. 2016). MITEs are non-autonomous DNA (Class II) TEs usually composed of ~100–800 pb, flanked by TIRs and adjacent to target site duplications (TSDs) (Fig. 3). Resembling the functioning of CRISPR systems during bacteriophage infection, MITEs also can generate sRNAs that mediate the RNAi silencing pathway during stress responses and environmental (hormone) signals (Yan et al. 2011). Indeed, the existence of MITE-derived CRISPR DRs indicates that site-specific TE insertions have contributed to evolution of CRISPR arrays (Mai et al. 2016). Diverse types of TEs exhibit preference insertional targets, whereby hot spots for many TE families have been correspondingly characterized (Castilho and Casadaban 1991; Feng et al. 2013). In the case of plant MITEs, these elements often exhibit preferential transposition into both AT dinucleotide and ATT trinucleotide genome signatures (Jiang and Wessler 2001; Jiang et al. 2003).

The transposition mechanism by which TEs are scattered into host genome challenged the mistaken concept that considered the genome as one fixed, immutable entity (Mourier 2016). Nowadays, it is well known that transposition comprises an important adaptive mechanism capable of providing both heritable and non-heritable variability, source of critical phenotypic plasticity. Under particular environmental conditions, active transposases mediate transposition and this may positively affect host gene expression by providing promoter sequences (Jordan et al. 2003; Feschotte

2008), modifying gene expression patterns (Cowley and Oakey 2013; Kim et al. 2015), changing local chromatin structure (Hollister and Gaut 2009), etc., thereby potentially resulting in adaptive phenotypes. For example, Class I retrotransposons sequences are activated during human neuronal differentiation and consequent amplification triggers chromosome arrangements capable of conferring somatic plasticity (Singer et al. 2010; Baillie et al. 2011). In maize, Qüesta and colleagues (2010) proposed that UV-B-radiation induces the transposition via modulation of chromatin structure and thus generates variation in the genome. Moreover, in *Arabidopsis*, it has been showed that stress heat conditions induce the transposition of the ONSEN (Class I) copia-like retrotransposon and TE accumulation was encouraged in plant small interference RNA (siRNA) deficient mutants (Matsunaga et al. 2012), whereby evidencing the critical role of RNAi during stress-mediated transposition. The eukaryotic RNAi machinery functions to epigenetically regulate the insertion of TEs by sequence-specific mechanisms and thus maintain genome stability (Buchon and Vaury 2006). Indeed, TE mobilization and epigenetic gene regulation mechanisms are deeply interconnected to each other, and nowadays, it is well known that transposition may trigger epigenetic modifications associated with more adaptable stress-tolerant phenotypes (Yang et al. 2005; Hou et al. 2012; Castelletti et al. 2014).

Control of host transcriptional activity by TE-based drives

The targeted insertional mutagenesis has emerged as an important strategy for deciphering the gene function by inducing large-scale mutations into genomic loci of interest. In cancer modeling, DNA Sleeping Beauty TE-based systems have been used to induce somatic-specific mutagenesis and, therefore, to identify essential genes involved in tumorigenesis (Molyneux et al. 2014). In plants, a transposon-tagging tool for genome-wide analysis based on the rice *mPing* MITEs was used in transgenic soybean (Hancock et al. 2011), showing this system preferential insertion for both nearby genes and AT-rich sequences. Hancock et al. (2011) observed an increased transpositional activity during specific developmental stages (cotyledon vs. globular stage), suggesting interestingly that insights into the developmental regulation pathways involved might be used to control transposition. Both results indicate how the genome fluidity can be efficiently manipulated via TE-based systems. In this sense, new molecular breeding strategies based on the induced activation of Class I retrotransposons have also already been suggested (Paszkowski 2015). TEs represent an important source of phenotype plasticity which is particularly interesting in view of the potentiality for addressing transposition through application of external stimuli. As a

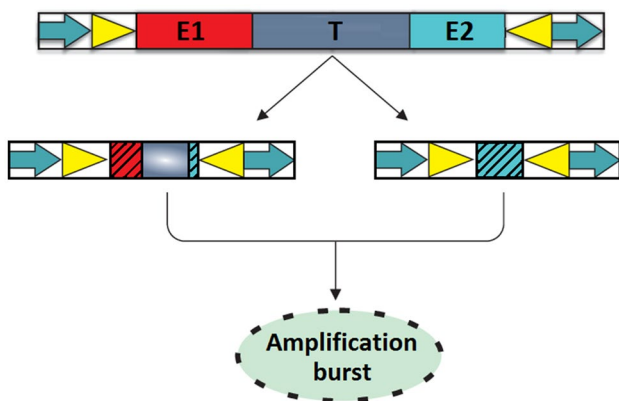


Fig. 3 Miniature inverted-repeat transposable elements (MITEs). MITE elements contain terminal inverted repeats (TIRs, triangles) adjacent to target site duplications (TSDs, arrows). It is believed that MITEs were originated from deletion derivatives of Class II TEs that lost the ability to drive transposition (i.e., transposases and functional enzymes), whereas maintaining the transpositional activity. MITE-derived sequences may not contain perfect TIRs or even lose the TSDs. An amplification burst of MITEs can dramatically increase the number of these elements in one host genome. *E1* and *E2* enzymes that mediate transposition, *T* transposase

consequence of this, we are currently able to engineer TE-based strategies to induce transcriptional control of target genes (Fig. 4). For such purpose, the versatile CRISPR/Cas9 system can be used to mediate HR-based genome editing and thus to introduce transposition insertion sites upstream target genes whose transcriptional regulation is desired. Alternatively, RNA-guided genome engineering constructs based on the use of Cas9 fused with transposase enzymes might be likewise employed to direct the insertion of target sequences. In this regard, targeted transposition using vector delivery systems consisting of the *piggyBac* transposase fused with ZF/TALE nucleases has already been reported (Kettlun et al. 2011; Li et al. 2013; Owens et al. 2013).

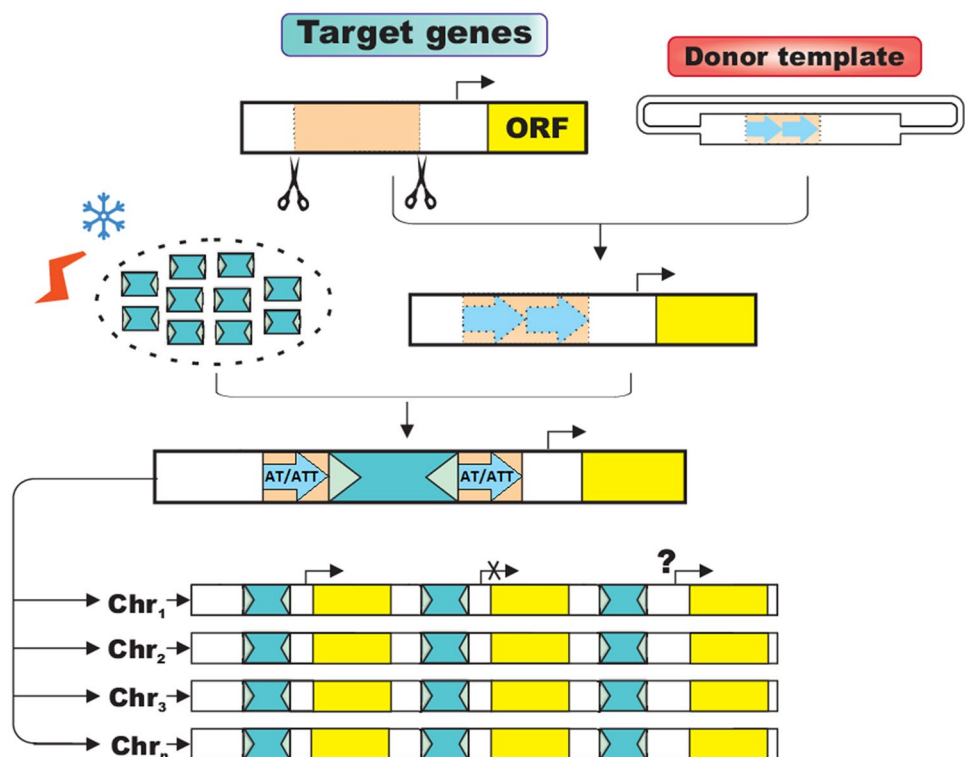
Miniature inverted repeat transposable elements (MITEs) and their interactions with the host genome

To design effective TE-based genome engineering strategies for control of host cell functions, insights into the particular type of TE should be mandatory. In this work, MITE elements are described, since these Class II DNA transposons exhibit a series of characteristics that make them especially suitable for designing TE-based drives. (1) MITEs are abundant repeat elements in eukaryote genomes and they have played critical roles during genome evolution (Lu et al. 2012; Ye et al. 2016). For example, MITE elements represent the most usual type of TEs in rice genome (Jiang et al. 2004). (2) Insertion of MITEs can both upregulate

and downregulate the expression of nearby genes (Chen et al. 2014). (3) Many active MITE families exist, among others, *mPing* and *mGing* in rice (Jiang et al. 2003; Dong et al. 2012), *Stowaway* in potato (Momose et al. 2010), and *AhMITE1* in peanut (Shirasawa et al. 2012). (4) Particular MITE families have been shown to mobilize during cell differentiation under certain conditions (Dong et al. 2012). (5) MITEs are preferentially inserted close to genes (Lu et al. 2012; Wessler et al. 1995). For instance, the Tourist-like and Stowaway-like plant MITE superfamilies show target site preference for integration into specific gene sequences (Jiang et al. 2004). (6) Since MITEs do not possess the enzymes required for their own transposition, they are usually shorter than autonomous DNA TEs (Jiang and Wessler 2001), thereby being more easily manipulated for designing efficient TE-based genome engineering strategies.

Although different families of MITEs have been identified in plants and animals, their origins are still unclear. It has been proposed that MITEs might have derived from ancestral DNA transposons which gave rise to related autonomous elements whose activity eventually resulted in the generation of non-autonomous deletion derivative elements. Subsequently, the amplification of these derivative elements would give rise to a collection of homogeneous non-autonomous TEs, e.g., a MITE subfamily (Feschotte et al. 2002a; see Fig. 3). This idea is supported by the sequence homology between MITEs and DNA transposons; however, most MITEs do not exhibit significant homology with previously

Fig. 4 MITE-based strategy to drive the transcription of target genes. The HR-mediated CRISPR/Cas9 genome editing system can be employed to generate MITE hotspots (in this case consisting of AT/ATT nucleotide signatures). Diverse environmental stimuli are capable of trigger selective amplification and movement of particular MITEs, which are thus accordingly integrated into CRISPR-targeted loci by taking advantage of the preferential insertion at genome-engineered hotspots. Such strategy makes use of the dual essential property of MITEs to either up or downregulate the transcription of nearby genes corresponding to *loci* in which they are hosted. References: open reading frame (ORF), yellow; MITE, light blue



characterized elements beyond the TIR sequences (Feschotte et al. 2002b). Similarly, mechanisms that trigger the temporal activation of MITEs via trans-acting transposases and/or “genome shock” phenomena such as, for example, cell culture or hybridization, remain to be elucidated (Lu et al. 2012).

In silico analyses show that MITE sequences are involved in RNA-based gene regulation, either by hairpin-like miRNA precursors (pre-miRNA) (Lorenzetti et al. 2016) or by siRNA biogenesis (Kuang et al. 2009). Interestingly, it has been recently demonstrated that sRNAs can be used by cell machinery for targeted gene expression control, not only downregulating but also upregulating the transcriptional activity (Li et al. 2006; Turner et al. 2014). Alternatively, insertion of MITEs can mediate regulation of gene expression by promoting the establishment of epigenetic modifications such as, for example, DNA and both histone methylation and acetylation marks. The currently active *mPing* MITEs represent a clear example of how TEs may act synergistically with the host modulating its functioning. The *mPing* family is present in high copies in the rice genome and exhibits preferential insertion into AT-gene-rich regions, avoiding exons while simultaneously choosing promoter regions. These elements are able to both upregulate or downregulate the expression of genes according to the localization of the insertion, although more than 80% of the *mPing* insertions did not exhibit detectable effects on the expression of nearby genes (Naito et al. 2009). Interestingly, Naito and colleagues (2014) proposed that active *mPing* elements should be benign to their hosts, because the consequent *mPing* amplification resulted in selective gene expression control, which is particularly useful in view of manipulating transcription of target genes. In crops, MITEs can modulate the transcriptional activity of essential genes and, therefore, MITE insertion represents an important source of genetic variation worthy to be considered for improvement of agronomic traits (Zhang et al. 2000; Patel et al. 2004; Yang et al. 2005; Li et al. 2014; Mao et al. 2015; Vaschetto 2016).

Concluding remarks

Most eukaryotic genomes are littered with TEs, and nowadays, it is well known that these elements have played critical roles in genome evolution (Casacuberta and Santiago 2003; Garfinkel et al. 2016). Although their mode of action resembles to selfish parasitic elements, TE dispersal may also be beneficial for hosts. Recent insights into the biology of TEs retake an initial position which postulated that environment-induced TE dispersion represent an important adaptive mechanism (McClintock 1984), and nowadays, it is recognized as a regulatory mechanism for both allocating

gene motifs and genome sequences involved in the establishment of epigenetic profiles. It is worthy of note that amplification of TEs induced under certain conditions can offer significant benefits to the host by modulating expression of non-linked genes involved in the same gene regulatory pathways, even having potential for generating de novo regulatory networks (Feschotte 2008).

Transposon-based vectors are very useful and effective tools for achieving targeted mutagenesis (gene disruption/replacement). The emergence of versatile CRISPR/Cas9 genome editing system has enabled to overcome long-standing technical limitations to give the next step towards the design of more efficient TE-based targeting systems. Moreover, the preferential integration induced through application of particular stress/environmental treatments represents a key milestone to maximize the versatility of such systems, since it allows the exploitation/use at will of the natural potential that TEs have to control the host. As it is stated before, MITEs exhibit a series of well-defined characteristics that make them an appropriate tool for engineering of TE-based systems. In particular, the *mPing* MITE family from rice genome have shown to be useful for achieving this objective, since the movement of *mPing* elements can be activated by application of different treatments such as cell culture (Jiang et al. 2003; Kikuchi et al. 2003), hybridization (Shan et al. 2005), or pressurization (Lin et al. 2006). In addition, since it has been demonstrated that transgenic *mPing*-based transposon-tagging systems remain active in the soybean genome (Hancock et al. 2011), the functional characterization of *mPing* family would also allow designing effective genome engineering strategies involving not only endogenous but also exogenous control of the host cell machinery.

Diverse host functioning mechanisms are controlled by TEs, while efficient CRISPR/Cas9 technologies can be used to deploy the potential that these elements have to influence cellular activity of the host. In crops, the climatic fluctuations associated with global warming require the design of molecular breeding strategies tailored to optimize the plant development under such scenarios of change (Hou et al. 2012); thereby, it has already been suggested that TE amplification may represent a solution for generating functional genetic diversity in the face of ever-changing environments (Naito et al. 2009). Moreover, nowadays, we are able to engineer CRISPR/Cas9-targeted loci to efficiently achieve site-specific TE integration, thereby allowing regulation of target genes by exploiting intrinsic TE capabilities for controlling transcription. In addition, since it is well known that TEs are associated with control of different host genome mechanisms including telomere maintenance, chromosomal rearrangements, gene duplication, and epigenetic regulation, analogous strategies can be eventually developed to manipulate such important functions. Consequently, the emergence

of versatile CRISPR/Cas9 genome editing tool raises TE-based strategies to endogenously coopt the host mechanisms, being therefore undoubtedly beneficial for the development of diverse biological applications.

Acknowledgements The author would like to thank Marcela Rivarola and Cristian Arellano for grammatical corrections. Also, I wish to thank Jimena Ingaramo who provided support in the preparation of the figures.

Compliance with ethical standards

Conflict of interest The author declares that has no conflict of interest.

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