Transposable Elements in Fungi

The availability of fungal genomes has led to the elucidation of the number and distribution of transposable elements (TEs) in several species of fungi. TEs are moderately repetitive DNA sequences dispersed throughout the genome. Such elements can move from one site to another within the genome, and their insertion can produce a broad spectrum of host mutations [1].

Transposable elements are hierarchically classified into class, subclass, order, superfamiliy, family and subfamily. There are two classes of TEs that differ in the presence (class I) or absence (class II) of an RNA intermediate. Class I TEs transpose via an RNA intermediate that is transcribed from a single copy of the genome and produces an RNA intermediate. Class I TEs transpose via an RNA intermediate. Class I TEs transpose via an RNA intermediate, which is encoded by the element itself. Class I have two major subclasses, the LTR (Long Terminal Repeat) retrotransposons and the non-LTR retrotransposons (LINEs, Long Interspersed Nuclear Elements, and SINEs, Short Interspersed Nuclear Elements), which are distinguished mainly by the respective presence or absence of LTRs at their ends. The class II TEs are divided into two subclasses. Subclass 1 are TEs that are transposed by integration and excision mechanisms and subclass 2 are TEs that duplicate before insertion. Subclass 1 contains two orders; the most well-known is the TIR (Terminal Inverted Repeated) order. This order contains nine superfamilies: Tc1-Mariner, Mutator, hAT, Merlin, Transib, P, PIF/Harbinger, CACTA and Crypton. Subclass 2 has two orders: Helitron and Maverick. Furthermore, there are groups of non-autonomous TEs that lack one or more of the genes essential for transposition, including MITEs (Miniature Inverted-repeat Terminal Elements) for class II, SINEs for non-LTR retrotransposons, and TRIM retrotransposons (Terminal-repeat Retrotransposon In Miniature) and LARDs (Large Retrotransposon Derivates) for LTR retrotransposons [2].

Copies or remnants of both class I (retrotransposons) and class II (DNA transposons) have been identified, and the number of distinct mobile elements continues to grow. Such repeated sequences vary in number as well as in size, and comprise 3 to 20% of the sequenced genome of most fungi. In contrast, the plant pathogenic fungus Blumeria graminis has a genome estimated at 174 Mb, and 85% of the genome includes TEs [3]. More than 215 genus-specific TEs were found in Laccaria bicolor in addition to many remaining degenerate copies. Additionally, 40 different TE families were detected in the genome of L. bicolor, and less than 5% nucleotide mutations had accumulated, which suggest that they are recently acquired elements, and possible activities can be inferred for such elements [4]. In general, the most abundant TEs in sequenced genomes of fungi are Gypsy, Copia and Tc1-Mariner (Figure 1).

Since their discovery during the late 1940s by Barbara McClintock [5], the importance of TEs in genome structuring and evolution becomes increasingly clear due to the completion of several genome sequencing projects. Some of the effects of transposable elements may be due to ectopic recombination among elements of the same family. Ectopic recombination occurs between homologous DNA sequences on different chromosomes. An analysis of TEs distributed across the Magnaporthe grisea genome has demonstrated extensive past ectopic recombination events. Thus, ectopic recombination events can aid in fitness because many host-specific genes are in transposable element-rich regions. Therefore, recombination events delete or alter the structure of such genes, which consequently alters their expression [6]. In Coprinus cinereus, the elements are in regions with medium and high recombination rates [7].

Certain species-specific genomic islands in Aspergillus fumigatus include a disproportionate number of TEs compared with syntenic areas, which may be involved in origination or modification of such areas. These areas include genes involved in carbohydrate transport and catabolism, secondary metabolite biosynthesis and detoxification. Lineage specific (LS) genes are a key point in many comparative genomics studies because such regions may be responsible for phenotypic differences between species and reflect
different capacities for niche exploitation [8]. Similarly, numerous (potentially related) multi-copy retrotransposons have been identified in *Phanerochaete chrysosporium* that affect expression of genes related to lignin degradation. Retrotransposons interrupt the gene sequence for putative members of the cytochrome P450 gene family and multicopper oxidase *mco3* genes. An additional element was found 100 bp upstream of the peroxidase gene. The intact coding regions for such elements suggest a recent transposition event [9].

Transposable elements are important for generating fungal variability and increased fitness in the environment. Transposition in response to environmental stress was proposed as an adaptive genomic response. Such stresses may be abiotic (temperature, irradiation and oxidative stress) or biotic, such as pathogen infection. TE activity in fungi under stressful conditions has been demonstrated in *Aspergillus oryzae* [10], *Ophiostoma novo-ulmi* [11], *Moniliophthora perniciosa* [12] and *Magnaporthe oryzae* [13].

Transposable elements in plant pathogenic fungi

Effectors are considered to be small secreted proteins that target and inhibit specific physiological processes involved in plant defense and enhance pathogen virulence without causing a general debilitation of cellular metabolism or cell death [1]. Many effector genes in several plant pathogens have been found embedded in genomic regions with many repeats and TEs [14]. Through studying the avirulence gene of *M. oryzae* AVR- *Pita* gene, Khang et al. [15], Chuma et al. [16] found that such genes are associated with various types of TEs. The activity of such elements and rearrangements caused by recombination can produce deletion, duplication, inversion and translocation. Certain changes may alter *AVR* gene structure or expression, thereby promoting the emergence of novel pathogen races.

Unique sequences involved in pathogenicity were discovered in the *Fusarium oxysporum* genome. Notably, such regions compose 74% of the TEs identified in the *Fusarium* genome, including 95% of the DNA transposons that might be involved in gene duplication events. Of these elements, the *pogo* family is best represented with more than 500 identified copies [17]. The analysis of the *Leptosphaeria maculans* genome revealed the presence of effector genes in TE-rich genome environments that contribute to sequence diversification and probably adaptation to new hosts or better adaptation to the existing host. The presence of effector genes in such dynamic genome environments has a dual advantage for phytopathogens, immediate adaptation to new source of resistance and long-term enhanced ability to duplicate and diversify effector genes to facilitate adaptation to the host or adaptation to new hosts [18,19].

Silencing of transposable elements

Given the potential damage to genome stability caused by transposon movement, many organisms have developed defense systems that suppress TEs activity [20]. Two particular defense mechanisms have received the most attention: repeat induced point mutation (RIP) and RNA silencing [21]. RIP is a gene silencing mechanism that mutates repetitive DNA sequences during the sexual cycle between fertilization and nuclear fusion. RIP prevents accumulation of TEs in fungi. This mechanism may also be related to the gradual divergence of duplicated genes in *Neurospora crassa*, which is essential for the development of new genes with novel functions [22]. In general, RIP induces GC to AT mutations in *Neurospora crassa*, is essential for the development of new genes with novel functions [22]. In general, RIP induces GC to AT mutations in *Neurospora crassa*, which is essential for the development of new genes with novel functions [22]. In general, RIP induces GC to AT mutations in *Neurospora crassa*, which is essential for the development of new genes with novel functions [22]. In general, RIP induces GC to AT mutations in *Neurospora crassa*, which is essential for the development of new genes with novel functions [22]. In general, RIP induces GC to AT mutations in *Neurospora crassa*, which is essential for the development of new genes with novel functions [22].

**Figure 1:** Structure of the major transposable elements found in the genome of fungi. In 1, the class I: LTR-Gypsy (A) and LTR-Copia (B) with their respective coding regions. The pol region contains the PR (protease), RT (reverse transcriptase), RH (RNase H) and IN (integrase) domains. The LTRs (Long Terminal Repeats) are indicated by wide arrows. Gypsy and Copia differ in the order of the regions that encode the reverse transcriptase and the integrase within the pol region. In 2, the class II is represented by Tc1-Mariner (C). The TIRs (Terminal Inverted Repeats) are indicated by small arrows.
Experimental observation of RIP requires a fungal species cultured under laboratory conditions and lineage transformation. However, processes similar to RIP can also be detected through in silico analyses for repeated elements in the genome using a software (RIPCAL) that can align repeated regions and analyze changes in dinucleotides frequencies related to RIP-like events [25]. Two indices, TpA/ApT and (CpA + TpG)/(ApC + GpT), are commonly used for in silico RIP detection. The TpA/ApT index is simple and measures RIP product frequency, TpA, with a tendency for false positives due to A:T rich regions. High TpA/ApT values indicate a strong RIP response. The (CpA + TpG)/(ApC + GpT) index is similar to TpA/ApT in principle, but it measures depletion of the RIP targets CpA and TpG. In this case, low (CpA + TpG)/(ApC + GpT) values strongly suggest RIP. Additional RIP indices can be defined as (CpN + NpG)/(TpN + NpA), which represents the rate of pre-RIP to post-RIP dinucleotide conversion for the characteristic dinucleotide mutation CpN to TpN and the reverse complement NpG to NpA [25]. A complementary method to the analysis of dinucleotide preference rations, developed by Bouvet et al. [26], can be utilized also. The method, called cumulative transition score (CTS), adds a visual component to analysis of RIPed sequences.

RIP is highly frequent in *N. crassa*, where no intact TE was found because a single pass through the sexual cycle causes mutations in up to 30% of CG pairs in duplicate sequences [27]. Alternatively, such mutations may have a low efficiency as in *Podospora anserina* [28] or be absent. Through sequencing of entire fungal genomes, RIP-like processes have among others also been detected in *P. anserina* [28], *Stagonospora nodorum* [29], *M. grisea* [6] and *Mycosphaerella fijiensis* [30], *Cochliobolus heterostrophus* [31], and *Mycosphaerella graminicola* [33]. However, differences in the intensity with which RIP acts between different transposons, within the same genome, have been reported [33,34].

Despite the in silico evidence of RIP in a wide range of fungi [35], the actual mechanism of RIP is still uncertain. For instance, the importance of the RID proteins for RIP in *N. crassa* was confirmed experimentally [24], but remains unclear in others fungi. In *O. novo-ulmi* and others related fungi, the RIP mechanism has been evidenced, but the RID gene does not appear to be present in the genome assembly [36]. The authors suggested three possible explanations: a recent loss of RIP function, an unidentified functional analog of RID maintaining RIP function or the RID gene may be present in the small assembly gaps. In fact, in many genomes where RIP was evidenced,
Transposable elements as molecular markers

In addition to an important role in genome evolution, TEcs have a broad applicability as a molecular marker system due to their ubiquity, abundance and dispersion throughout the genome and conserved regions [43]. In many species, markers based on TEs can be successfully used to study genetic diversity and variability, such as SSAP (sequence-specific amplified polymorphism) used in Fusarium oxysporum [44], REMAP (retrotransposon-microsatellite amplified polymorphism) in mushroom species [45] and IRAP (inter-retrotransposon amplified polymorphism) in M. fijensis [46]. IRAP and REMAP are exceptional TE markers because they are simple and efficient systems that require only one PCR experiment followed by electrophoresis to resolve the products (Figure 2). IRAP and REMAP examine polymorphisms in regions where retrotransposons are inserted. IRAP studies polymorphisms between retrotransposons, whereas REMAP is used for the analysis of polymorphisms between retrotransposons and microsatellites (SSR - single sequence repeat). Both techniques use primers designed against conserved retrotransposon regions, such as LTRs. For IRAP, the primer pairs anneal to known and conserved retrotransposon regions. In REMAP, one primer that anneals to an LTR is used in combination with primers designed to anneal to SSRs, thereby detecting retrotransposon insertions near SSRs [47]. Polymorphisms are detected through presence or absence of loci. The absence of a band indicates the absence of a retrotransposon at a particular locus. Thus, the presence or absence of the TE at a given locus can be used as a marker to study diversity [48].

Conclusions

The availability of fungal genomes has facilitated the identification and characterization of TEs, which have generated important information on transposition mechanisms, such as activation of such elements, their horizontal and vertical distribution, gene silencing, host genome organization and evolution, insertion of such elements into gene regulatory networks, as well as utilization of such elements as molecular markers. TEs likely play a strong role in evolution, especially for plant pathogenic fungi, because the activity of transposons and rearrangements can potentially alter gene structure or expression and, therefore, facilitate to the emergence of new pathogen races.

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