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# ***P* Element Temperature-Specific Transposition: a Model for Possible Regulation of Mobile Elements Activity by pre-mRNA Secondary Structure<sup>1</sup>**

**A. Gulyaev<sup>a</sup>, T. Redchuk<sup>b</sup>, A. Korolova<sup>c</sup>, and I. Kozeretska<sup>c\*</sup>**

<sup>a</sup>*Leiden Institute of Chemistry, Leiden, 2300 RA Netherlands*

<sup>b</sup>*Palladin Institute of Biochemistry, ul. Leontovicha 9, Kyiv, 01030 Ukraine*

<sup>c</sup>*Taras Shevchenko National University of Kyiv, ul. Volodymirs'ka 60, Kyiv, 01601 Ukraine*

\**e-mail: iryna.kozeretska@gmail.com*

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**Abstract**—*P* element is a DNA transposon, known to spread in genome using transposase activity. Its activity is tissue-specific and normally observed at high temperatures within 24°C to 29°C. Here, we present a predicted RNA secondary structure domain of *P* element pre-mRNA which could potentially regulate the temperature sensitivity of the *P* element activity. In canonical *P* elements, the structure is a small hairpin with double-helical part interrupted by a symmetric loop and a mismatch. In *M* type *P* elements, the A.A mismatch is substituted by an A–U base pair, stabilizing the structure. The hairpin structure covers the region involving the IVS–3 5' splice site and both pseudo-splice sites F1 and F2. While the IVS–3 and F1 binding sites of U1 snRNA are located in the double-stranded part of the structure, the F2 site is exposed in the hairpin loop. The formation of this structure may interfere with landing of U1 snRNA on IVS–3 site, while F2 is positioned for the interaction. Alignment of *P* element sequences supports the proposed existence of the hairpin, showing high similarity for this region. The hairpin structure, stable at low temperatures, may prevent correct IVS–3 splicing. Conversely, temperature-induced destabilization of the hairpin structure may result in the splicing at the proper IVS–3 splice site. Taking into account the increasing amount of data demonstrating the important influence of RNA folding on phenotypes determined by alternative splicing a model for possible regulation of the activity of mobile elements by pre-mRNA secondary structure seems intriguing.

**Keywords:** *P* element, RNA structure, alternative splicing, transposase

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## INTRODUCTION

Mobile genetic elements are widespread in eukaryotic genome and play a significant role in mutagenesis, genome regulation, and evolution [1–4]. *P* element is a DNA transposon that spread in wild populations of *Drosophila melanogaster* after most common laboratory strains had been isolated in the early 20th century [5]. Wild stocks carrying *P* element are referred to as P strains, and lab stocks that lack these elements are referred to as M strains [5, 6]. Crosses between P strain males and M strain females thus lead to a phenomenon called P–M hybrid dysgenesis, which is characterized by *P* element mobilization and reduced fertility in F1 progeny, restored in females with aging [7]. Reciprocal crosses between P strain females and M strain males produce genetically identical female progeny, but these hybrids are viable and fertile due to maternal deposition of *P* element piRNAs [8]. *P* element transposition is tissue- and temperature-specific; transposition occurs at high rates in the germ line

but is not detected in somatic tissues. Furthermore, *P* element activity in the germ line is normally observed at temperatures within 24 to 29°C and absent at temperatures lower than 24°C [9].

*P* element is known to spread in genome using its transposase activity. The transposase pre-mRNA, encoded by *P* element, undergoes alternative splicing in a tissue-specific fashion. The canonical transposase gene consists of 4 exons. In germline cells, all the three introns are excised with the subsequent translation of an 87-kDa active protein. Translation of *P* element mRNA retaining the third intron in somatic tissues leads to the synthesis of a truncated 67-kDa product. The truncated protein not only fails to work as a transposase, but also inhibits the transposition [10]. The intron retention is determined by a 97 kDa protein named *P* element somatic inhibitor (PSI), which is specifically expressed in somatic cells and functions as a tissue-specific splicing repressor [11]. Splicing assays, mutagenesis and RNA binding studies indicate that PSI binds to unprocessed transposase pre-mRNA and recruits U1 snRNP to pseudo-splicing

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sites located 12 to 31 bases upstream of the 5' end of the 3d intron, precluding its excision [12–14].

As mentioned above, in germline cells splicing occurs only at relatively high temperatures above 24°C. In spite of the fact that the origin of PSI–determined tissue specificity is now pretty well understood, little is known about the temperature dependence of the process.

While the temperature effect was first reported over 30 years ago, the nature of this mechanism remains unclear, perhaps due to the absence of proper methods to address this question. On the other hand, alternative splicing is known to be frequently regulated by RNA structures folded in pre–mRNA [15]. In particular, splicing efficiency may be sensitive to the thermodynamic stability of structures folded in the regions of splice sites [16].

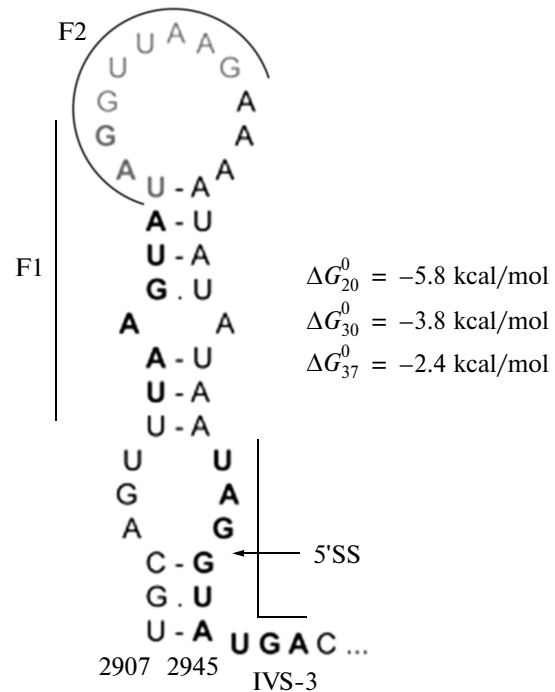
In this study, we present a predicted RNA secondary structure domain of *P* element pre–mRNA which could potentially regulate the temperature sensitivity of the *P* element transposase activity in germ cells.

## MATERIALS AND METHODS

RNA secondary structures were predicted on the basis of both comparative structure analysis and folding thermodynamics. In order to identify the structures conserved in related *P* elements, thus more likely to be functional, we used the RNAalifold algorithm, designed for prediction of common structures in RNA sequence datasets [17]. For development of initial models, representative *P* element sequences of different groups, such as canonical, were aligned and the multiple alignments were used as an input in the RNAalifold program. The regions of the *P* element intron 3 together with flanking exon sequences were used. The predictions yielded by RNAalifold algorithm were further analysed using free energy calculations by mfold program [18] and multiple alignments of *P* element sequences available in GenBank. This allowed us to estimate thermodynamic stability of predicted structures and the extent of their conservation in *P* element subfamilies.

## RESULTS

The most conserved structure in the area of intron 3 (IVS–3) was predicted by RNA–alifold algorithm (see Methods) in the 3'–proximal part of exon 3, in the region of the 5' splice site (Fig. 1). In canonical *P* elements, the structure is a relatively small hairpin with double-helical part interrupted by a symmetric loop and a mismatch. In M type *P* elements, the A.A mismatch is substituted by an A–U base pair, apparently stabilizing the structure (Fig. 2). In the elements of *O* type and *K* type, some of the base pairs are disrupted, destabilizing the hairpin (not shown).



**Fig. 1.** *P* element RNA secondary structure prediction for the region around 5' splicing site of IVS-3. The canonical sequence of *P* element from *D. melanogaster* X06779, coordinates of F1, F2 pseudo SS and accurate 5'SS marked according [19].

Interestingly, the hairpin structure covers the region involving the sequences of the IVS–3 5' splice site and both pseudo-splice sites F1 and F2 [12, 13] (Fig. 1). While the IVS–3 and F1 binding sites of U1 snRNA are mostly located in the double-stranded part of the structure, the F2 binding site is exposed in the hairpin loop. Obviously, the formation of this structure may interfere with landing of U1 snRNA on one of the three binding sites: most likely the IVS–3 and F1 sites are hidden in the stem, while F2 site is positioned for the interaction with U1 snRNA. Remarkably, the hairpin is significantly destabilized within the range of temperatures known to switch on the splicing process: while at 20°C the free energy is –5.8 kcal/mol, its value is –3.8 kcal/mol at 30°C and only –2.4 kcal/mol at 37°C, as calculated using mfold program with the version 2.3 thermodynamic parameters [18].

Of course, the (temperature-dependent) formation of the hairpin in the 5' splice site region is not only dependent on its own folding free energy, but on the possible alternative conformations that may interfere with the hairpin folding. For instance, an alternative stem-loop structure is also possible in the same region, involving a larger part of IVS–3 (Fig. 2). Apparently, its folding would prevent the formation of the hairpin shown in Fig. 1.

Multiple alignment of *P* element sequences (Fig. 2) supports the proposed existence of the first hairpin



**Fig. 2.** Alignment of the sequences in the region of IVS–3 5' splice site in canonical and M type *P* elements. Representative sequences were selected according to the phylogenetic tree of Loreto et al. [20]. Paired regions of the predicted RNA structures in canonical *P* element (according X06779) are shown by brackets. Canonical: *D. melanogaster* X06779 2860–2980 (Dmel), *D. nebulosa* M17424 1860–1980 (Dneb), *D. willistoni* AY578739 1834–1954 (Dwilcons1), *D. sturtevantii* AY578782 1691–1811 (Dstucons1), *D. sturtevantii* AY578776 1834–1954 (Dstucons2), *D. sturtevantii* AY578781 1792–1912 (Dstu7), *D. mediopunctata* AF313770 1827–1947 (Dmed). M type: *Lordiphosa miki* AF313772 478–598 (Lmik), *D. Helvetica* AF313771 1846–1966 (Dhel), *D. bifasciata* X60990 1885–2005 (Dbif), *Scaptomyza pallida* M63342 1878–1998 (Spal). Substitutions as compared to the reference Dmel are marked by grey boxes (for the putative hairpin structure regions only). F2—the second pseudo–SS and the true 5'SS are marked accordingly [19].

2907–2945 in the canonical and M type elements, showing rather high sequence similarity for this region among the analyzed genes. Only two substitutions led to mispairings: 553 T > A transition in *L. miki* and 1906 A > G in *D. mediopunctata*. The rest of the nucleotide differences are observed in regions predicted to be loops in the reference sequence. Moreover, transversions 554 A > T (bp numbering according to AF313772) even closed small internal loop (A.A mismatch) in the hairpin stem of M type structure. Similarly, 526 G > A (also observed in M type species) converts the non–Watson–Crick GU pair into the canonical AU. The presence of trans-versions increasing the hairpin stability in M type elements is in accordance with the splicing patterns of *P* element-derived mRNAs of *D. bifasciata* and *S. pallida* and could possibly explain the immobility of the M type *P* elements subfamily [21].

As can be seen from the alignment, the alternative hairpin 2911–2971 region is less conservative among the analyzed species.

**DISCUSSION**

Since up to 85% of some eukaryotic genomes studied so far consist of mobile element [22], and sequences similar to drosophila *P* element have been detected in the genomes of yeast, fish, birds, mammals, and even humans [23, 24], interest in understanding of the mechanisms of *P* element regulation reaches far beyond the biology of *D. melanogaster*.

It has long been known that the structure of pre-mRNA may have a strong effect of on spliceosome assembly, shown both in vitro and in vivo [25]. A variety of such examples have already been described in literature for different unrelated species, revealing their ubiquitousness. A well-known illustration of the pre-mRNA secondary structure influence on splicing is the human gene *tau* [26, 16]. In this case, a stem-loop structure controls recognition of the 5' splicing site of an intron by spliceosome. Destabilization of the stem loop opens an access of splicing machinery to the splice site.

Similarly, the structure predicted in the *P* element pre-mRNA (Fig. 1) may suppress recognition of the IVS-3 5' splice site. Furthermore, this folding exposes one of the pseudo-splice sites in the hairpin loop, presumably facilitating U1 snRNA binding to the F2 pseudo splice site sequence. Thus, the hairpin structure, stable at low temperatures, may prevent correct IVS-3 splicing. At the elevated temperatures, the hairpin becomes less stable and/or is disrupted in favour of an alternative folding. This is expected to result in both increased efficiency of IVS-3 splice site recognition and a decrease in the F2 site binding by U1 snRNA.

An increased binding of U1 snRNA to a 5' splice site located in the loop of pre-mRNA secondary structure has been recently shown for the serotonin receptor 2C gene [27]. In this case, a dependence of U1 snRNA binding on RNA structure makes splicing sensitive to RNA conformational changes: the switching between alternative structures, triggered by the presence of a small molecule—pyrvinium palmoate, regulates splicing efficiency. Similarly, temperature-induced destabilization of the structure exposing the F2 pseudo splice site of *P* elements in the hairpin loop (Fig. 1), may result in the splicing at the proper IVS-3 splice site.

Such a transition may involve formation of alternative structures competing with the hairpin formation. We could readily identify a potential for formation of such structures (Fig. 2), especially because the hairpin structure does not have a low folding free energy. These structures are less conserved than the hairpin in the F1-F2 pseudo splice site region, and may be specific for various *P* element subfamilies. On the other hand, conformational transitions in the splice site regions may also involve complex tertiary structures such as e.g. pseudoknots, suggested to regulate the splicing of influenza virus mRNAs [28, 29]. We cannot exclude a possibility of similar effects of 3D RNA folding on the U1 snRNA binding to *P* element 5' splice site.

Apparently, the existence of the putative structures described in our study and their possible effect on the splicing of the 3rd intron still has to be confirmed by mutational studies. Taking into account the increasing amount of data demonstrating the important influence of RNA folding on phenotypes determined by alternative splicing [15, 30], a model for possible regulation of the activity of mobile elements by pre-mRNA secondary structure seems intriguing.

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