Moving through the Stressed Genome: Emerging Regulatory Roles for Transposons in Plant Stress Response

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The recognition of a positive correlation between organism genome size with its transposable element (TE) content, represents a key discovery of the field of genome biology. Considerable evidence accumulated since then suggests the involvement of TEs in genome structure, evolution and function. The global genome reorganization brought about by transposon activity might play an adaptive/regulatory role in the host response to environmental challenges, reminiscent of McClintock's original ‘Controlling Element’ hypothesis. This regulatory aspect of TEs is also garnering support in light of the recent evidences, which project TEs as “distributed genomic control modules.” According to this view, TEs are capable of actively reprogramming host genes circuits and ultimately fine-tuning the host response to specific environmental stimuli. Moreover, the stress-induced changes in epigenetic status of TE activity may allow TEs to propagate their stress responsive elements to host genes; the resulting genome fluidity can permit phenotypic plasticity and adaptation to stress. Given their predominating presence in the plant genomes, nested organization in the genic regions and potential regulatory role in stress response, TEs hold unexplored potential for crop improvement programs. This review intends to present the current information about the roles played by TEs in plant genome organization, evolution, and function and highlight the regulatory mechanisms in plant stress responses. We will also briefly discuss the connection between TE activity, host epigenetic response and phenotypic plasticity as a critical link for traversing the translational bridge from a purely basic study of TEs, to the applied field of stress adaptation and crop improvement.

Keywords: transposable elements, plant stress tolerance, epigenetic, regulatory role, crop improvement

INTRODUCTION

The solution of the C-value paradox, a well-known enigma dating from the earliest days of genome biology, led to the recognition of repetitive DNA, while simultaneously raising questions regarding its nature and possible function (Ohno, 1972; Doolittle and Sapienza, 1980; Orgel and Crick, 1980). Within the repetitive DNA, which constitutes the bulk of most eukaryotic genomes, transposable elements account for the major part. Furthermore, there exists a positive correlation of TE content with the organism genome size, though the individual TE classes vary as regards their presence
in the respective genome (El Baidouri et al., 2014). TEs have been widely regarded as genomic parasites, with a tendency to self-perpetuate at the expense of host genomic stability (Doolittle and Sapienza, 1980; Orgel and Crick, 1980). However, systematic analysis across different plant systems has revealed their potential role in regulating host gene expression and genomic rearrangements (Lisch, 2012). The prevailing view considers the evolution of epigenetic processes as a defense against TEs and viruses, and TE involvement in any essential and adaptive host functions as matter of mutual coadaptation on part of the host and the TE (Kidwell and Lisch, 1997; Slotkin and Martienssen, 2007). However, this viewpoint is now being challenged in favor of TEs as the agents of evolutionary innovation and phenotypic plasticity (Fedoroff, 2012).

Initial clues regarding the nature of transposable elements came from a series of breeding experiments in maize, centered on a breakage region associated with variegated kernel color (McClintock, 1950). Unable to map these regions, McClintock surmised that these elements were in fact “mobile,” contributing to host genome dynamics. The Ac/Ds system thus opened a whole new dimension in the field of genetics, challenging the idea of a static genome with a more dynamic outlook. Since then, the overwhelming evidence accumulated via a series of diverse studies has recognized these selfish DNA as useful parasites instead (Magalhaes et al., 2007; Hilbricht et al., 2008; Hayashi and Yoshida, 2009; Studer et al., 2011; Butelli et al., 2012; Guan et al., 2014; Li et al., 2014; Mao et al., 2015). As the evidence of their involvement in host genome evolution and function grows, one part of their role continues to be highlighted, i.e., in plant stress response to environmental challenges (Johns et al., 1985; Hirochika, 1993; Hirochika et al., 1996; Grandbastien et al., 1997; Bui and Grandbastien, 2012). Although TE activation under stress is considered a consequence of epigenetic deregulation (Slotkin and Martienssen, 2007), certain studies have hinted at a regulatory role for TEs, echoing the original controlling element hypothesis (McClintock, 1984; Bui and Grandbastien, 2012; Lisch, 2012). In this review, we attempt to explore these regulatory themes and associated applied aspects.

**TRANSPOSABLE ELEMENTS (TEs): CLASSIFICATION AND ORGANIZATION IN HOST GENOME**

In general, TE classification is based on type of chromosomal movement: Conservative (cut and paste) or Replicative (copy and paste) and the nature of the transposing unit. Though attempts have been made to devise a unified system of classification which tries to combine both the phylogenetic and enzymatic aspects (Figure 1A, Wicker et al., 2007; Kejnovsky et al., 2012), classification becomes rather difficult in the lower orders and most likely will require rigorous analysis of the aforementioned aspects (Piégu et al., 2015). The first class, known as retrotransposons, consists of an RNA molecule as the transposing unit which encodes proteins structurally homologous to retroviral gag- and pol-encoded proteins, facilitating their reverse transcription and subsequent integration (Kejnovsky et al., 2012). Retrotransposons are the predominant TEs in larger plant genomes such as maize, wheat and sugarcane and are further divided into those flanked by long terminal repeat (LTR) and those devoid of them (Figure 1A). The LTR retroelements are further divided into two major groups: Ty1-copia (Pseudoviridae) and Ty3-gypsy (Metaviridae), both of which are widely distributed across angiosperms. The Class II elements, on the other hand, transpose via a DNA intermediate and possess terminal inverted repeats (TIRs), which serve as sites of excision and reintegration by an element-encoded transposase (Finnegan, 1989). Non-classical transposons include helitrons, which transpose via a rolling-circle mechanism with the aid of a RepHel (Replicate-Helicase) protein with both ligation initiator and DNA helicase domains, plus occasionally a replication protein A like single-stranded DNA-binding protein (Kapitonov and Jurka, 2001; Lal et al., 2003). So far their insertion has been reported in maize, where it accounts for about 2% of the genome (Kapitonov and Jurka, 2001; Lal et al., 2003; Gupta et al., 2005), although these have been computationally predicted in *Arabidopsis*, Rice (Kapitonov and Jurka, 2001; Xiong et al., 2014), as well as other plant genomes, such as *Sorghum*, *rapeseed, Medicago*, and solanaceous crops (Xiong et al., 2014). A third group is the Penelope like Elements, which are considered sufficiently distinct from the two major TE classes (Gladyshiev and Arkhipova, 2007) and have been found recently in conifers (Lin et al., 2016). Another non-canonical TE family, the DIRS-like elements (named after *Dictyostelium* intermediate repeat sequence), have still not been described in higher plants (Kejnovsky et al., 2012). Apart from the autonomous TEs, non-autonomous TEs such as TRIMs (Terminal-Repeat Retrotransposons In Miniature) and LARDs (Large Retrotransposon Derivative) have also been discovered (Witte et al., 2001; Jiang et al., 2002) which owe their mobility to trans-complementation by their autonomous counterparts.

The whole genomic TE complement, also referred to as the mobilome, constitutes a hefty chunk of plant genomes. Retrotransposons, in particular, can be several MBs in size and are known to be present in very high copy numbers, for example, the barley BARE-1, and Maize Opie-1, Cinful-1 and Huck2 reach up to 20,000–200,000 copies (SanMiguel et al., 1996; Vicent et al., 1999; Meyers et al., 2001), a reflection of their replicative mechanism of transposition. In what can be considered a direct answer to C-value paradox, it is now recognized that the respective changes in transposon content, especially retrotransposons, constitute a major source of genome size variation (Figure 1B). This is particularly supported by insights from cereal crops such as maize and sugarcane, where they account for approximately 40–75% of their genome respectively (Schnable et al., 2009; de Setta et al., 2014). DNA transposons generally form a relatively small portion of mobilome in higher plant genomes. Noticeable exceptions are MITEs such as the rice mPing (Jiang et al., 2003; Nakazaki et al., 2003; Naito et al., 2009) and the maize mPIF element, whose copy number reaches upto, 6000 (Zhang et al., 2001). In case of rice, MITEs, such as Stowaway like elements account for almost 2% of the genome (Mao et al., 2000) Despite this, the small size of MITEs (200–500 kb) mean that their contribution to plant genome
FIGURE 1 | (A) General Classification of currently known plant transposable elements. According to the broadest classification system, TEs are divided into two classes on the basis of respective transposing unit: retrotransposons (class 1) and DNA transposons (class 2). Penelope like elements form a distinct group (Gladyshev and Arkhipova, 2007) and have been described only recently in conifers. Further classification is based on the classical /non-classical mode of transposition, and the presence /absence of LTR, TIR sequences, though the phylogenetic origin of TEs is quickly gaining prominence as a key criterion for TE systematic (Modified from Wicker et al., 2007). Due to the lack of sufficient information about Helitron transposition, non-autonomous elements have not been indicated here. The presence of certain non-classical TEs such as Mavericks/Polintons have not been conclusively shown in plant systems yet and is thus (Continued)
size is relatively minor (Lee and Kim, 2014). The non-LTR retrotransposons, also forma substantial part of certain plant genomes (Casacuberta and Santiago, 2003), examples being TS, a SINE with over 50,000 copies in tobacco (Yoshioka et al., 1993) and Del2, a LINE with upto 250,000 copies in Lilium (Leeton and Smyth, 1993). Despite the lack of autonomy, certain LARDs such as Dasheng have upto, 1000 copies in the maize genome (Jiang et al., 2002).

Another key aspect of transposon genomics is the organization of TEs within the genome. Retrotransposons, particularly members of Ty3/gypsy and Ty1/copia super families are known to be present in high copies in centromeric heterochromatic regions with relatively lower rates of recombination (Arabidopsis Genome Initiative, 2000; He and Dooner, 2009), but are also associated with genes (White et al., 1994). Insights from barley genome showed that the major chunk of mobilome and other repeat structures is situated in random BACs, but less so in the gene bearing BACs (International Barley Genome Sequencing Consortium, 2012). As with rice genome, MITE’s organization differs from that of LTR retrotransposons in that these are preferentially located within euchromatic regions as nested insertions (Zhang et al., 2000; Jiang and Wessler, 2001; International Rice Genome Consortium, 2005). In maize some of these insertions produced novel transcripts through shuffling of different host exons, paving the way for the evolution of novel proteins (Morgante et al., 2005; Du et al., 2009).

**TEs AS FUNCTIONAL MOBILE TRACES IN HOST GENOME**

As per McClintock’s original genome shock hypothesis, TEs when triggered can bring about large-scale chromosomal rearrangements which might collectively shape the stressed host genome and facilitate adaptive evolution, as demonstrated in cereal genomes (McClintock, 1984; SanMiguel and Bennetzen, 1998; Vicent and Schulman, 2002; Du et al., 2009; Yu et al., 2011; Barbaglia et al., 2012; Ben-David et al., 2013) as well as eudicot genomes such as tomato (Bolger et al., 2014). The extent of their pervasion is demonstrated by the fact that TEs can directly control plant gene expression through interruption of regulatory motifs or by insertion of new regulatory circuits (Feschotte, 2008; Bonchev and Parisod, 2013). Evidence accumulated till date shows that TE insertion can function at multiple levels to regulate host expression and bring about interesting phenotypes, such as abiotic stress tolerance, fruit color, and so on (Lisch, 2012).

While TE activity appears to be fairly stable in most experimental populations, certain genomic, and physiological stresses, such as regulatory derepression through mutations in regulatory components, hybridization, tissue culture, and abiotic and biotic stresses can lead to a hike in the transposon activity (Hirochika, 1993; Hirochika et al., 1996; Petit et al., 2010). These situations reflect the transient TE-induced spikes in mutation rates in natural populations such as barley (Hordeum spontaneum) and in some cultivars of rice (Oryza spp.) which have been reported as a common occurrence (Kalander et al., 2000; Naito et al., 2006; Ungerer et al., 2006). Considering the fact that artificial populations experience none of the vast array of biotic and abiotic stress that natural population are faced with regularly, these TE bursts and associated rate of null mutations are more likely to be accurate (Lisch, 2012).

**TEs Are Potent Mutagens of the Plant Genomes**

TEs are potent mutagens and hence the most direct consequence of TE activity is the generation of loss-of-function mutants, exemplified by the famous wrinkled pea phenotype, resulting from the insertion of an Ac/Ds TE in the starch branching enzyme SBEI (Bhattacharyya et al., 1990). The TE-induced mutations are similar to other null mutations but for the rate at which they arise, being subject to the activity and copy number of the TE in question (Lisch, 2012). Apart from disrupting the function of host genes, TEs can also disrupt positive or negative regulatory regions, to bring about interesting new twists in host gene expression. This is exemplified by the maize Vgt1, a conserved non-coding sequence (CNS) CNS present roughly 70 kb upstream of a gene that encodes an AP2 transcription factor which is a negative regulator of flowering (Salvi et al., 2002). An insertion of a MITE into the conserved portion of Vgt1 is tightly associated with flowering time variation in maize (Salvi et al., 2002). Similarly, in rapeseed, BrFT2, an ortholog of the Arabidopsis FLOWERING LOCUS T (FT) gene is involved in regulation of flowering time (Zhang et al., 2015). A loss of function allele was generated via insertion of a retrotransposon in the second intron of BrFT2 in one of the parental lines, disrupting its expression The RILs carrying only the mutated BrFT2 allele showed delayed flowering regardless of growing seasons when compared to RILs carrying the wild-type BrFT2 allele. The consequence of TE insertion in regulatory regions like enhancers or repressors, is also exemplified by the insertion of Mutator elements into a CNS present in the first intron of the knotted1 gene in maize which leads to its ectopic expression (Salvi et al., 2007). Additional examples include the certain horticultural traits such as seedless apples (Yao et al., 2001) and flower-color variation in morning glory (Park et al., 2007).

**TEs Can Modulate Native Gene Expression Patterns**

While TE insertion can functionally impair the host gene, regulation may also come in the form of upregulation/repression of the gene expression profile or altered tissue specificity. A classic
example is the maize tb1, which encodes a transcription factor which represses branching (Studer et al., 2011). The presence of a retrotransposon, *Hopscotch*, present as far as 60 kb upstream of the tb1 locus, greatly enhances its expression resulting in a dramatic reduction in the number of branches relative to the progenitor species. This example clearly demonstrates how TE-induced changes in gene expression, are literally “far-reaching” and have been a shaping influence in the evolutionary domestication of maize (Studer et al., 2011; Lisch, 2012). The alteration in tissue specificity due to TE activity is exemplified by the pigmentation alleles in maize. The insertion of an active retroelement in the first exon of the maize b1 gene shifted the expression pattern from vegetative tissues to the seed, creating the B-Peru allele (Selinger and Chandler, 2001). Further transposon insertion in this locus, created the B-Bolivia allele, which further reduced the expression of b1 to yield a more variegated phenotype.

**TEs Can Introduce New Genetic Information**

Additionally, TEs can also bring some novelty to the genome architecture to introduce new coding information. Many Transposon families (such as *Mutator*, Ac, and mPing) tend to insert into the 5′ region of genes, which, combined with the elemental information TEs carry, can give rise to interesting new patterns of gene expression (Pan et al., 2005; Naito et al., 2009; Vollbrecht et al., 2010; Lisch, 2012). In rice, a recent burst of mPing activity was detected (Naito et al., 2009). The comprehensive sequence analysis of 1664 mPing insertion sites in rice genome and their impact on the expression profile of 710 genes revealed a surprising avoidance of exon insertions and a preference for insertion into 5′ flanking regions instead. Moreover, an increase in expression was observed in case of 156 genes. Thus, TEs act as a natural reservoir of regulatory information for host gene-expression networks.

**TEs Can Move Host Genes**

Gene movement is another mechanism by which transposons exert control over the host gene expression, as it transports host genes in an entirely new setting of regulatory regions, methylation backgrounds, and chromatin landscapes that might cause neofunctionalization or silencing, depending upon the altered genomic context (Lisch, 2012). This may constitute another mechanism by which TEs can act as agents of evolutionary revolution (Fedoroff, 2012; Lisch, 2012). The visibly phenotypic traits again provide an example, in this case the oval shape of Roma tomatoes (van der Knaap et al., 2004). Changes in fruit shapes between some varieties of *Solanum lycopersicum* and its round-fruited wild relative *Solanum pimpinellifolium* are largely caused by the variation at the sun locus. The movement of *IQ domain 12 (IQD12)*-the key gene at the sun locus to *DEFL1*, a fruit specific gene causes *IQD12* to be expressed in fruit now, bringing about the round shape fruit which is typical of the Roma variety (van der Knaap et al., 2004).

**TEs Affect Epigenetic Makeup of Host Loci**

So far the prevailing view on epigenetic processes and TE amplification has been that the extensive epigenetic arsenal has evolved in plant systems to suppress TE bursts and such the TE activity observed under stress conditions is due to epigenetic deregulation instead (Slotkin and Martienssen, 2007). An alternate view which is now gaining prominence highlights TEs as the mediators of epigenetic responses instead (McClelland, 1984; Miroouze and Paszkowski, 2011; Fedoroff, 2012). Thus, along with the co-transcriptional regulation, the TE-facilitated post-transcriptional regulation of expression of neighboring gene, can effect far-reaching phenotypic outcomes. This is exemplified by the *FLOWERING WAGENINGEN* (FWA) locus in *A. thaliana* (Kinoshita et al., 2007). The presence of a SINE element immediately upstream of the FWA locus causes it to be epigenetically silenced in vegetative tissues by small RNA processing and DNA methylation. In mutants where the small RNA processing or DNA methylation can result in ectopic expression of FWA, a late flowering phenotype results, which manifests itself as a stable epimutation. In addition, allelic variation in *FLOWERING LOCUS C* (FLC), a central repressor of flowering, contributes to differences in flowering behavior among natural *Arabidopsis* accessions. The insertion of a 1.19 kb non-autonomous Mutator-like TE in first intron of FLC locus renders FLC-Ler (Landsberg Erecta) to siRNA mediated repressive chromatin modifications, reminiscent of the “controlling Element” hypothesis (McClelland, 1950; Liu et al., 2004). The consequent attenuation of FLC expression causes delayed flowering, which is speculated to provide a fitness advantage as the extension of vegetative development can lead to more robust plants with high seed yield (Strange et al., 2011).

**Exaptation of Transposon Sequences into Host Genes**

From an evolutionary angle, the co-option of adaptive features naturally selected for one role, for a new role is called exaptation. Concurrent with this viewpoint, TE sequences, such as exons and binding sites can be directly exapted for specific phenotypic functions in the organism (Hoen and Bureau, 2012). There are multiple examples of TE sequence exaptation in animal systems (Volff, 2006). In plants fewer such examples exist, yet they are critical to plant growth and development. These include FAR1 and FHY3, two transcription factors involved in phy-A signaling, domesticated from the maize TE Jittery (Lin et al., 2007), Mustang(MUGI-8) derived from the MULE superfamily which plays an important role in flower development and reproductive fitness (Cowan et al., 2005; Joly-Lopez et al., 2012), the *Arabidopsis* DAY SLEEPER, important in plant body development (Bundock and Hooykaas, 2005) and the barleyGaryand sugarcane SchAT, members of hAT transposase-like gene family (Muehlbauer et al., 2006; Sinzelle et al., 2009; de Jesus et al., 2012).

**Transduplication: Host Gene Capture by Moving TEs**

Further mechanisms by which transposons can modulate the host transcriptional machinery are transduplication, which has
the potential to create new genes. This mechanism holds special significance in case of maize, where 2791 gene fragments are known to be captured by Helitron elements in the past (Lal and Hannah, 2005; Du et al., 2009; Dong et al., 2011). While many of these gene fragments might be non-functional pseudogenes, some of these may be expressed, generating potential for neofunctionalization. In addition, Pack-MULE has been known to acquire host genes on frequent basis (Jiang et al., 2004; Juretic et al., 2005). As sufficient data on the evolutionary selection of these new gene is not available, further analysis of such gene capture events would be required to establish the functionality of these new genes unambiguously (Lisch, 2012).

**TWO TO TANGO: PLANT STRESS AND TRANPOSONS**

Given the sessile habit of plant systems, the phenomenon of environmental stress, and the corresponding response holds a special significance. As stress impacts the plant at various levels, the plant response to it also manifests at multiple levels of organization (molecular, tissue, anatomical, and morphological), such as the adjustment of the membrane system, alterations in cell wall architecture, transcriptional, and metabolic response (Shinozaki et al., 2003; Hirayama and Shinozaki, 2010). Through the history of evolution, plants have amassed an impressive array of sophisticated response machinery: sensors, signaling components, regulatory components, transcription factors, and effectors that function to “sense, respond, and adapt”- working in an orchestrated fashion to bring about a multifarious phenotypic response (Shinozaki et al., 2003; Hirayama and Shinozaki, 2010). It has also become evident that different stress response pathways intersect at many points and their crosstalk gives way to an intricate network which constitutes not only the immediate component of stress alleviation, but also possibly of memory (Bruce et al., 2007; Walter et al., 2013; Kinoshita and Seki, 2014; Avramova, 2015).

The activation of transposons under plant stress is a well-known phenomenon (Johns et al., 1985; Hirochika, 1993; Hirochika et al., 1996; Turcich et al., 1996; Grandbastien et al., 1997; Bui and Grandbastien, 2012). The first instance of a TEs being induced by stress was the maize Bs1, in response to barley stripe mosaic virus infection (Johns et al., 1985). Since then, several studies have revealed the activation and/or transposition of TEs in response to a diverse array of stresses (Table 1). Abiotic (irradiation, temperature) and biotic (culture tissues or infections by viruses or pathogens) stresses are known to awaken quiescent TEs in plants (Grandbastien et al., 2005; Bouvet et al., 2008). Irradiation is known to induce biological events like mutagenesis and genomic instability. Activation of several retrotransposons has been shown which is variable and subject to radiation type. For example, ion beam, a type of high linear energy transfer (LET) radiation causes relatively few but large, irreparable DNA lesions (Tanaka, 1999). Hence, it is expected that dormant transposons may be induced by ion beam irradiation in rice. UV-B caused activation of cryptic TEs could also increase the mutation rate (Walbot, 1999). Gamma irradiation impact on transposon activation has been also shown in yeast, where it acts on multiple levels in Ty1 life cycle. Ty1 transcription and transposition is repressed in diploid cells and ste12 mutants, where both yet gamma-irradiation affects both an increase in Ty1 RNA levels as well as its transposition (Sacerdot et al., 2005). Interestingly, the stress induced retrotransposon response has been shown to be genotype specific (Ansari et al., 2007; Long et al., 2009; Lopes et al., 2013; Berg et al., 2015).

The global upregulation of transposon activity in response to stress is a well-established fact. The “Controlling Element” hypothesis placed a special emphasis on their potential role as genomic modules controlling host gene expression (Bui and Grandbastien, 2012). In McClintock’s own words “The real point is control. The real secret of all of this is control. It is not transposition.” According to this hypothesis, environmental stress boosts TE expression and transpositional activity, leading to extensive genomic changes that facilitate the adaptation of populations and species facing changing environments (McClintock, 1984). The coevolutionary angle on these transpositional bursts suggested that while such bursts of activity, is self-perpetuating in intent, it might bring about genetic variation on a global scale. For the host, this extensive genome restructuring facilitates stress adaptation and ultimately evolution (Kidwell and Lisch, 1997). However, a more regulatory view of TEs is garnering support in light of recent evidences which establish these as “distributed genomic control modules” (Bui and Grandbastien, 2012; Fedoroff, 2012). The manner in which the TEs participate in this could also be different. The TE occupying heterochromatic regions e.g., the LTR retrotransposons, by virtue of their size, stress responsive promoters, and replicative mechanism of transposition, could sense the physiological or environmental cues affecting alterations in the overall genome architecture (Kidwell and Lisch, 1997). The TEs closely associated with genic regions could be involved in actively reprogramming of host transcriptional networks, affecting specific expression characteristics of individual genes ultimately fine-tuning the host response to specific stimuli (Shapiro, 2005; Lisch, 2012).

**Transposing toward a Regulatory Role**

While aforementioned studies staunchly prove the activation of TE under various environmental challenges, solid evidence for a regulatory role of TEs in host stress adaptation is still scarce. A series of recent studies have pointed out this facet of transposon activity, highlighting their potential role as regulators of host gene expression for host stress amelioration (Magalhaes et al., 2007; Hilbricht et al., 2008; Hayashi and Yoshida, 2009; Bui and Grandbastien, 2012; Butelli et al., 2012; Tsuchiya and Eulgem, 2013; Guan et al., 2014; Li et al., 2014; Mao et al., 2015). Based on the studies certain common mechanistic themes have emerged which explain most of the naturally present transposon modulation of host stress responses (Figure 2).
TABLE 1 | Selected examples demonstrating stress-induced activation of transposable elements under various environmental challenges.

| TE     | Plant system | Tissue/Organ                  | Stress/Environmental stimulus | Transcriptional regulation and/or transposition | Methodology                           | References                  |
|--------|--------------|-------------------------------|--------------------------------|------------------------------------------------|---------------------------------------|-----------------------------|
| MITE   |              |                               |                                |                                                |                                       |                             |
| mPing  | Oryza sativa | Leaves of callus regenerants  | Cell culture                   | Transposition                                  | Transposon display                   | Ngezahayo et al., 2009      |
|        | Oryza sativa | Leaves                       | Inter-specific hybridization    | Transposition                                  | Locus excision display; Southern blotting | Wang et al., 2009           |
|        | Oryza sativa | Leaves of M0 generation       | Gamma Irradiation               | Transposition                                  | Transposon display                   | Nakazaki et al., 2003       |
|        | Oryza sativa | Anther-derived calli          | Anther culture                  | Transposition                                  | Southern Hybridization                | Kikuchi et al., 2003        |
|        | Oryza sativa | Leaves of introgressed lines  | Interspecific hybridization     | Transposition                                  | Locus excision display; Transposon display | Shan et al., 2005           |
|        | Oryza sativa | Leaves                       | Hydrostatic pressurization      | Transposition                                  | Locus excision display; Transposon display | Lin et al., 2006            |
| mGing  | Oryza sativa | M1 generation seedlings, Calus regenerants, Scutellum-derived calli | High dose γ-ray irradiation, anther and scutellum callus culture | Transposition                                  | Transposon display                   | Dong et al., 2012           |
| ZmTPApong | Zea mays    | Anther derived calli         | Cell culture                  | Transcriptionally activated: cell culture and Transposition: regenerated progeny | cDNA-AFLP                         | Barret et al., 2006          |
| Minos  | Triticum aestivum | Leaves                                    | Interspecific hybridization | Transposition                                  | 454 Pyrosequencing, Excision assay, transposon display | Yaakov et al., 2012        |
| DNA TRANSPONOS |             |                               |                                |                                                |                                       |                             |
| TCUP   | Zea mays     | Leaves                        | Long-term tissue culture and aacetylhydine treatment | Transposition, Transcriptional upregulation     | Transposon display; RT-PCR            | Smith et al., 2012           |
| Pong   | Oryza sativa | Leaves of introgressed lines  | Interspecific hybridization     | Transposition                                  | Locus excision display; Transposon display | Shan et al., 2005           |
|        | Oryza sativa | Leaves                       | Hydrostatic pressurization      | Transposition                                  | Locus excision display; Transposon display | Lin et al., 2006            |
| Pong   | Oryza sativa | Leaves of M0 generation       | Gamma Irradiation               | Transposition                                  | Transposon display                   | Nakazaki et al., 2003       |
| Mx, PIF | Zea mays     | Cultured cells                | Cell culture                  | Transcript abundance | EST library                           | Vicent, 2010                |
| Tam3   | Antirrhinum majus | Low temperature            |                                | Transposition                                  | Locus excision display               | Uchiyama et al., 2008       |
| RETROTRANSPOSONS |            |                               |                                |                                                |                                       |                             |
| Athila | Arabidopsis thaliana | Suspension cultures            | Gemivirus proteins treatment   | Transcriptional upregulation                    | Northern blotting, and RT-PCR        | Buchmann et al., 2009       |
| TLC1/Treltyc1 | Solanum chilense | Leaves                             | Prolonged heat stress, wounding, ethylene, auxins, abscisic acid, jasmonate, salicylic acid, H₂O₂ | Transcriptional upregulation | Quantitative PCR; Northern blotting, RT-PCR | Pecinka et al., 2010; Tapia, 2005 |
| Osr7, Tos17 and Osr23 | Oryza sativa | Leaves                       | Interspecific hybridization     | Transposition                                  | Southern blotting                   | Wang et al., 2009           |
| TLC1.1 | Solanum chilense | Leaves                             | Ayclic acid, abscisic acid, methyl jasmonate, hydrogen peroxide and 2,4-D | Transcriptional upregulation | RT-PCR, Promoter/GUS Fusion       | Salazar et al., 2007        |

(Continued)
| TE       | Plant system       | Tissue/Organ          | Stress/Environmental stimulus                              | Transcriptional regulation and/or transposition                  | Methodology                  | References       |
|----------|--------------------|-----------------------|-----------------------------------------------------------|----------------------------------------------------------------|----------------------------|-------------------|
| HRET1    | Hibiscus syriacus  | Leaves                | wounding                                                  | Transcriptional upregulation                                    | Northern Blotting            | Jeung et al., 2005|
| LORE1    | Lotus japonicus    | Progenies of callus-regenerants | Callus culture                                          | Transposition and transcriptional upregulation                   | SSAP, Quantitative PCR       | Madsen et al., 2005|
| Hopscotch-like | Saccharum officinarum | Pollen of callus-regenerants | Callus culture                                             | Transcriptional upregulation                                    | RT-PCR                      | Fukai et al., 2010|
| Morgane  | Triticum aestivum  | Leaves                | fungal infection various biotic/abiotic stresses and nitrogen stress | Presence in EST abundance                                      | EST library                 | de Araujo et al., 2005|
| CCR      | Oryza sativa       | Aerial tissue         | Callus, drought, cold and fungal inoculation             | Increased EST abundance                                          | RT-PCR, Northern blot        | Neumann et al., 2007|
| CIRE1    | Citrus sinensis    | Leaves                | Wounding and auxin treatment                             | Transcriptional upregulation                                    | Semi quantitative PCR         | Rico-Cabanas and Martinez-Izquierdo, 2007|
| Various  | Oryza sativa       | Aerial tissue, suspension culture | Fungal culture, drought and salt stress                  | Transcription upregulation                                      | Microarray                   | Jiao and Deng, 2007|
| Enka, Romani | Triticum aestivum    | Roots                | Fungal inoculation and mycoxin treatment               | Genotype-dependant transcriptional upregulation                  | differential display analysis and RT-PCR | Ansari et al., 2007|
| BAGY1    | Hordeum vulgare    | Leaves                | Seneserase                                                | Transcriptional upregulation                                   | RT-PCR                      | Ay et al., 2008    |
| AtCopeg1 | Arabidopsis thaliana | Whole seeding | Nutrition starvation, salt stress, cytokinin, abscisic acid | Uregulation: nutrition starvation, salt stress and cytokinin, downregulation: abscisic acid | AFLP-TP analysis | Duan et al., 2008    |
| LORE2    | Lotus japonicus    | Leaves                | Cell culture                                             | Transcriptional upregulation and transposition                  | Southern and northern blotting | Fukai et al., 2008|
| Reme1    | Cucumis melo       | Leaves                | UV light                                                 | Transcriptional upregulation                                    | RT-PCR                      | Ramallo et al., 2007|
| CLCo1    | Citrus lemon       | Leaves                | Wounding and salt stress, cell culture                   | Transcriptional upregulation                                    | RT-PCR, Quantitative PCR     | Felice et al., 2009|
| Various  | Vitis vinifera     | Berry fruit           | Post-harvest withering (dehydration)                     | Transcriptional upregulation                                   | AFLP-TP analysis             | Zamboni et al., 2008|
| MERE1    | Medicago truncatula | Cali, callus regenerated plants | Tissue culture                                           | Uregulation: cali and transposition: cali-regenerated plants   | RT- POR, Transposon Display   | Rakocivic et al., 2009|
| Osr23, Osr36, Osr42 | Oryza sativa    | Leaves                | Spaceflight                                              | Genotype-dependent transpositional activation                   | Southern blotting            | Long et al., 2009  |
| FaRE1    | Fragaria x ananassa | Leaves              | Auxins and abscisic acid treatment                       | Transcriptional upregulation                                    | RT-PCR and northern blotting | He et al., 2009    |
| At2G06045 | Arabidopsis thaliana | Whole seeding | Salt, osmotic, cold and heat stress abscisic acid (ABA) treatment | Salt, cold, osmotic stress and ABA: transcriptional upregulation and heat stress: transcriptional downregulation | whole-genome tiling arrays | Zeller et al., 2009 |
| Ttd1a    | Triticum durum     | Salt and light stress | Transcriptional upregulation and transposition           | Transcriptional upregulation and transposition                  | Semi quantitative PCR and SSAP profiling | Woodrow et al., 2010|

(Continued)
| TE | Plant system | Tissue/Organ | Stress/Environmental stimulus | Transcriptional regulation and/or transposition | Methodology | Reference |
|----|--------------|--------------|-------------------------------|-----------------------------------------------|-------------|-----------|
| Various | Lolium perenne | Pseudostem tissue | Pathogenic fungal interaction | Interaction dependant Up- or downregulation | RNA Seq | Eaton et al., 2010 |
| ONSEN/Atcopia78 | Arabidopsis thaliana | Whole plant | Prolonged heat stress | Transcriptional upregulation | cDNA microarray | Pecinka et al., 2010 |
| Callus | Heat stress | Transcriptional upregulation and Transposition | Quantitative PCR and Southern Blotting | Matsunaga et al., 2012 |
| Progeny of stressed mutants | Heat stress | Transcriptional transposition | Northern blotting and transposon display | Ito et al., 2011 |
| Leaves | Elicitation with bacterial flagellin | Transcriptional upregulation | Quantitative PCR | Yu et al., 2013 |
| Various | Arabidopsis thaliana | Whole seedlings | Cold-heat temperature shifts | Transient upregulation | cDNA microarray | Pecinka et al., 2010 |
| Zénit | Zea mays | Seedings | Water deficit | Transcriptional Up- or down-regulation | cDNA microarray | Lu et al., 2011 |
| Various | Oryza sativa | Roots and shoots | Atrazine treatment | Element specific-transcriptional Up or downregulation | RNA Seq | Zhang et al., 2012 |
| Tcs1, Tcs2 | Citrus sinensis | Fruit juice | Cold stress | Transcriptional upregulation | Quantitative PCR | Butelli et al., 2012 |
| Corky | Quercus suber | Leaves | Wounding | Transcriptional upregulation | Quantitative PCR | Rocheta et al., 2012 |
| FIDEI | Arachis spp | Leaves and roots | Infection with C. Personatum and gradual drought stress | Transcriptional upregulation | RNA Seq | Guimaries et al., 2012 |
| Various | Coffea arabica | Leaves | Drought stress | Genotype-specific response | Macroarray | Lopes et al., 2013 |
| ATGP1, EVD | Arabidopsis thaliana | Leaves | Elicitation with bacterial flagellin | Transcriptional upregulation | Quantitative PCR | Yu et al., 2013 |
| Various | Arabidopsis thaliana | Arsenic stress and phosphate starvation | Transcriptional upregulation | Microarray | Castrillo et al., 2013 |
| Various | Oryza sativa | Early microspore stage anthers | Cold stress | transcriptional up or downregulation depending on the genotype | Microarray | Ishiguro et al., 2014 |
| Various retrotransposons | Pinus sylvestris | Heat stress, aphids infestation, and abscisic acid salicic acid treatment. | transcriptional upregulation | iPBS | Voronova et al., 2014 |
| Queinti | Nicotiana tabacum | Cell suspensions, leaves | Fungal elicitation-Cryptogein treatment, ROS and H2O2 | Upregulation byCryptogein, downregulation by ROS and H2O2 | Semi quantitative and Quantitative PCR | Arca et al., 2014 |

**SINES**

- Alu- SINEs | Tritium aestivum | Leaves | Inter-specific hybridization | Transposition | site-specific PCR and transposon display | Ben-David et al., 2013 |
FIGURE 2 | (A) Control of host stress response genes via transposition in regulatory and coding regions. (1) TE insertion into promoter can simply lead to loss of promoter function, keeping with their mutagenic activity, (2) TE insertion can enhance host gene expression or alter gene expression patterns, such as tissue specificity, (3) TE insertion confers stress-responsiveness to host promoter through its own cis-elements, (4) TE insertion may refunctionalise defunct host gene

(Continued)
TE Insertion into Regulatory Regions Facilitates Stress Inducible Gene Expression

Viewed mechanistically, the inherently mobile nature of transposons suggests movement into the genic regions as an obvious mechanism of gene regulation. One of the most striking examples of transposons acting as a regulatory element under stress is the blood orange fruit trait, a consequence of the LTR-driven transcriptional activation of the Ruby Myb gene, an activator of anthocyanin synthesis (Butelli et al., 2012). The Ruby locus is inactive in naval blond oranges, but the insertion of the Tcs1 retrotransposon in its promoter, drives its expression in a cold-dependent fashion in Sicilian blood oranges, whereupon the Tcs1 3' LTR provides transcriptional start and regulation. Furthermore, in some Sicilian blood orange accessions Tcs1 has undergone recombination to form a solo-LTR, sufficient to maintain Ruby expression. Interestingly in Chinese blood orange variety, there exists an upstream insertion of Tsc2, another copy of the same retrotransposon, however in reverse orientation to Ruby, suggesting that regulatory motifs have been preserved either in the U3 or the U5. This study demonstrates how two parallel, yet independent, LTR recruitments influence the host gene expression to perform similar functions, despite the possibly artificial selection of blood orange phenotype (Butelli et al., 2012; Lisch, 2012). It was speculated that the observation of the cold dependence of this desired phenotype led to the artificial implementation of cold in the early selection steps, which could have mobilized this particular retrotransposon (Bui and Grandbastien, 2012; Lisch, 2012).

In rice, the GSTL2 promoter further demonstrates the effect of transposon insertion on host stress responses (Hu et al., 2011). The plant glutathione S-transferase genes play an important in herbicide detoxification. The characterization of its promoter region through deletion analysis revealed that it contains two transposons; a partial Ds-like element and a stowaway-like element resulting in a reduced expression from this promoter, but this reduction was balanced by the presence of enhancer elements (Hu et al., 2011). Moreover, the study suggested that the original promoter of this gene was constitutive in expression, and the cis-elements responsible for its hormone and herbicide responsible expression might have been attributed by the TEs instead (Hu et al., 2011). In rice, the differential transcriptomic analysis of rice seedlings exposed to iron excess revealed a large no. of LTR retrotransposons to be upregulated. While no direct role of transposons in iron excess-responsive gene upregulation could be established, the authors found certain common cis regulatory elements in the promoters of upregulated host genes and LTRs of retrotransposons, which warrants further investigation (Finatto et al., 2015).

TE insertion in other regulatory regions also brings about stress inducible expression of associated genes. A haplotype study conducted in wheat (*Triticum aestivum*) comparing the expression of TaHSP16.9-3A in a heat-tolerant and susceptible cultivar (TAM107, and Chinese Spring respectively), showed its enhanced expression in TAM107 under heat stress (Li et al., 2014). The heightened expression of the TaHSP16.9-3A was found to be due to the presence of a tourist-like MITE into its 3' UTR, which enhanced its transcription. Remarkably, the effect of MITE insertion and the subsequent higher expression was consistent throughout several wheat varieties which possessed the mutant haplotype. Although the 3' UTR is well-known as the determinant of mRNA stability, this study suggested a possible cis-element like role for MITEs in heat stress responsive enhancement of HSP expression (Li et al., 2014).

A key insight in TE activity with host epigenetic processes in context of plant stress is provided by ONSEN, in *Arabidopsis thaliana*. ONSEN is an Copia-type LTR retrotransposon that is transcriptionally activated under heat stress, whose transposition in *Arabidopsis* involves an epigenetic mechanism (Ito et al., 2011; Matsunaga et al., 2012). Transposition of ONSEN is more frequent in the RdDM complex mutants exposed to heat stress, suggesting a role for RdDM machinery in the prevention of transgenerational propagation of ONSEN (Ito et al., 2011). The newly synthesized ONSEN copies decay post stress, suggesting that the stress-induced transcriptional activation of ONSEN may trigger a feedback mechanism that re-establishes methylation and silencing of TEs following recovery from stress. Despite the post-stress decay of newly synthesized transcripts, a burst in transposition was detected in the progeny of the progeny of the RdDM complex mutants. This was attributed to the persisting memory of stress, as a consequence of the faulty epigenetic reponse in these mutants (Ito et al., 2011). An *Arabidopsis* locus with ONSEN insertion in Col-O ecotype was known to be heat responsive (Lim et al., 2006) but the transcriptional response of the same locus in ecotype Zurich, devoid of ONSEN insertion, was considerably low (Ito et al., 2011), suggesting the genotype specificity generally observed in TE responses (Ansari et al., 2014).
Through an artificial system designed to induce experimental bursts of ONSEN in a controlled fashion it was observed that even the loci affected by such stress-induced TE bursts acquired heat responsiveness (Ito et al., 2011). The promoter regions of ONSEN contains a heat shock element (HSE) shown to bind host heat shock transcription factors and permitting its heat inducible expression (Cavrak et al., 2014). This example successfully demonstrates a possible mechanism by which a TE can modulate the host transcriptional circuits in response to stress (Ito et al., 2011; Fedoroff, 2012).

**TE Generated Mutant Alleles in Host Stress Response**

TEs are potent mutagens, and create definite genetic variations at the insertion loci, which could be selectively neutral, fatal, or beneficial for the host. The tight evolutionary pressure governing plant genomes ensures the removal of lethal insertions by failure of survival, the neural and beneficial ones may get selected and fixed in the host genome. The potential role of TEs as regulators of host gene expression is particularly supported by data from the rice genome. Nearly one-sixth of rice genes are associated with retrotransposons (Krom et al., 2008), whereas 58% are associated with a MITE (Lu et al., 2012), suggesting that a large proportion of rice gene promoters appear to contain a TE. For instance, a gene called Pit confers resistance to rice blast disease in the cultivar K29 (Hayashi and Yoshida, 2009). The comparative sequence analysis of the Pit allele between K29 and a susceptible cultivar, Nipponbare revealed that the Pit allele in K29 contains both a DNA transposon dDart, and a long terminal repeat (LTR)-retrotransposon, named Renovator. These results, combined with transgenic studies chiefly attributed the resistance phenotype to the presence of the LTR retroelement (Hayashi and Yoshida, 2009). A similar study was recently made in cucumber, where the locus in question, CsaMLO8, is a candidate powdery mildew susceptibility gene residing in the QTL conferring hypocotyl resistance (Berg et al., 2015). The comparison of alleles of CsaMLO8 from resistant and susceptible genotypes showed the presence of 2 non frameshift deletions of 172 and 74 bp in case of resistant genotypes. These deletions result from aberrant splicing of the CsaMLO8 transcript caused by the presence of a non-autonomous LTR retroelement, leading to the complete and partial loss of the exon 11, which is highly conserved among the susceptible genotype. Similar findings have been reported in pea (Humphry et al., 2011).

Metal tolerance is a root associated trait and a recent study revealed TE insertion as a contributory factor. In sorghum, the principal aluminum tolerance locus in sorghum is AltSB, which encodes three ORFs, out of which only one—a MATE (multidrug and toxic compound extrusion), shows high enough expression in roots where Aluminium quenching occurs (Magalhaes et al., 2007). SbMATE is an aluminum responsive citrate transporter whose coding sequence is the same in both the Aluminium tolerant and susceptible cultivars, the only significant variation being the presence of a Tourist—like MITE which confers root specific expression to this transporter (Magalhaes et al., 2007). In rice, stress activated TEs have been showed to be involved in host response to cadmium toxicity (Ishikawa et al., 2012). Using accelerated carbon atoms, Ishikawa and coworkers recovered three mutants showing drastic reduction in accumulated grain cadmium levels. They mapped these changes to a region housing two putative heavy-metal transporters. Sequence comparison with the WT loci indicated OsNRAMP5 as the center of these changes. While all the three lines experienced mutation in this region, one line—lcd-kmt1, had a 433-bp insertion identical to mPingA1 in the exon X replacing the terminal 32 bp, the remaining insertion being spliced out with intron X. The disruption of this cadmium transporter caused a significant reduction in grain cadmium levels, without much change in the agro-morphological quality characteristics of the parent variety (Ishikawa et al., 2012).

The deleterious effect of TE insertion in host stress genes have also been demonstrated in soybean (Guan et al., 2014). Using a fine mapping approach in the population derived from commercial cultivars Tiefeng 8 and 85–140 (salt-tolerant and salt-sensitive parent respectively), Guan and coworkers identified GmSALT3 (salt tolerance-associated gene on chromosome 3), a major gene encoding an ER-localized cation/H+ exchanger protein that contributes to a salt tolerance phenotype through sodium exclusion in shoots (Guan et al., 2014). They showed that the disruption of the third exon by a 3.78-kb Copia-like retrotransposon, produced a truncated transcript contributed to salt-sensitivity in the cultivar 85–140 (Guan et al., 2014). Further attesting to the capability of TEs to rewire innate transcriptional circuits is the Arabidopsis COPIA-R7. Inserted in the first intron of RPP7 (Resistance to Peronospora parasitica 7), COPIA-R7 introduces an alternative polyadenylation site; the critical balance between the functional and nonfunctional RPP7 then decides the host responsiveness to the pathogen transcripts (Tsuchiya and Eulgem, 2013).

**TE Driven Genetic Polymorphism in Plant Defense**

The plant resistance genes are known to be subject to high levels of polymorphism and are subject to adaptive evolution. For example, the rice Xa21 gene family houses as many as 17 TEs (Song et al., 1997, 1998); even a closely linked marker pTA818 encodes a transposon protein (He et al., 2000). Moreover, some of these insertions have given rise to novel proteins upon insertion. This is demonstrated by Xa21D, which shows 99.9% similarity to the resistance conferring allele, before and after a Retrofit insertion, but encodes a truncated protein lacks the transmembrane or kinase domain present in its progenitor. Furthermore, TE insertion may arise spontaneously in these resistance genes, such as the dLute insertions in L6 flax rust resistance gene (Luck et al., 1998). Their insertion and subsequent reversion or imprecise excision serves as an evolutionary device in the generation of high allelic variability, a fundamental prerequisite for emergence of new pathogen race specificities (Richter and Ronald, 2000). One such example is the evolution of the ALP-A3 gene in diploid wheat (Akhunov et al., 2007). The original progenitor, ALP-A1 specifies an acireductonedioxygenase-like protein, and has undergone duplication, with the new duplicated gene acting as the parent of the next duplication. ALP-A3, while having the complete coding region, has seemingly lost the promoter region, instead
being driven by a promoter sequence originating from a CACTA element (Akhunov et al., 2007).

**Source of Small RNA Regulating Plant Abiotic Stress Response**

Almost 23.5% of all small RNAs identified from rice are derived from MITEs (Lu et al., 2012). TE mediated regulation of plant stress-responsive gene expression also occurs due to the generation of small non-coding RNA that originate from TEs in response to stress. This is exemplified by the generation of a cluster of small RNAs (smRPi1LTR) derived from the Copia95 retrotransposon LTR region under phosphate (Pi) starvation, in *Arabidopsis* (Hsieh et al., 2009). A comparative analysis of the expression pattern between Columbia and Landsberg alleles indicated that smRPi1LTR is a newly evolved small RNA, resulting due to rapid rearrangement of LTR and is an intermediate small RNA species transitioning from siRNAs to microRNAs (miRNAs). The Pi-responsive small RNAs and their target genes are possibly involved in the development or regulation of adaptive response to phosphate starvation (Hsieh et al., 2009). The extent of pervasiveness of TE-derived siRNA in host gene expression is demonstrated by the *Athila6*-derived siRNA854 which shares partial complementarity with the target gene *UPB1b*. The latter is speculated to function in the translational inhibition of TE activity under stress, and *Athila6* might have retained the siRNA854 sequence as a mechanism to silence *UPB1b* instead (Hsieh et al., 2009). The same study estimated the total number of genes targeted by TE derived siRNAs to lie between 20 and 300. In rice, Small RNA siR441 and siR446 are derived from the MITE Stowaway-1 through OsDCI3a. These exhibit a high degree of similarity to a 21 nucleotide sequence present in the 3′ UTR of target stress responsive genes including MAIF1 (Yan et al., 2010).

In maize, critical insights into the role of TEs as determinants of natural variation for stress tolerance, exists in form of a MITE inserted in the promoter of a NAC gene ZmNAC111 (Mao et al., 2015). This example is particularly interesting because this 82-bp MITE insertion in the promoter down regulates its expression, not only though the insertional disruption, but also by acting as the source of 21–24 nt siRNA, which lead to silencing of this locus by the RdDM pathway & H3K9 dimethylation. The ZmNAC111 overexpression maize lines exhibit enhanced drought tolerance at the seedling stage, improves water-use efficiency and upregulation of drought-responsive genes under water stress (Mao et al., 2015). In sugarcane, differential mapping patterns was observed for sRNA and miRNA which showed peaks of RNA mapping in a region downstream to an LTR retrotransposon. Promoters within the 3′ LTR region may be driving expression of this region in an allelic dependent manner (de Setta et al., 2014). Thus TEs, acting through small non-coding RNA can provide a widespread regulatory influence over host gene expression pertaining to stress tolerance.

Despite the predominance of LTR retroelements and MITEs in stress regulation, non-LTRs are not completely far behind. For example, in the resurrection plant *Craterostigma plantagineum* activation tagging identified *Craterostigma* desiccation tolerant (CDT-1), a dehydration-related ABA-inducible gene which confer desiccation tolerance without ABA pre-treatment of the callus (Furini et al., 1997). Related to a family of dehydration responsive retroelements, CDT-1 represents an evolutionary novelty; it is an intron-less multi-copy gene flanked by direct repeats and possesses a very small ORF which is not translated into a protein; instead it encodes a double-stranded 21-bp short interfering RNA (siRNA), which regulate the expression of ABA- and dehydration-responsive genes leading to desiccation tolerance (Hilbricht et al., 2008). Interestingly, the default expression of CDT-1 is only under dehydration stress in vegetative tissues, and upon ABA treatment in the callus. CDT-1 thus serves to illustrate the critical link between environment and genome; CDT-1 is transcribed and transposed under dehydration, and the increasing numbers of such insertions into a stress responsive transcribed locus will eventually lead to a progressively higher level of CDT-1 siRNA, ultimately triggering desiccation tolerance (Hilbricht et al., 2008). This example is particularly important as it demonstrates how TEs are critical in plant evolution and stress response—not only TEs drive stress induced genome dynamics, but also function as the material for evolutionary innovation directed toward plant stress tolerance.

**Recruitment of TE Coding Sequences as Host Stress Response Genes**

While many examples involving TE protein exaptation exist in mammalian and other vertebrate systems (Vollf, 2006; Hoen and Bureau, 2012), those implicated in plant stress response are still scarce. One such example is provided by the Rim2 gene, which is expressed in response to fungal elicitors, implying a role in plant defense (He et al., 2000). Interestingly, the Rim2 protein exhibits considerable sequence homology to the TNP2 transposase of the CACTA TE family. It is present in at least four copies in the modern rice varieties, suggesting that the progenitor element was mobile at some time point prior to becoming inactive (He et al., 2000). Similarly, the *Arabidopsis* AtCopeg1 (Copia evolved gene 1), has evolved from AtCopia95_1, and encodes AtCopia95 polypeptide (Duan et al., 2008). Initially identified as a salt inducible transcript, AtCopeg1 is an intron containing gene, having two alternative 3′ ends with specific expression in leaves and roots. Further, it shows strong fluctuations in response to phosphate (Pi) starvation, nitrate, potassium, or iron starvation. Moreover, external treatment with hormones and hormone analogs, was also discovered to bring about marked changes in its expression, suggesting a possible involvement in the cross talk between various hormone and nutrient stress responsive pathway (Duan et al., 2008).

Despite the existing wealth of information regarding upregulation of TEs under various stresses and environmental challenges, a cause and effect relationship between TE activation and stress adaptation is yet to be established. The aforementioned examples have provided an insight into the possible mechanisms of TE-mediated control of host stress response machinery (Figure 2). Viewed together, it seems likely that the relationship between TE activation and stress response pathways is an
intricately complex one, and it will take extensive evidence to prove TEs as McClintock’s de facto controlling elements.

From Parasitic to Phenotypic: Toward an Applied Front for TEs

The abovementioned studies highlight TEs as the yet untapped reservoirs of genetic variation and phenotypic diversity. The question that arises now is how can TEs be exploited for crop improvement and increasing crop productivity. An interesting and relatively less explored theme is the selectivity in transposition. The rice mPing, an active TE in rice, was shown to be exclusively in single copy regions (Jiang et al., 2003). Furthermore, the comprehensive sequence analysis of, 1664 mPing insertion sites in rice genome and their impact on the expression profile of 710 genes revealed a surprising avoidance of exon insertions and a preference for insertion into 5′ flanking regions instead. The lack of deleterious effects associated with such a TE burst demonstrates how TEs can establish stress-responsive networks (Naito et al., 2009). Tf1, an LTR retrotransposon in Schizosaccharomyces pombe, exhibits a propensity for integration into promoters of stress response genes (Feng et al., 2012). Moreover, only the host genes subject to heat induction themselves, could be activated by Tf1 integration, suggesting a synergy of Tf1 enhancer sequence with the stress response elements of target promoters (Feng et al., 2012). While such a selectivity component in TE activity is in agreement with the proposed regulatory role, the area is still nascent and would require further investigation. A recent review suggested the development of controlled transposition for crop improvement (Paszkowski, 2015). The stress-induced, TE mediated distribution of regulatory elements, can potentially create and fine-tune many regulatory networks responsive to the original stress. More importantly however, the dynamic and possibly heritable restructuring of the host epigenetic landscape can create stress-inducible epiallelic switches. Based on this information, can the deployment of an inducible TE create selective changes to stress responsive QTLs in stress-susceptible cultivars, thus rewiring the transcriptional circuits for tolerance? Can the controlled transposition system be used for creating epigenetic diversity capable of conferring stress tolerance? Such inducible diversity can theoretically be paving the path for phenotypic plasticity—the ability to adjust to seasonal variations so as to maintain homeostasis during development (Rathcke and Lacey, 1985; Paszkowski, 2015). The latter is crucial for the successful adaptation and survival of crop cultivars facing stress. Therefore, controlled transposition in crop plants may potentially generate cultivars with novel variation in response to a particular stress, with implications for sustainable agriculture (Bloomfield et al., 2014).

A second area where studies of TE can directly assist the crop improvement programs is through molecular marker approach. This is especially significant in the case of polyploidy crops with frequent stress-induced genomic rearrangements. The presence of TEs, often close to or within the stress responsive QTLs, especially plant defense genes, along with the traditional attributes of a molecular marker, makes them the markers of choice for diversity studies and trait mapping (Kalendar et al., 2011; Alzhohairy et al., 2014). The mapping of “spikelet-tipped bristles” (stb), which is involved in determining grain number per panicle in foxtail millet, Setaria italica (L.) P. Beauv represents one such example (Sato et al., 2013). Using a combination of transposon display and SSR markers in the F2 population of a cross between a Taiwanese and Japanese landrace, stb1 was successfully mapped to chromosome 2 (Sato et al., 2013). In recognition of their utility in mapping studies and diversity analysis, certain TE based markers, such as IRAP (Inter Retrotransposon Amplification Polymorphism), REMAP (Retrotransposon-microsatellite amplified polymorphism) and SSAP (Sequence Specific Amplification Polymorphism) have been used to some extent in crop improvement programs (Table 2, Manninen et al., 2000; Queen et al., 2004). A more high-throughput application of TE based polymorphism in diversity analysis was shown in soybean for GmCHX1, previously referred to as GmSALT3 (Guan et al., 2014). The same locus was further probed through whole-genome resequencing (WGRS) on 106 diverse soybean lines and three major structural variants and allelic variation were identified (Patil et al., 2016). These SNPs were then utilized for the design of six KASPar (Competitive allele-specific polymerase chain reaction) assays. The haplotype analysis of 104 soybean lines revealed a strong correlation between the genotype and salinity associated phenotype. The high resolution (>91%) and polymorphism (>98%) offered by

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**Table 2: Selected examples demonstrating the application of transposon markers in crop improvement for disease resistance.**

| Marker technique | Transposon element | Plant species | Study | References |
|------------------|--------------------|--------------|-------|------------|
| SSAP             | Tao1, Tao2         | Anacardium occidentale L. | Assessment of polymorphism in segregating F1 progeny of genetic cross for nut size, resistance to anthracnose and black mould | Syed et al., 2005 |
| IRAP and REMAP   | Bagy2, Willma, Sumana, Sabrina, Haignt | Triticum dicoccoides | MAS (Marker assisted selection) for stripe rust resistance | Mandoulakani et al., 2015 |
| IRAP and REMAP   | Reme1              | Cucumis melo L. | Development of markers associated with the melon populations resistant to ZYMV and potential use for introgression in MAS | Mandoulakani and Bernoussi, 2015 |
| IRAP and REMAP   | LTR 6150 d Nikita Carica papaya | Carica papaya | Identification of papaya breeding material resistance to PRSV and potential application for use in Molecular Assisted Breeding (MAB) and MAS | Rashid et al., 2014 |
these markers demonstrate how TE-generated polymorphism can be utilized in crop improvement programs.

The generation of genetic variability in crops with a narrow genetic base, or targeting traits that are not normally present in the crop germplasm, can be accomplished by the use of irradiation technology (Tanaka, 1999). Exposure to irradiation generates mutations on a global scale, which constitutes a genomic shock-like situation (McClintock, 1984). Under such conditions, TEs are mobilized globally creating widespread changes in the host genome, some of these changes might be beneficial and selected through evolutionary sieve to affect heritable changes in the host genome (McClintock, 1984; Ito et al., 2011; Fedoroff, 2012). This rationale serves as the basis of in vitro mutagenesis system for crop improvement, where such beneficial induced mutations are selected and maintained through multiple cycles, a prominent example being the low-cadmium rice (Ishikawa et al., 2012). This system holds special significance for polyploid crops, for which breeding is difficult, such as sugarcane. We have successfully applied the in vitro mutagenesis system to obtain high sugar- (Mirajkar et al., 2016) and salt-tolerant mutants (Nikam et al., 2015). Taking into view that TE mobilization is one of the major events under radiation exposure, further efforts toward can be directed toward enhancing the transposon mobility and transposition efficiency so as to generate heritable mutations with a definite, selectable phenotype.

CONCLUSIONS

Long viewed as “parasites” hazardous to host survival, TEs have since been recognized for their role in plant evolution and stress responses (Bui and Grandbastien, 2012; Fedoroff, 2012; Lisch, 2012). Multiple studies spanning various plant systems, in particular, ONSEN, and mPing, suggest that TEs not only rewire host transcriptional circuits in times of stress, but the extensive genomic rearrangements mediated by such TE bursts shapes genome architecture, ultimately leading to speciation and evolution of plant genomes (McClintock, 1950; Fedoroff, 2012; Lisch, 2012; Bonchev and Parisod, 2013). The host epigenetic processes are thought to have evolved to counter and repress their activity, highlighting their still prevalent image as deleterious to host. However, it is now recognized that under stress, the relaxation of such epigenetic constrains permits TEs to perpetuate their stress-inducible regulatory sequences to hostgenes. TEs also spread their epigenetic signatures to the surrounding new loci, and combined with the generation of regulatory small RNA contribute to epigenetic regulation of host genes as well. As seen in case of ONSEN and mPing, the stress induced transcription and transposition of TEs epigenetic feedback mechanism is at play that re-establishes methylation and silencing of TEs following recovery from stress. This permits a truly dynamic reshaping of the host epigenetic landscape under prevailing environmental stimulus. The recognition of the underlying active regulatory potential of TEs could pave the way for their successful application in crop improvement programs in multiple ways. Clearly, the connection between TEs and host stress response is an exceedingly complex and multifaceted one, and needs further exploration for their potential to be fully realized.

PERSPECTIVES

The ever-growing world population places a proportionately large demand on agriculture for subsistence. Maintaining the crop production is proving to be a hard task in the face of changing climatic scene. The concept of genetic diversity and phenotypic plasticity thus becomes crucial, in the context of developing crops resilient to such climate changes, as the lack of adaptive plasticity can drive a particular cultivar to decline. From a molecular perspective, this phenotypic plasticity is conferred by epigenetic processes, speculated to have evolved to counter viruses and TEs. The latter are known to contribute to the transcriptional and epigenetic landscape of their immediate neighboring regions, and are emerging as active regulators of host stress response. In times of changing environments, the rapid phenotypic plasticity conferred by host epigenetic responses may translate in stress-adaptation. Controlled transposition—the deployment of inducible TE to specific host loci for rewiring transcriptional and epigenetic network scan therefore lead to generation of novel genetic variation and phenotypic plasticity aimed toward stress tolerance. The usefulness of TEs for mapping studies and mutation breeding only adds to their applicability in crop improvement programs. It is being realized that investigation of TE regulation of host transcriptional networks and epigenetic responses is critical for traversing the translational bridge from a purely basic study of TEs to the applied field of stress adaptation and crop productivity.

AUTHOR CONTRIBUTIONS

PN took the lead in writing with inputs from AR during the writing process. PS coordinated the entire work and finalization of manuscript.

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Bui and Grandbastien, 2012; Fedoroff, 2012; Lisch, 2012). Multiple studies spanning various plant systems, in particular, ONSEN, and mPing, suggest that TEs not only rewire host transcriptional circuits in times of stress, but the extensive genomic rearrangements mediated by such TE bursts shapes genome architecture, ultimately leading to speciation and evolution of plant genomes (McClintock, 1950; Fedoroff, 2012; Lisch, 2012; Bonchev and Parisod, 2013). The host epigenetic processes are thought to have evolved to counter and repress their activity, highlighting their still prevalent image as deleterious to host. However, it is now recognized that under stress, the relaxation of such epigenetic constrains permits TEs to perpetuate their stress-inducible regulatory sequences to hostgenes. TEs also spread their epigenetic signatures to the surrounding new loci, and combined with the generation of regulatory small RNA contribute to epigenetic regulation of host genes as well. As seen in case of ONSEN and mPing, the stress induced transcription and transposition of TEs epigenetic feedback mechanism is at play that re-establishes methylation and silencing of TEs following recovery from stress. This permits a truly dynamic reshaping of the host epigenetic landscape under prevailing environmental stimulus. The recognition of the underlying active regulatory potential of TEs could pave the way for their successful application in crop improvement programs in multiple ways. Clearly, the connection between TEs and host stress response is an exceedingly complex and multifaceted one, and needs further exploration for their potential to be fully realized.

PERSPECTIVES

The ever-growing world population places a proportionately large demand on agriculture for subsistence. Maintaining the crop production is proving to be a hard task in the face of changing climatic scene. The concept of genetic diversity and phenotypic plasticity thus becomes crucial, in the context of developing crops resilient to such climate changes, as the lack of adaptive plasticity can drive a particular cultivar to decline. From a molecular perspective, this phenotypic plasticity is conferred by epigenetic processes, speculated to have evolved to counter viruses and TEs. The latter are known to contribute to the transcriptional and epigenetic landscape of their immediate neighboring regions, and are emerging as active regulators of host stress response. In times of changing environments, the rapid phenotypic plasticity conferred by host epigenetic responses may translate in stress-adaptation. Controlled transposition—the deployment of inducible TE to specific host loci for rewiring transcriptional and epigenetic network scan therefore lead to generation of novel genetic variation and phenotypic plasticity aimed toward stress tolerance. The usefulness of TEs for mapping studies and mutation breeding only adds to their applicability in crop improvement programs. It is being realized that investigation of TE regulation of host transcriptional networks and epigenetic responses is critical for traversing the translational bridge from a purely basic study of TEs to the applied field of stress adaptation and crop productivity.

AUTHOR CONTRIBUTIONS

PN took the lead in writing with inputs from AR during the writing process. PS coordinated the entire work and finalization of manuscript.

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Conflict of Interest Statement: The authors declare that the research was conducted in the absence of any commercial or financial relationships that could be construed as a potential conflict of interest.

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