Intricate RNA: RNA Interactions in U12-Dependent Nuclear Pre-mRNA Splicing

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INTRICATE RNA: RNA INTERACTIONS IN U12-DEPENDENT NUCLEAR
PRE- mRNA SPLICING

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Coding regions or exons of most human genes are interrupted by noncoding intervening regions or introns. Removal of nuclear precursor messenger RNA (pre-mRNA) introns by RNA splicing is an essential step in eukaryotic gene expression. Two types of nuclear pre-mRNA introns are known as U2-dependent or major type and U12-dependent or minor type. Nuclear pre-mRNA introns are removed by two distinct sets of ribonucleoprotein complexes or spliceosomes, which are formed by five small nuclear RNAs (snRNAs) for each spliceosome. U6atac and U12 snRNAs are central to U12-dependent spliceosome and play essential roles in the removal of U12-dependent introns. U6atac and U12 snRNAs bind to the 5’ splice site and branch site, respectively of an U12-dependent intron. In addition, it has been predicted that, U6atac and U12 snRNAs interact inter-molecularly to form helix I structure, which appears to be an essential element of the minor spliceosome. We have been studying U6atac and U12 inter-molecular base-pairing interaction using an in vivo mutation suppression assay. In this study, we have characterized U6atac and U12 mediated helix I intermolecular interactions and have shown in vivo existence of the predicted structure. In addition, we have also identified a region of U6atac snRNA which appears to be a structural
analog of U12 snRNA stem III element. This element is important for the function of U12 snRNA and functions by binding to a RNA binding 65K protein, which is unique to minor spliceosome. We show that, analogous stem-loop of U6atac snRNA also interacts with 65K - RNA binding protein. However, functional significance of this interaction remained unclear. In summation, we have characterized sequential and dynamic RNA-RNA interactions between U4atac-U6atac and U6atac-U12 snRNAs. Our data show that, extensive and obligatory RNA-RNA interactions are critical to the splicing of U12-dependent introns.
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LIST OF ABBREVIATIONS

ATP Adenosine Tri-Phosphate

cDNA complementary Deoxyribo nucleic Acid

CHO Chinese Hamster Ovary

CTP Cytidine Tri-Phosphate

DM Double Mutant

DMEM Dulbecco’s Modified Eagle’s Medium

dNTP deoxy Nucleotide Tri-Phosphate

EDTA Ethylene Diamine Tetra-acetic Acid

EMSA Electrophoretic Mobility Shift Assay

FBS Fetal Bovine Serum

GTP Guanosine Tri-Phosphate

PAGE Polyacrylamide Gel Electrophoresis

Pre-mRNA Precursor messenger Ribonucleic Acid

RRM RNA Recognition Motif

RT-PCR Reverse Transcriptase Polymerase Chain Reaction

snRNA small nuclear RNA

TBE Tris-Borate-EDTA

UTP Uridine Tri-Phosphate

WT Wild Type
CHAPTER I

INTRODUCTION

1.1. Mechanism of Gene Expression

The genes can be compared to books in a library. Just like a book may be taken off a shelf and chosen to be read, similarly, a gene can be expressed to produce a functional RNA and protein molecules in the cell. Gene expression initiates at transcription. Here, the gene (DNA) is copied to produce an RNA molecule. This primary transcript or precursor mRNA essentially has the same sequence as the gene. These sequences are called exons and introns. Of the exons and introns, only the exons carry the information essential for protein synthesis. So, the intron sequences need to be removed from the primary transcript by nuclear pre-mRNA splicing. Splicing results a mature transcript of RNA (messenger RNA) that contains only exons (Figure 1.1). Subsequent step of gene expression is called translation. Here, the mRNA is translated into a sequence of amino acids, which eventually define a protein. Just as some books in a library are well thumbed and others are not equally consulted, likewise, there are multiple mechanisms to regulate transcription and translation, which result to differential gene expression (Genetics, Third Edition, 1997, Robert F. Weaver and Philip W. Hedrick).
1.2. Eukaryotic Pre-mRNA Processing

Splicing is the process of modification of RNA in which introns are removed and exons are joined. Typically, precursor transcript of a eukaryotic messenger RNA undergoes through a series of reactions to become a mature mRNA transcript. These reactions are catalyzed by a complex, called spliceosomal complex, constituting of small nuclear ribonucleoproteins (snRNPs) (Genetics, Third Edition, 1997, Robert F. Weaver and Philip W. Hedrick). So, spliceosome is, generally, a huge RNA-protein complex composed of five snRNPs. RNA components of snRNPs interact with the introns (Figure 1.2). Within an intron, there is 5’ splice site (ss), 3’ splice site (ss) and a branch site. 5’ss is called splice donor site. 3’ss is called splice acceptor site.

Two distinct sets of spliceosomes have been identified which contain different snRNPs. Moreover, the type of splicing depends on structure...
of the spliced intron and the catalytic components for splicing (Dietrich RC et al, 1997).

1.3. Biochemical Pathway of Spliceosomal Splicing

Eukaryotic pre-mRNA undergoes two sequential transesterification reactions, catalyzed by snRNPs. First, 2’OH of branch point nucleotide (Adenosine) within the intron performs a nucleophilic attack at first nucleotide of 5’ss of intron. A lariat intermediate is formed. Secondly, 3’ OH of the released 5’ exon performs a nucleophilic attack at last nucleotide at 3’ss of intron. Flanking exons are joined (Figure 1.3). The final products of splicing are: ligated exons (mature transcript) and free intron (lariat debranches), while the snRNPs are recycled (Genetics, Third Edition, 1997, Robert F. Weaver and Philip W. Hedrick).

1.4. Introduction of Introns into Eukaryotic Genomes

In higher eukaryotes, there is a large variation in the
length and number of introns. One example is the longest human gene coding for the muscle-specific protein dystrophin, which spans 2.5 million bases, with its exons accounting to < 1% of total length, but, average intron length is 26,000 bases (Pozzoli et al. 2002).

Introduction of introns into eukaryotic genomes follows two hypotheses (Darnell 1978, Doolittle 1978, Basu et al., 2007): according to the “intron early” theory, reading frames are discontinuous and so the prokaryotes lost their introns with the increasing compactness of their genomes; whereas, according to “intron late” theory, introns are getting inserted into genes throughout eukaryotic evolution. In addition, the “exon theory” of gene evolution postulated by Gilbert (1987) is based on the idea that, each exon was a separate gene coding for a discreet protein domain and proteins remained functional long before the splicing mechanisms evolved. It was considered that “splicing” event is responsible for formation of longer genes with many exons.

Taken together, the removal of introns from the pre-mRNA is, undoubtedly, a significant event of gene expression.

**Figure 1.3:** Biochemistry in pre-mRNA splicing: 3’OH of branch point (BP) of Adenosine (A) in the intron attacks 5’ splice site (ss). Lariat is formed. 3’OH of 5’ exon triggers a second trans-esterification at 3’ss. Exons are joined. (Reference: Molecular Biology by Robert Weaver).

**Figure 1.4:** Major and minor class introns and snRNAs involved: Consensus 5’splice site, branch point sequence and 3’ splice site are shown for major and minor class introns. (Reference: Molecular Biology by Robert Weaver).
1.5. Spliceosomal Introns

Spliceosomal introns have an almost invariant sequence GT at 5’ SS end of the intron. An almost invariant sequence AG at 3’ SS terminates the intron sequence. Upstream from AG, there is usually a region high in pyrimidines (C and T). It is called Poly-Pyrimidine Tract (Figure 1.4). Upstream of Poly-Pyrimidine tract is the branch point (BP), which usually includes an adenosine (A) nucleotide (Sheth et al., 2006). Majority of introns with GT-AG as flanking sequences, at 5’ SS and 3’ SS, respectively, account for canonical type of intron splicing. For such introns, U1, U2, U4, U5 and U6 snRNPs constitute the major spliceosomal complex. Their spliceosomal machinery is called major spliceosome. Their splicing process is termed as U2-dependent or major spliceosomal pathway. With short stretches of highly conserved sequences, they account for more than 99% of splicing.

Between years 1989 and 1997, introns with non-canonical flanking splice sites, as for example, in the proliferating cell nucleolar protein P120 gene in Human, were reported. This subset of introns has long and nearly invariant consensus sequences (Jackson, 1991). The first minor class introns found had AT-AC as terminal di-nucleotides instead of the canonical GT-AG, and these introns represent < than 1% of all introns in human. This splicing system uses a different set of snRNAs and it was confirmed by mutational and biochemical analyses (Hall and Padgett, 1996, Tarn and Steitz, 1996a and 1996b, Burge CB, Padgett RA and Sharp PA, 1998). Later, U12-type introns with non-canonical terminal dinucleotide (CT-AC, GG-AG and GA-AG) were also reported (Dietrich RC et al., 1997). Nowadays, their spliceosomal machinery is called minor spliceosome, constituting U11, U12, U4atac, U5 and U6atac snRNPs. Their splicing process is termed as U12-type or U12-
dependent or minor spliceosomal pathway. U12-type introns are found in most multicellular organisms. Higher proportion of U12-type introns occur in more complex eukaryotes, as revealed from computational analyses on intron frequencies (Levine and Durbin, 2001, Schneider et al, 2004, Alioto 2007). However, simple eukaryotes like nematode (Trichinella spiralis), protists (Physarum polycephalum) and fungi (Phytophthora sp.) also have U12-type introns, but, another nematode (Caenorhabditis elegans) is the most notable exception where components of U12-pathways are absent.

The terminal di-nucleotides, however, do not distinguish between the major or U2-type and minor or U12-type spliceosomal introns. As mentioned earlier, many U12-type introns have GT-AG as flanking di-nucleotides. Similarly, many introns with AT-AC as flanking di-nucleotides are spliced by U2-type spliceosome (Wu and Krainer, 1999). However, the persistent conservation of U12-type intron locations between human and plant over such a broad evolutionary distance indicates that U12-type introns have important role in gene expression (Zhu and Brendel, 2003). But, U12-type introns are less efficiently spliced than U2-type introns since pre-mRNA with unspliced U12-type introns are found to be abundant in cells (Patel et al, 2002).

### 1.6. Splicing Fidelity

Splicing mechanics depends on the accurate recognition of introns. In complex eukaryotes, splice sites are often degenerate to demarcate discreet intron boundaries without other signals. So, alternative splicing is common.
For example, over 90 percent of human genes are alternatively spliced (Wang et al., 2008). Long consensus sequences of U12-type 5’ss and BP is sufficient for recognition of splice sites, but due to low abundance of U12-type introns, even long sequences are not adequate to locate the introns.

Although splicing patterns are rarely alternative splicing for U12-type introns because of their long consensus sequence and the co-operative intron recognition by U11/U12 di-snRNP, one example of alternative splicing event is on the mRNAs of U11/U12 48K protein. Also, there are instances, where an intron can be spliced by U2-type spliceosome or U12-type at slightly differing splice sites.

Moreover, the *prospero* gene of *Drosophila* (fruit fly) contains a “twintron” consisting of a U2-type intron within a U12-type intron. The splicing of twintron is regulated by a purine–rich element which maneuvers the function of both types of spliceosomes (Hall and Padgett, 1994, Borah et al., 2009).

Another means of shuffling of exons among different mRNA molecules, is first identified in *Trypanosoma*, where the regulated splicing is termed as trans-splicing (Bonen, 1993).

1.7. **Major Spliceosomal Components and Their Biogenesis**

MS (mass spectrometry) sequence of spliceosomal complexes shows that the spliceosome is a muti-megadalton ribonucleoprotein complex and can be defined as the largest intracellular machine (Deckert et al., 2006, Jurica and Moore, 2003).

Five snRNAs namely, U1, U2, U4, U5 and U6 are the core components of U2-type or major spliceosomes (Krainer and Maniatis 1985). U1,
U2, U4 and U5 snRNAs, each has U-(uridine) rich conserved sequence, the Sm region, which binds to seven common proteins, namely, B/B’, D1, D2, D3, E,F and G (Bringmann and Luhrmann, 1986). U6 snRNP lacks Sm proteins but it has Sm-like proteins, namely Lsm 2-8 (Seraphin, 1995, Salgado-Garrido et al, 1999). snRNP – specific proteins like the p110/SART3 of U6snRNP restore the U4-U6 snRNAs base-pairing (Bell et al, 2002). DExD/H box proteins like U5-100K, with ATPase and RNA helicase activity are required for conformational changes during spliceosome formation (Staley and Guthrie, 1999). Many non-snRNP proteins mediate RNA-RNA interactions, as for example, U2AF-65 anneals U2 snRNA with branch point sequence (Valcarcel et al, 1996). Moreover, auxiliary proteins are also required to bind to exonic and intronic splicing enhancers in addition to the spliceosomal core proteins. The composition of proteins in major spliceosomal snRNPs are pictorially represented (Figure 1.5).

![Figure 1.5: Composition of proteins in major spliceosomal snRNPs (Human): “Sm” denotes Sm proteins (B/B’, D3, D2, D1, E, F and G. “Lsm” proteins are (Lsm 2-8) associated with each snRNP. U4/U6.U5 tri-snRNP contains two sets of Sm proteins and one set of Lsm proteins.](image)
In multicellular organisms, spliceosomal snRNPs transiently localize in cytoplasm during their biogenesis (Will and Lührmann 2001, Patel and Bellini 2008). After transcription by RNA polymerase II, each of U1, U2, U4 and U5 gets a mono-methylated cap and are exported from the nucleus (Ohno et al. 2000). In cytoplasm, seven Sm proteins assemble on Sm site, which is assisted by SMN complex consisting of seven Gemin proteins (2-8) (Pellizzoni et al. 2002). Next, the cap modifies to 2, 2, 7-tri-methyl-guanosine (m3G) structure (Mattaj 1986). Both the cap and Sm complex contain nuclear localization signals (NLS) that re-import snRNPs back to nucleus, being mediated by the import adapter snurportin1 (Fischer and Lührmann, 1990, Huber et al. 1998, Narayan et al. 2004). Now, snRNPs acquire their specific proteins and are internally modified by pseudo-uridylation and 2’-O-methylation (Yu et al. 2001). Newly assembled snRNPs accumulate in sub-nuclear organelles termed Cajal bodies (Sleeman and Lamond 1999, Jády et al. 2003). Cajal bodies may also be the sites for recycling of the U4/U6.U5 tri-snRNP (Stanek et al. 2003, Schaffert et al. 2004). Mature snRNPs are enriched in interchromatin granule, which are presumed as storage sites for splicing factors (Matera and Ward, 1993, Lamond and Spector 2003).

U6 snRNA is transcribed by the RNA polymerase III (Kunkel et al. 1986) and is believed not to be exported out of the nucleus. The poly-uridine tail of nascent U6 is transiently bound by the La protein (Rinke and Steitz 1985, Pannone et al, 1998). The triphosphate cap is modified into a gamma-monomethyl phosphate cap (Shimba and Reddy 1994). As the other spliceosomal snRNAs, U6 is also modified by pseudo-uridylation and 2’-O-methylation by small nucleolar RNPs, probably in the nucleolus (Tycowski et al, 1998). U4 and U6 snRNPs interact by base

1.8. Minor Spliceosomal Components and Their Biogenesis

U1, U2, U4 and U6 snRNAs of U2-type or major spliceosome are functional analogues for U11, U12, U4atac and U6atac snRNAs of U12-type or minor spliceosome, respectively, whereas U5 is shared between both splicing machinery (Montzka and Steitz 1988, Hall and Padgett 1994, Tarn et al. 1995, Hall and Padgett 1996) (Figure 1.6).

Proteomic analysis of purified spliceosomal complexes indicate that, over 170 proteins are found associated with the major spliceosome, although many proteins are also shared by the minor spliceosome. Unlike U1 and U2, which exist mainly as mono-particles, U11 and U12 form a di-snRNP through protein-protein interactions (Wassarman and Steitz 1992b) and lack U1-specific proteins but contain most proteins present in U2 snRNP, such as the SF3b complex. U11, U12 and U4atac snRNPs contain Sm protein complex.

In addition, U11/U12 snRNP contains seven unique, novel proteins not found in the U2-dependent spliceosome, designated 65K, 59K, 48K, 35K, 31K, 25K, and 20K, all of which are essential for cell viability (Will et al. 1999, Will et al. 2004). 59K, 48K, 35K and 25K are detected also in U11 mono-particles. 65K binds U12 snRNA and 59K binds U11 snRNA, thus bridging U11 and U12 snRNPs (Benecke et al. 2005). 35K recognizes 5' splice site just like its homolog, U1-

U11/U12-65K, -35K and -31k proteins, which are found to contain RNA Recognition Motifs (RRMs), are the essential snRNA binding proteins, which are presumably involved in di-snRNP formation and intron bridging.

The minor spliceosome tri-snRNP U4atac/U6atac.U5 forms similarly (Tarn and Steitz 1996a, Tarn and Steitz 1996b) like major tri-snRNPs. Protein components of the U6atac/U4atac.U5 tri-snRNP, also, do not differ from those in the major tri-snRNP (Luo et al., 1999, Nottrott et al., 2002, Schneider et al., 2002). Just like U6snRNA, U6atac snRNA lacks Sm binding site and instead, it is assembled with Sm-like proteins, the Lsm 2-8 complex, which replaces the La protein and serves as NLS (Achsel et al. 1999, Schneider et al., 2002, Spiller et al., 2007).

In sum, it is important to note that, proteins involved in the splicing play critical role in recognizing and pairing of splice site and ensure the dynamicity as well as conservation of the splicing event.

1.9. Catalytic Reactions during Splicing

RNA catalyzes splicing reactions and the spliceosome becomes a ribozyme (Kruger et al. 1982, Cech 1986; Valadkhan 2007). For example, U2 and U6 snRNAs catalyze a splicing-like reaction free of proteins in vitro (Valadkhan and Manley 2001, Valadkhan et al. 2009). RNA molecules in catalytic core of the spliceosome form structures which look identical to the conformation of self-splicing Group II introns. So, spliceosomal introns have presumably evolved from
self-splicing introns (Sharp 1985, Toor et al. 2008).

The spliceosome might have originated through dispersion of the self-splicing intron ribozyme core into many small RNAs (Jarrell et al. 1988, Sharp 1991). The fact that, catalytic domain of self-splicing group II intron functionally replaces the metal binding stem loop in U6atac snRNA supports this presumption (Shukla and Padgett 2002).
1.10. U2-type / Major Spliceosomal Assembly and Dis-assembly

RNA components of snRNPs interact with the intron along with many other proteins which are required for the spliceosomal assembly. After final products of splicing are formed, snRNPs are recycled for next round of splicing. In U2-type, these are the following sequential events (Figure 1.7):

In E complex, splice site consensus sequence is first recognized by U1snRNP and U1 snNRA binds to GT at 5’ splice site (ss). SF1 (splicing factor 1) binds to branch point sequence. Accessory proteins, U2AF subunits 65K and 35K, bind to Poly-pyrimidine tract and 3’SS, respectively (Mount et al, 1983, Black et al, 1985, Zamore and Green, 1989, Abovich and Rosbash, 1997, Staley and Guthrie, 1998, Smith et al, 2008). The base pairing interactions of U1 with 5’ss spans -1 to +6 nts. with respect to 5’ss.

In A complex (pre-spliceosome complex), U2snRNP displaces SF1 and binds to branch site with ATP hydrolysis and bulges out the branch point adenosine (Parker et al. 1987, Query et al. 1994).

of splicing (Newman and Norman 1992, Sontheimer and Steitz 1993). The protein composition of the activated spliceosome is also re-modeled now (Makarov et al. 2002).

In C1 complex, U4 is released. U6/U2 catalyzes trans-esterification. 5’end of intron ligates to A on intron. Lariat forms. In C2 complex, U2/U5/U6 remains bound to lariat. 3’ss is cleaved. Exons are spliced with ATP hydrolysis. Final products are spliced RNA and de-branched intron lariat. Some reports suggest that the spliceosome may exist as a preassembled complex. For example, all five spliceosomal snRNPs have been detected in yeast and HeLa cell nuclear extracts (Stevens et al. 2002, Malca et al. 2003). However, a preformed penta-snRNP is not a prerequisite for splicing in vitro (Behzadnia et al, 2006).

Figure 1.7: Spliceosomal formation and activity: The formation of splicing complexes during U2-dependent or major-, and U12-dependent or minor spliceosomal pathways are pictorially represented in the left and right boxes, respectively. Figure reference: Cindy L. Will and Reinhard Luhrmann. Cold Spring Harb Prospect Biol.
1.11. U12-type / Minor Spliceosomal Assembly and Dis-assembly

U12-type spliceosomal assembly is almost similar to U2-type cascade of interactions (Figure 1.7). Unlike U1 and U2 in major spliceosome, U11 and U12 snRNAs form a di-snRNP and recognize the 5’ss and BP co-operatively (Frilander and Steitz, 1999). The first three nucleotides of 5’ ss are recognized by U11-48K protein (Turunen et al., 2008). So, U11 base pairs from +4 to +8 nts. at 5’ss. The U4atac/U5/U6atac tri-snRNP enters to form the B complex. U11 and U4atac snRNAs are displaced. U6atac and U12 snRNAs base-pairs at 5’ss (Yu and Steitz, 1997, Frilander and Steitz, 2001).

Important molecular structures and interactions remain conserved during spliceosomal assembly (Kolossova and Padgett, 1997, Yu and Steitz, 1997, Incorvaia and Padgett, 1998). It has also been proved that, splicing activity is restored when the conserved intra-molecular stem loop of U6atac snRNA is replaced by a chimeric form of U6 snRNA (Shukla and Padgett, 2001).

Moreover, U4atac snRNA could also be replaced by U4 snRNA, if appropriate base-pairing could be restored with U6atac snRNA (Shukla and Padgett, 2004).

1.12. Dynamic RNA-RNA Interactions in Helices of Spliceosomal snRNAs

Iterative conformational changes in snRNAs at different stages of splicing are adapted for effective RNA-RNA interactions. RNAs of Group II introns are found to fold intricately for self-splicing and undergo same stereochemical reactions as the spliceosomal snRNAs with its pre-mRNA substrate.

U2 and U6 snRNAs form three helices, namely Ia, Ib and
II (Madhani and Guthrie, 1992). Helix I is presumed to get disrupted after first step of splicing but re-forms in the next step (Mefford and Staley, 1995; Sashital et al, 2004). Likewise, U2 snRNA forms Stem IIC to promote first catalytic step of splicing and then toggles to another mutually exclusive Stem IIa conformation required for exon ligation (second step of splicing) (Hilliker et al, 2007; Perriman and Ares 2007). In U12-type splicing, U6atac snRNA base pairs with U12 snRNA to interact at helix Ia and helix Ib whose sequences are almost identical in the two splicing systems (Shukla and Padgett 2004).

1.13. Homologous RNA Elements in Catalytic Triad

Both spliceosomal and self-splicing group II introns require the function of a highly conserved, metal-ion (Mg$^{2+}$) -binding catalytic triad (AGC) present in RNA stem-loop elements located in U6 or U6atac snRNAs and also in Domain 5 (D5 Stem 1) of Group II introns (Figure 1.8) (Madhani and Guthrie, 1992; Chanfreau and Jacquier, 1994; Peebles et al,1995; Abramovitz et al,1996; Konforti et al,1998; Yu et al,1995; Yean et al, 2000; Gordon and Piccirilli,2001; Villa et al,2002; Shukla and Padgett,2002; Griffiths-Jones et al,2005; Davila Lopez et al,2008; Butcher and Brow, 2005).

Sequence analyses of a large number of genome for spliceosomal snRNAs have identified snRNA homologies within distant lineages (Lopez et al., 2008). For example, 3’ end domain of U6atac snRNA shows substantial conservation across different species. Within the 3’ end domain, the 5’ stem-loop and the 3’ stem-loop structures are well conserved among divergent eukaryotic lineages (Figure 1.9). 3’ end domain of human U6atac snRNA shows ~65% identity with that in plant. However, the good homology existing between human (Homo sapiens), protist (Physopthora infestans) and plant (Arabidopsis thaliana) in their 5’ stem-loop and 3’ stem-loop structures of U6atac snRNA 3’ end domain predicts the functional significance for such phylogenetic conservation among divergent lineage.

![Figure 1.9: Predicted secondary structures of U6atac snRNA 3’ end domain among divergent eukaryotic lineages: Conserved elements are marked in red, blue and black lines. Figure reference: Shukla Grant and Lopez et al, 2008.](image)

1.15. Evolutionary History of the Two Spliceosomes

Major (U2)–type introns are small ubiquitous in
eukaryotes but U12-type introns are not. Minor spliceosomes remove a small fraction (less than 0.5%) of introns, referred to as U12-type introns (Lin CF et al, 2010). But, analogous structures and similar enzyme kinetics of major (U2) and minor (U12) spliceosomes in the same living organism, suggest their common ancestry. To understand the evolution of the splicing machinery and of the spliceosomal RNAs, phylogenetic distribution of both have been systematically examined.

Earlier studies have proposed (Burge et al, 1998) that a “subtype switch” from AT-AC to GT-AG could serve as an intermediate stage for conversion from U12-type introns to U2-type introns. As an evidential proof, one of the nineteen U12-type introns was detected in fruit fly (Drosophila melanogaster) to be the first known example for U12-type to U2-type intron conversion. But, the loss of U12-type introns was found to be more frequent than such conversion. Moreover, natural U12-type introns with non-canonical terminal dinucleotide (CT-AC, GG-AG and GA-AG) were also reported (Dietrich RC et al, 1997). Investigation revealed an early origin of U12-type spliceosomal RNAs being present in a variety of evolutionarily distinct phyla. U12-type introns were found to be highly conserved among vertebrates.

Other notable information was that, there has been a significant loss of U12-type of introns in invertebrate lineages, compared to vertebrate lineages. From the fact that U12-type introns and snRNAs exist in both plants and animals, it was proposed (Burge et al, 1998) that, the U2- and U12-types splicing pathways diverged from one in the course of speciation (fission) and later merged (fusion) on one organism. By fission-fusion hypothesis, it was reasoned that, the similar secondary structure and interaction between U2- and U12-types of spliceosomal snRNAs are not due to convergent evolution.
In contrast, the endosymbiotic theory mentions of a unicellular eukaryotic ancestor, carrying introns and a spliceosome, diverged to two lineages which gained cumulative differences in spliceosomal components and intron consensus sequences. With time, these two split lineages fused by endosymbiosis and gave rise to the ancestor of present day eukaryotes.

The homologous proteins in U2- and U12-types of spliceosomes could be a result of duplication and specialization of genes for proteins that were primarily shared (Lynch and Richardson, 2002). It was argued that the two splicing systems originated from two Group II type self-splicing introns, and the spliceosomal proteins were similar between the functionally analogous snRNAs (Russell AG et al, 2006). Supporting to this hypothesis, it was suggested (Basu et al, 2008) that U12-type introns first populated ancestral genome to a certain intron density and their successive invasions got restricted due to limited space available for further insertion of new genes. Moreover, analyses of proto-splice sites in human and plant (Arabidopsis thaliana) genomes confirmed that, U2-type introns predate U12-type introns.

It is noteworthy that, there are multiple instances in species or branches, where not only U12 spliceosomal components are missing, the U12-type introns are lacking, as well. A few examples are the Entamoeba histolytica, green algae and fungal lineage, Basidiomycota. In addition, there are instances, where U12-type RNAs were present in very early fungal evolution while they are lacking in Microsporidia (Fungi). Again, U12-type splicing does not occur in Caenorhabditis elegans (nematode), whereas, both U12 introns and U12 snRNAs are present in Trichinella spiralis, which is a deep branching of the nematode tree.
U12 introns have been identified from BLAST searches. In addition, EST sequences have been retrieved using NCBI Entrez to obtain a database of genomic sequences. Investigations reveal that phylogenetic distribution of U12 introns and spliceosomal RNAs support to an early origin of U12 dependent splicing (Davila Lopez M et al, 2008). It is not yet known if the loss of minor spliceosome could result as a consequence of the strong pressure to reduce genome size. But, it has been identified in many similar instances, as discussed above, where U12-dependent splicing was lost somewhere during the course of eukaryotic evolution. Hence, it is concluded that U12-type splicing has a comparatively marginal role and, therefore, getting disposed of in many phylogenetic groups. Moreover, the phylogenetic distribution of U12 introns is directly correlated to the U12 snRNA distribution (Bartchat et al, 2010). But the origins of the two parallel splicing systems remain elusive till date.

1.16. Significance of Study of Minor Spliceosomes

Study of splicing is important in its relevance to diseases. Mutations lead to aberrant splicing. For example, cis-acting mutations in cystic fibrosis trans-membrane conductance regulator gene cause exon skipping (Pagani et al, 2005). Abnormal tau exon 10 splicing causes Alzheimer’s disease (Liu and Gong, 2008). Splicing abnormalities are common in cancer, as for example, oncogenic isoforms of ribosomal protein S6 kinase –β1 increase (Qi Ma et al, 2011).

Abnormal U4atac/U5/U6atac snRNP complex causes retinitis pigmentosa, which characterizes retinal degeneration leading to blindness (Makarova et al, 2002).
Spinal muscular atrophy (SMA) causes childhood lethality, due to mutations in the genes encoding motor neuron protein, SMN. SMN deficit causes reduction in spliceosomal snRNP levels, particularly U11 snRNA (Livio Pellizzoni, 2007).

Four point mutations in U4atac snRNA of minor spliceosome are found in patients with MOPD 1 (Microcephalic Osteodysplastic Primordial Dwarfism type 1) (Huiling He et al, 2011).

The list is inexhaustible but fact remains, that the splicing of minor introns plays critical roles in human development.
CHAPTER II

MATERIALS AND METHODS

2.1. Mutant Construction

The single nucleotide and adjacent two or three nucleotide mutations in U12, U11, U6atac and U4atac snRNAs were generated from expression plasmid background by site directed mutagenesis with Change-IT mutagenesis kit (USB Corporation). Using 10X buffer, 5 µM of 5’phosphorylated mutagenic oligonucleotides (primer), dNTPs and 15 ng of plasmid, the PCR product was subsequently treated to Dpn I, to degrade the template plasmid. The plasmid was transformed into E. coli DH5α competent cells and plated on Ampicillin (100mg/ml) contained in agar plates. Each colony was picked from plate and grown overnight in an autoclaved glass tube with 5ml Luria-Bertani broth (Amresco, OH) and 5 µl of Ampicillin (100mg/ml), in 37 °C at 250 rpm incubator shaker. Zippy miniprep kit (Zymo Research) was used to make plasmid mini-preps from overnight cultures. Once the sequences of mutant snRNAs were confirmed by DNA sequencing, maxi-preps were done using Qiagen Plasmid Maxi Kit. Construction of single nucleotide mutants and adjacent two and three nucleotides mutants are listed below.
Mutants For Helix Ia Region of Interaction:

The single nucleotide mutations on U12 snRNA were A9U, A10U, C11G, U12G and U13A. The single nucleotide mutations on U6atac snRNA were A21U, G22U, G23C, U24A and U25A. The single nucleotide mutations on U4atac snRNA were A63U, A64U and C65G. All mutants were named after the nucleotide position where mutations were done.

The adjacently two nucleotide mutations on U12 snRNA were AA9/10UU, AC10/11UU, CU11/12GG, and UU12/13GA. The adjacently two nucleotide mutations on U6atac snRNA were AG21/22UU, GG22/23UC, GU23/24CA and UU24/25AA. The adjacently two nucleotide mutations on U4atac snRNA were AA63/64UU, and AC64/65UG. All mutants were named after the nucleotide position where mutations were done.

Mutants For Helix Ib Region of Interaction:

The single nucleotide mutations on U12 snRNA were C5G, C5U and C5A, and, U6C, U6G and U6A. The single nucleotide mutations on U6atac snRNA were A26G, A26C and A26U, and, G27A, G27C and G27U. The single nucleotide mutations on U4atac snRNA were C61U, C61G and C61A, and, U62C, U62G and U62A. All mutants were named after the nucleotide position where mutations were done.

Mutants For U6atac snRNA 3’ SL Region:

Mutations for U6atac snRNA stem and loop nucleotides at 3’SL region were generated following the same principle and procedure, using Change-IT site directed mutagenesis kit. (USB Corporation). U11 snRNA mutants, also generated likewise, are listed in Appendix.
2.2. *In vivo* Genetic Suppression Assay

One snRNA expression plasmid, which contains mutations in a region, can restore the complementarity to its interaction with a second snRNA expression plasmid, provided that, this second snRNA expression plasmid contains complementary nucleotide mutations in its region of interaction with the first mentioned snRNA expression plasmid. The mutations in two interacting snRNA expression plasmids can reactivate correct splicing, after co-transfection of both expression plasmids, which express the snRNAs carrying compensatory mutations in their region of interaction.

The same principle behind *in vivo* genetic suppression assay was applied to P120 mini-gene plasmid, which contains exons 5, 6, 7 and 8 and introns E, F and G of the human nucleolar protein P120 gene. The 5’ss (splice site) of U12-dependent intron F, if contains CC5/6GG mutation, abolished U12-dependent splicing of intron F, *in vivo*. On co-transfection with U11 snRNA GG6/7CC suppressor mutant and U6atac snRNA GG14/15CC suppressor mutant, the U12-dependent splicing to the P120 CC5/6GG splice site mutant was restored. So, the restoration of correct U12-dependent splicing to the 5’ss mutant, by compensatory mutants of U6atac snRNA and U11 snRNA, allowed us to test the effect of other alterations in U12, U11, U6atac and U4atac snRNAs on their function, *in vivo*. P120 mini-gene construct, P120 CC5/6GG (mutation only at 5’ss), P120DM mutant (double mutations, at both 5’ss and branch site), U11 snRNA expression plasmid, U6atac snRNA expression plasmid, U4atac snRNA expression plasmid were the starting materials for this assay.

P120 mini-gene construct (P120 WT, wild type) and the U11 snRNA GG6/7CC suppressor mutant and U6atac snRNA GG14/15CC suppressor mutant, cloned in pALTER expression vector, were the combined gift from Dr. Richard
2.3. CHO Cell Maintenance and Transfection

CHO (Chinese Hamster Ovary) cell line was obtained from American Type Culture Collection (Manassas, VA). Cells were cultured in 100 mm plate with 10 ml complete CHO media, containing DMEM (Dulbecco’s Modified Eagle’s Medium, 1X) supplemented with 4.5 g/l glucose and L-glutamine (CCF, OH), antibiotics (100units/ml penicillin G sodium and 100µg/ml streptomycin sulfate), 5% Fetal Bovine Serum (FBS, Atlanta Biologicals, Inc., Lawrenceville, GA), 1 mM L-proline (Sigma, MO) and 10 mM HEPES (SIGMA, MO). All cells were kept in a humidified incubator at an atmosphere of 5% CO$_2$ at 37 °C.

For transfection, a 100% confluent 100 mm plate was split at 1:8 ratios in 100 mm plates, using 10 ml complete CHO media. Cells were ~70% confluent after 24 hours. Morphologically healthy cells were transfected using 5 ml complete CHO media containing 12 µl polybrene (10 mg/ml Sigma, MO) and 1 µg plasmid DNA per plate. 9 µg pUC19 was used as carrier in order to equalize amount of DNA being added. In wells where both U11 suppressor plasmid, U6atac suppressor plasmid and U4atac suppressor plasmid were transfected together, pUC19 vector was omitted. For the followed 8 hours, the transfected plates were maintained in 37 °C in a humidified 5% CO$_2$ incubator, with gentle swirling every one hour. After 8 hours, media from wells were aspirated and cells were treated with 5 ml of 30% DMSO (Dimethyl sulfoxide, prepared in CHO media) and incubated at room temperature for 5 minutes. Plates were washed with 5 ml of CHO media. Final step was replenishing each plate with 10ml of CHO media and putting the plates back to 37 °C in a humidified 5% CO$_2$ incubator. The cells were harvested after 36 hours of incubation from post-DMSO
shock time-point.

2.4. Total RNA Isolation

Isolation of total RNA from cultured CHO cells was done using High Pure RNA Isolation kit (Roche). Cells were suspended in 200 µl PBS in 15 ml tubes. 400 µl of Lysis buffer (pH= 6.6) containing Tris-HCl was added to each tube and contents are vortexed. Contents from each 15 ml tube were transferred to separate polypropylene High Pure Filter tube fitted to a collection tube. Each tube assembly was then centrifuged for a short spin at 8000 x g for 60 seconds. Flow through was discarded and the collection tube was fitted back to each respective filter tube. 90 µl of DNase incubation buffer containing NaCl, Tris-HCl and MnCl$_2$ was mixed with 10 µl of 10 KU DNase I (suspended in Elution Buffer) for removal of contaminant DNA. The contents were incubated for 20 minutes at temperature between 15°C and 25 °C. 500 µl Wash Buffer containing ethanol was then added to each tube assembly followed by a centrifugation for a short spin at 8000 x g for 60 seconds. Flow through was discarded and the collection tube was fitted back to each respective filter tube. The wash buffer step was repeated followed by centrifugation at 13000 x g for 2 minutes to ensure complete removal of residual impurities and residual wash buffer. Flow through was again discarded and, for each time, the collection tube was fitted back to respective filter tube. After a final wash, collection tubes were finally discarded and the filter tubes were fitted to 1.5 ml tubes before the elution step. 100 µl of Elution Buffer was added and each tube was centrifuged at 8000x g for 60 seconds. The micro-centrifuge tubes containing the eluted RNAs were immediately transferred to -80 °C until further use. Each isolated RNA was quantified under spectrophotometer and absorbance was recorded at 260/280 nm.
Furthermore, the integrity of RNA was checked on a denaturing 1% MOPS-Agarose gel.

2.5. Reverse Transcriptase- Polymerase Chain Reaction (RT-PCR)

Gene Amp Thermostable rTth Reverse Transcriptase RNA PCR kit (Applied Biosystems, CA) was used to perform reverse transcription step to a total RNA of 0.4 µg per reaction. Recombinant Thermus thermophilus (rTth) DNA polymerase was used to reverse transcribe RNA to cDNA in presence of Mn^{+2} ions and also for PCR amplification for cDNA synthesis from RNA templates. A master mix of 10 µl reaction volume was made using p120 min-gene specific reverse primer, E-7-8-2 (5’-CTTCTAAGAAGTCCACCAGCTCAGA-3’) at 70 °C for 15 minutes in PTC-100 thermal cycler (BioRad laboratories, CA).

Next step was addition of 40 µl of master mix containing p120 mini-gene specific E-5-6 primer 5’-GGCCCGGAAGCTGCTGCTGGGGATC-3’ to the 10 µl RT product kept on ice. A final reaction volume of 50 µl containing dNTPs and MgCl₂ was ready for PCR with amplification for 39 cycles, consisting of a denaturation step at 95 °C for a minute, annealing and extension step at 55 °C for 1.05 minutes and a final extension step for 7 minutes at 68 °C in a PTC-100 thermal cycler (BioRad laboratories, CA). Positive control pAW109 RNA was provided with control primers DM151 and DM152.

2.6. Nested PCR

Hi Fidelity PCR Kit (Roche, IN) was used to set up a 25 µl total reaction volume for analyzing the in vivo splicing phenotypes.

PCR was done using p120 mini- gene specific primers E6
long 5’- TTGTGCTGCCCTGCTGGGGAGATG-3’ (forward) and E7 long 5’- TGAGCCCCAAAATCAGCAGAATTCC-3’ (reverse). PCR program was set with denaturation at 95 °C for 3 minutes, followed by 29 cycles of denaturation at 95 °C for a minute, annealing step at 60 °C for 1.05 minutes, extension at 68 °C for a minute and a final extension step at 68 °C for 7 minutes. PCR products were analyzed by ethidium-bromide-stained 3 % Agarose (2 % Agarose: 1 % NuSieve agarose) in 1X TBE gel, ready to be observed on Typhoon Scanner.

2.7. *In vitro Transcription*

Maxi script Kit (Ambion, TX) was used to set up *in vitro* transcription of DNA from PCR products. At room temperature, 50 µl reaction volume was made with the following reagents added sequentially to each 1.5 ml micro-centrifuge tube: 12.5 µl of Nuclease free sterile water, 5 µl 10X Transcription Buffer (with DTT, Dithiothreitol), 2.5 µl 10 mM ATP, 2.5 µl 10 mM UTP, 2.5 µl 10 mM GTP, 2.5 µl 10 mM CTP, 10 µl (2 µg) linearized DNA template, 2.5 µl [α-³²P] UTP 800 Ci/mmol (10 mCi/ml- MP Biomedicals, OH) and finally 5 µl T7 RNA Polymerase (10 U/µl). All reactions were incubated at 37 °C for 2 hours. 2 µl RNase-free DNase I (2 U/µl) was added to the reaction mixture to remove DNA template, mixed thoroughly, and incubated for 20 minutes at 37 °C. Gel loading Buffer II (Ambion, TX) was added to the contents in each reaction tube. Samples were loaded on the wells of pre-electrophoresed 5% polyacrylamide/ 8M Urea gel prepared from 40% A: B (19:1), TEMED and APS (Ammonium per sulfate). The gel was run behind a β-blocker screen.

As control, pTRI-Actin was also transcribed with T7 polymerase and loaded in one well, and, expected size of its transcripts is 304 bases.
Finally, the gel was covered with a saran wrap carefully and exposed to phosphor imager screen for about 1 minute and scanned. Bands were subsequently excised from the gel. The eluted RNA in 30 µl Elution Buffer was kept in 1.5 µl micro-centrifuge tubes with caps sealed with parafilm and then placed on a rotator for 3 hours. The supernatants were collected in fresh 1.5 µl micro-centrifuge tube, with 900 µl ethanol and 40 µg glycogen (Roche, IN) in each. Overnight precipitated RNA was taken out from -20 °C, centrifuged at 13000 rpm for 20 minutes. Pellets were re-suspended in nuclease-free H2O and stored at -20 °C in β –blocker, until further use.

2.8. GST-FP (GST-Fusion Protein) Expression

2X YT media was prepared with 16 g Tryptone, 10 g yeast extract, 5 g NaCl in 900 ml distilled water and pH was adjusted to 7.0 with NaOH. Total volume of 1 liter of 2xYT was sterilized by autoclaving for 20 minutes. On cooling, 1 ml of 100 mg/ml Ampicillin stock solution (final concentration 100 µg/ml) was added before use. pGEX-6p1 vector containing *E. coli* strain BL-21 was grown overnight in mini cultures of 5 ml of 2xYTA media. On following day, 5 ml overnight culture was transferred to a 250 ml flask with ampicillin added. The culture was kept in incubator shaker at 37 °C with 250 rpm until A600 reached nearly 0.8, indicating exponential growth. 10 ml of the culture was now segregated from the 250 ml culture.

To the 240 ml culture, 5 µl of 1M IPTG/50 ml culture was added. Induction of lactose analog, IPTG (Isopropyl β-D Thio-Galactoside) was expected to be observed by GST-FPs (Glutathione S-Transferase Fusion Protein) expression. 240 ml culture (with IPTG in a 250 ml flask) and the 10 ml culture (without any IPTG) were put back to incubator shaker at 37 °C with 250 rpm.

At 3 hours of post-IPTG induction, cultures were
separately centrifuged at 2000x g for 5 minutes and pelleted. Laemelli Buffer (SDS reducing buffer) with β-mercaptoethanol was added to the pellets. Resulting solutions were heated in a beaker of boiling water for 5 minutes and immediately chilled for 2 minutes on ice, and, centrifuged at 13000x g for 20 minutes at 4°C. Supernatants were rescued from the viscous pellets and bromophenol blue was added before loading the wells of a SDS-PAGE gel with 6% stacking gel and 13% resolving gel.

A BioRad Precision Plus Protein™ Kaleidoscope Marker was loaded on one well. The proteins were electrophoresed at ~200 Volts for about 45 minutes till the bromophenol blue dye travels to the bottom of the resolving gel. The gel was stained in Coomassie blue dye (R-250) and de-stained to detect the induced expression of fusion protein.

2.9. GST-FP (GST-Fusion Protein) Purification

Using ~ 250 ml bacterial culture (O.D. = 1.5-3.0 at A_{600}), pelleted the bacterial cells by centrifugation and supernatant was removed. The protein purification was performed by using B-PER GST-Fusion Protein purification Kit (Thermo Fischer Scientific, PA).

Cell pellet was suspended in 10 ml of B-PER reagent by pipetting up and down to make a homogeneous cell suspension. The mixture was shaken at room temperature for 10 minutes. To separate soluble proteins from insoluble ones and cell debris, the mixture was centrifuged at 14,000 rpm with Beckman JA17 rotor for 20 minutes. The protein extracts containing about ~90 % of soluble proteins in the supernatant was transferred to a new tube. A second extraction of the pellet with additional B-PER reagent was done to recover remaining soluble protein. The
immobilized glutathione was swirled to make homogeneous gel slurry. 1 ml of the slurry was pipetted to tube containing protein extract. The contents in tube were shaken for 10 minutes at room temperature to allow binding, followed by centrifugation at 2500x g for 5 minutes. The supernatant was discarded and gel was re-suspended with 0.25 ml of wash buffer. The gel slurry of ~ 0.75 ml was transferred to a column fitted to a collection tube and centrifuged at 2000x g for 2 minutes and the column was transferred to a new collection tube. The non-bound proteins were incubated for 5 minutes with 0.5 ml of wash buffer and washed away by centrifugation at 2000x g for 2 minutes. An elution buffer was prepared from 15 µg reduced glutathione added to 2 ml of wash buffer. The GST-FP was eluted by incubation for 5 minutes with 0.5 ml of the prepared elution buffer and subsequent centrifugation at 2000x g for 2 minutes. The spin column was transferred to a new collection tube.

A total of 4 elute fractions were taken and purified elutes of protein were stored in 4 °C until future use. The purified protein was checked for quality on 13 % SDS-PAGE.

2.10. Protein Estimation by Bradford Assay

The determination of concentration of the solubilized protein was done by addition of Bradford Solution (BioRad Laboratories, CA) to the protein solution and subsequent measurement at 595 nm with a spectrophotometer and a relative measure of protein concentration by a standard curve, keeping Bovine Serum Albumin (BSA 1X, 0.1 mg/ml) as standard.

For 500 µl determination, disposable polystyrene micro-
cuvettes were used, samples were mixed with Bradford reagent and waited for 2 minutes and measurement was read thereafter. A plot of O.D. 590 nm versus µg (ug = µg) of BSA standard was made. Triplicates of the sample were made using the following calculation:

<table>
<thead>
<tr>
<th>Bradford (ul)</th>
<th>Sterile Water [ul]</th>
<th>BSA100ug/ml [ul]</th>
<th>Total ug BSA</th>
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<tr>
<td>100</td>
<td>400</td>
<td>0</td>
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<td>100</td>
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<td>100</td>
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<tr>
<td>100</td>
<td>370</td>
<td>30</td>
<td>3</td>
</tr>
<tr>
<td>100</td>
<td>350</td>
<td>50</td>
<td>5</td>
</tr>
<tr>
<td>100</td>
<td>400</td>
<td>X ml sample protein</td>
<td>X</td>
</tr>
</tbody>
</table>

2.11. EMSA (Electrophoretic Mobility Shift Assay)

10x Binding buffer, prepared from NaCl, Tris HCl and RNase free water, was added to 1.5 µl of radiolabelled RNA (~ 20,000 counts). RNA was incubated for 1 minute at 90 °C and transferred to room temperature for 3 minutes. Next, 2x Gel Shift buffer or Incubation buffer, prepared from Tris acetate and Magnesium acetate, was added to the same tube containing RNA. Final volume was adjusted with DEPC water. Now, incubation time given was 7 minutes to initiate RNA folding.

On ice, the purified proteins were kept for quantitation. Then, protein aliquots in different 1.5 µl micro-centrifuge tubes, containing increasing protein concentrations, were thawed at room temperature. Then, NaCl, Na₂HPO₄ and
NaH$_2$PO$_4$ were added to these aliquots. Final volume was adjusted to same in all tubes, by adding DEPC water.

Next, RNA and protein were mixed at room temperature. RNA-protein reaction mixture was given mild orbital shake for 20 minutes, with 1.5 ml micro-centrifuge tube-tops sealed with parafilm. 10 µl of loading buffer, containing glycerol and bromophenol blue and β-mercaptoethanol, was added to make a final loading mixture. One 1.5 µl micro-centrifuge tube will have radiolabelled RNA and all ingredients in it, but the protein, as control for EMSA.

Final reaction mixture tubes were loaded on wells of a 5% polyacrylamide gel, prepared with 40% A: B (19: 1), TBE, MgCl$_2$, APS and TEMED in distilled water. 0.5x TBE was the running buffer. RNA-protein complexes formed resulted in the retarded mobility in the gel and showed the shift. Contrastingly, free RNA migrated faster in the gel during electrophoresis for 2 hours at 100 volts. Gel shift results were analyzed by phosphor imager screen.
CHAPTER III

RESULTS AND DISCUSSION

3.1. Compensatory Mutants of U12 and U6atac snRNAs At Helix I Intermolecular Interaction Region

To study the interactions of the snRNAs during minor class intron splicing, generation of compensatory mutants was required. Mutants could experimentally enable us to study the base-pairing interactions of the snRNAs. Base pairing between U2 and U6 snRNA components were previously demonstrated essential to form helix Ia and Ib as active sites of spliceosomal machinery. So, we mutated functionally analogous sites in U12 and U6atac snRNAs, to test if mutants abrogate splicing (Figure 3.1). The mutants were generated from the expression plasmid backgrounds of U12, U6atac and U4atac snRNAs, using mutagenic oligonucleotides. A schematic of the single nucleotide mutants of U12, U6atac and U4atac snRNAs is illustrated in Figure 3.2.

3.2. Compensatory Mutants affect in vivo U12-dependent Intron Splicing Efficiency

The predicted helix Ia region of U12 snRNA WT spans
between 9-13 nts (nucleotides). Five U12 snRNA mutants generated were named as, A9U, A10U, C11G, U12G and U13A. WT U6atac snRNA presumably uses 25-21 nts to base-pair with U12 snRNA helix Ia region. To test if predicted U12 snRNA interacts with corresponding U6atac snRNA, five compensatory mutants of U6atac snRNA were generated and named as, U25A, U24A, G23C, G22U and A21U (Figure 3.2). Likewise, 63-65 nts of U4atac snRNA Stem I region is predicted to base pair with U6atac snRNA along 25-23nts. So, three compensatory mutants of U4atac snRNA were generated and named as, A63U, A64U and C65G (Figure 3.2). For purpose of simplicity, each nucleotide of interacting U12 snRNA, that has been mutated to create a compensatory mutant, is grouped with the corresponding compensatory mutants of U6atac snRNA and U4atac snRNA. Then, a Set number is assigned to the team of 3 compensatory mutants. All team members grouped within a particular Set is predicted to rescue U12-dependent splicing upon co-transfection of all snRNAs which belong to that Set (Figure 3.2).
For example, U12 A9U mutant is expected to abolish U12-dependent splicing. On co-transfection with U6atac U25A and U4atac A63U mutant, the U12-dependent splicing by U12 A9U mutant is expected to be restored. Then, we can conclude that, the restoration of correct U12-dependent splicing to the U12 snRNA mutant is possible by compensatory mutant of U6atac snRNA and compensatory mutant of U4atac snRNA.

This is the strategy of proposed compensatory sequential base-pairing alterations. It allows us to test the effect of the alteration of one nucleotide in U12, U6atac and U4atac snRNAs on their function, in vivo. It will also prove if one nucleotide mutation in U12 snRNA is enough to impair splicing. Moreover, if splicing gets restored, when all compensatory mutants are put together, we will conclude that, these three snRNAs cross-talk with one another at their respective mutated nucleotide position (Figure 3.2 C: Table).

![Figure 3.2: U4atac/U6atac snRNAs base-pairing in schematic representation.](image)

- **Predicted base-pairing between U4atac and U6atac snRNAs prior to U6atac/U12 snRNAs base-pairing at Helix 1a interaction region.**
- **Construction of mutants of U12, U6atac and U4atac snRNAs at Helix 1a interaction region.**

![Table 3.2 (C): Tabular representation of the base-pairing of U12, U4atac and U6atac snRNAs at helix 1a interaction region along with their expected outcomes.](image)
3.3. Compensatory Mutants of U12 and U6atac snRNAs At Helix Ib Intermolecular Interaction Region

The predicted helix Ib region of U12 snRNA WT spans from 4-6 nts (nucleotides) (Figure 3.3). Three mutants generated from single nt. substitution at position 5 were named as, C5A, C5U and C5G. Three more mutants were generated from single nt. substitution at position 6 and named as, U6A, U6G and U6C. WT U6atac snRNA presumably uses 28-26 nts to base-pair with U12 snRNA helix Ib region. So, three compensatory mutants of U6atac snRNA were generated from single nt. substitution at position 27 and named as, G27U, G27C and G27A. Three more mutants generated from single nt. substitution at position 26 were named as, A26U, A26C and A26G. Likewise, 60-62 nts of U4atac snRNA Stem I region is predicted to base pair along 28-26 nts. of U6atac snRNA. So, three compensatory mutants of U4atac snRNA were generated from single nt. substitution at nt. position 61, and named as, C61A, C61G and C61U. Three more mutants were generated from single nt. substitution at position 62, and named as, U62A, U62G and U62C. A schematic representation (Figure 3.3) illustrates predicted base pairing of U12, U6atac and U4atac snRNAs at helix Ib. As done with helix Ia, likewise, a Set number was assigned to each team of interacting U12, U6atac and U4atac snRNAs compensatory mutants of helix Ib region. All team members grouped within a particular Set is predicted to rescue U12-dependent splicing upon co-transfection of all components belonging to that Set.

3.4. In vivo Genetic Suppression Assay Using P120 Mini-gene Intron F Can Detect Predicted Base Pairing of U12-U6atac and U6atac-U4atac snRNAs

Controlling the spliceosomal machinery by nucleotide base-pairing interaction within the helix region could be an effective cellular strategy to
Regulate gene expression. The in vivo requirements for the nucleotide base pairing of U6atac, U12 and U4atac snRNAs for catalytic reactions during minor class intron splicing can be observed by the phenotypic expression of splice variants by performing the in vivo genetic suppression assay, where the P120 minigene with U12-dependent intron F was utilized (Figure 3.4).

To analyze the splicing of U12-dependent introns in vivo, Hall and Padgett expressed a fragment of the human P120 gene containing a 99 nucleotides long AT-AC intron in CHO cells, using a transient transfection assay. The transiently expressed AT-AC intron is correctly spliced in the transfected CHO cells. Suppression of a loss-of-function mutation in one molecule of a proposed base pairing RNA interaction by introduction of compensatory mutations in the other proposed RNA interacting molecule is the principle of the in vivo genetic suppression assay.

Our assay is similarly designed to study the existence of RNA-RNA base-pairing in vivo and also to detect the splice sites of the pre-mRNA. The co-expression of U11 Wild Type (Wt) snRNA with P120 Wt and U6atac snRNA Wt will
generate a single band product as fully spliced. The co-expression of suppressor allele of U11 snRNA carrying the GG6/7 CC compensatory mutation to base-pair with P120 5’ss mutation will eliminate splicing and will generate a single band as unspliced product. Only, the co-expression of suppressor allele of U11 snRNA carrying the GG6/7 CC compensatory mutation to base-pair with P120 5’ss mutation activate splicing in vivo when co-transfected with U6atac mutant suppressor allele carrying the GG 14/15 CC compensatory mutation and will restore the spliced band to almost WT level (Figure 3.4C).

Similarly, co-expression of suppressor allele of U11 snRNA carrying the GG6/7 CC compensatory mutation to base-pair with P120 Double Mutant with a 5’ss mutation as well as a branch point sequence mutation, will eliminate splicing and will generate a single band as unspliced product (Figure 3.4C). Only, the co-expression of suppressor allele of U11 snRNA carrying the GG6/7 CC compensatory mutation to base-pair with P120 5’ss mutation activate splicing in vivo when co-transfected with U6atac mutant suppressor allele carrying the GG 14/15 CC compensatory mutation and the U12 suppressor mutant snRNA and will restore the spliced band to almost WT level.

Using the in vivo genetic suppression principle, our complementary mutants were grouped in sets and were co-transfected in Hela cells and were analyzed on 3% agarose gel, after RNA isolation followed by RT-PCR of the respective cDNAs, and, subsequent nested PCR steps. Optimization of the transient transfection conditions was performed to minimize non-specific products in each experiment. The results were verified by repeating the experiment, each time using appropriate controls in separate well on the gel. Along with the final in vivo products, the DNA ladder (100bp marker) is also loaded in one well to check the spliced product size.
3.5. Single Nucleotide Mutants of U12, U6atac and U4atac snRNAs At Helix Ia Interaction Region Could Not Prevent Splicing

Splicing levels (Figures 3.5 and 3.6) indicate that single nucleotide mutation could not limit the activity of the U12-type splicing by inhibiting the base pair interaction at helix 1a region. Plenty of in vivo spliced products were detected using single complementary mutant, when negligible or no splicing was our expected outcome. The efficiency of U12-type of splicing was almost equivalent to that of normal splicing in vivo by substitution mutants on transfected constructs.

3.6. Splicing Efficiency Is Almost Unaltered With Single Nucleotide Mutation in Helix Ib Region of Interaction

Minimal or insignificant change was observed in the amount of splicing levels from the single nucleotide mutants in helix Ib region, as compared to the fully spliced mRNA.

This establishes the fact that, single nucleotide mutation was not enough to inhibit splicing (Figures 3.6 and 3.7).
3.7. Analyses of Single Mutations From in vivo Spliced Phenotypes

Our data show that base-pairing interactions occur between
Figure 3.7: In vivo single nucleotide splice variants of helix 1b region –interacting snRNAs (SETS 6 and 7): After reverse transcription and PCR amplification, the products were analyzed by gel electrophoresis on a 3% agarose gel with 1X TBE.

Figure 3.8: In vivo single nucleotide splice variants of helix 1b region –interacting snRNAs (SETS 8, 9, 10 and 11): After reverse transcription and PCR amplification, the products were analyzed by gel electrophoresis on a 3% agarose gel with 1X TBE.

U12 and U6atac snRNAs (Figures 3.5, 3.6, 3.7 and 3.8). Splicing could not be
completely eliminated with mutation of a single nucleotide at helix interaction region of any one of the participating snRNAs. The \textit{in vivo} spliced phenotypes indicate that, the mutation of single nucleotide is not enough to abrogate splicing. None of the sets of compensatory mutants of U12, U6atac and U4atac snRNAs could completely eliminate splicing. Instead, most of the \textit{in vivo} phenotypes show that single nucleotide mutation was tolerated with little or no splicing in each Set.

As expected, no band was observed in case of negative control (mock transfection). Wild type lane had completely spliced product. The co-expression of suppressor allele of U11 snRNA carrying the GG6/7 CC compensatory mutation, P120 Double mutant, U6atac mutant suppressor allele carrying the GG 14/15 CC compensatory mutation and U12 suppressor mutant snRNA, activated splicing \textit{in vivo} and restored the spliced band to almost WT lane.

From the positive controls in the experiment, we could conclude that our \textit{in vivo} splicing assay has worked. However, there was no significant change visible on the splicing efficiency in all Sets of single nucleotide mutant co-transfected. Unlike the \textit{in vitro} splicing assay, where all intermediate stages and products of splicing could be resolved on same gel, using \textit{in vivo} splicing system could only allow us to obtain information only from the final products of splicing. Moreover, the splicing reaction kinetics could also be studied and different complexes formed during splicing could also be identified by the \textit{in vitro} splicing assay. So, an \textit{in vitro} splicing assay is recommended which could reflect the situation observed \textit{in vivo}.

Our results also suggest us to choose adjacently two nucleotide mutations in the interacting snRNAs and check if there is any obstruction of the splicing event with double nucleotide mutations in the region of the base-pairing interactions in helix Ia and Ib. Since RNA-RNA interactions are phylogenetically
conserved and are essential for splicing to occur, we hypothesize that, adjacent two nucleotide mutations in the interacting snRNAs might be sufficient to disrupt the Watson – Crick base pairing near helix region and therefore, will prevent splicing.

3.8. Double Nucleotide Mutations of U4atac, U6atac and U12 snRNAs In Helix Ia Region Inhibit in vivo U12-dependent Splicing

To study the events of snRNAs during minor class intron slicing, it was necessary to recapitulate the splicing phenomenon by using adjacent two nucleotide mutations in the interacting snRNAs. Then, restoration of splicing in spite of adjacently two nucleotide changes will indicate that the mutation in interacting nucleotides is enough to stop splicing catalysis. On other hand, if splicing gets restored, when all compensatory mutants will be put together, we can conclude that, these three snRNAs cross-talk with one another at their respective mutated nucleotides positions.

The predicted helix Ia region of U12 snRNA WT spans from 9-13 nts (nucleotides), and, four mutants generated were named as, AA9/10UU, AC10/11UU, CU11/12GG, and UU12/13GA. WT U6atac snRNA presumably uses 25-21 nts to base-pair with U12 snRNA helix Ia region. So, four compensatory mutants of U6atac snRNA were generated and named as, UU24/25AA, GU23/24CA, GG22/23UC and AG21/22UU. Likewise, 63-65 nts of U4atac snRNA Stem I region is predicted to base pair with U6atac snRNA along 25-23nts. So, two compensatory mutants of U4atac snRNA were generated and named as, AC64/65UG and AA63/64UU. A schematic representation (Figure 3.9) of U12, U6atac and U4atac snRNAs illustrate their predicted base pairing. A Set Number was assigned to each group of interacting U12, U6atac and
U4atac snRNAs compensatory mutants. All mutants grouped within a particular Set are predicted to rescue U12-dependent splicing upon co-transfection of all components that belong to that Set.

3.9. Analyses of in vivo Splicing Phenotypes from Double Mutations of U12, U6atac and U4atac snRNAs In Helix Interacting Region

One snRNA component bearing mutation in two adjacent nucleotides in the helix region of interaction was sufficient enough to inhibit splicing of a transfected construct. U12-type splicing does not occur if the in vivo environment is deficient of any one of the snRNA bearing at least two adjacent nucleotides mutation at the helix interaction region. However, the efficiency of splicing was significantly noticeable on co-transfection of all double nucleotide compensatory mutants on transfected constructs in vivo (Figure 3.10).

Our results could provide information to understand the structure – functional relationships in the interacting snRNAs at the helix region. The
base-pairing of, particularly, U12 and U6atac snRNAs are found to be more central to the \textit{in vivo} splicing reactions. Although all the base pairs are important for function, the identities of the bases may not be so important, because the complementary sequences between U12 and U6atac and U4atac snRNAs could execute splicing reaction in an analogous fashion as performed by the WTs of U12 and U6atac and U4atac snRNAs.

3.10. Different snRNA Elements Might Share Common Minor Spliceosome Specific Components during U12-dependent Intron Splicing

Recent studies have suggested that, apart from the helix region, and with the pre-mRNA, U6atac and U12 snRNAs interact extensively with each other by Watson-Crick base pairing. Previous findings in our lab, using an \textit{in vivo} genetic suppression assay, have identified that the Intra-molecular Stem Loop (ISL) structure of U6 snRNA can functionally replace U6atac snRNA ISL. Moreover, the 3’
RNA element of U6atac snRNA has been proved to be sufficient enough to guide U6 snRNA to minor spliceosome, indicating that, only 3’ stem-loop (SL) of U6atac snRNA region is capable enough to activate minor class intron splicing (Shukla and Padgett, 2009).

From this information, we can presume that, RNA substructure(s) within the 3’ RNA element of U6atac snRNA may have functional significance as critical component(s) which recruit spliceosomal components specifically to U12-dependent introns. Hence, disruption of such a critical RNA element would severely impair minor class intron splicing.

Summarizing the important break-through discoveries mentioned above, it is evident that, U6atac snRNA might be a good candidate to explain its interaction with some minor spliceosome-specific factors, most likely the components of U11/U12 di-snRNP and U4atac/ U5/U6atac tri-snRNPs, which must be required to bind the U12-type intron and prevent the minor spliceosome from being interfered by major spliceosome.

Recently, we have identified one potent critical substructure at 3’ end of U6atac ISL region, between nucleotides 91 to 109, whose sequence alignment corresponds to U12 snRNA SLIII (nts. 109-125) (Figure 3.11). Furthermore, it is known that, human 18S U11/U12 snRNP exists as a di-snRNP and contains a set of seven proteins that are not found in U2 snRNP, suggesting that, the recognition of 5’ splice and branch sites of U12-dependent introns is distinct from U2-dependent introns. Of these seven novel proteins, 65K RNA binding protein of U11-U12 di-snRNP complex is essential for cell viability (Will et al. 1999, Will et al. 2004) and forms a bridge between U11-59K protein and U12 snRNAs. It is also known to contain two RNA Recognition Motifs (RRMs) and is presumed to be involved in di-snRNP formation and
intron bridging. At this point, we also know that, the minor spliceosome tri-snRNP U4atac/U6atac.U5 forms similarly like major tri-snRNPs (Tarn and Steitz 1996a, Tarn and Steitz, 1996b). So, protein components of the U6atac/U4atac.U5 tri-snRNP do not differ from those in the major tri-snRNP (Luo et al, 1999, Nottrott et al, 2002, Schneider et al, 2002). On the contrary, U11 and U12 do not act as mono-particle but act as a di-snRNP. Since 65K protein of U11/U12 di-snRNP is only involved in minor splicing, so it is important to investigate that 65K protein might play a significant role during in recognizing and pairing of splice site and ensure the dynamicity as well as conservation of the minor splicing event.

65K RNA-BP appears to bind to apical stem–loop region of U12 SLIII and which has identical nucleotide sequences with U6atac ISL region, between nucleotides 91 to 109. Hence, we focused on two parameters: U12 snRNA SLIII nts (109-125) corresponding to 65K-C-RRM protein binding region, and sequence similarity of U6atac 3’ element (nts. 91-109) with U12 snRNA SLIII (nts. 109-125).

We envision that sequence in U6atac snRNA identical to U12 snRNA might accomplish an important strategy involving the 65K protein, during a proper assembly of minor spliceosomal complex and a productive minor class intron splicing. The U6atac snRNA nucleotide sequences, from 91 to 109, within the 3’ element might, therefore, be one critical sub-structure which could be indispensable for catalytically active spliceosomal rearrangements during snRNP assembly of U12-dependent intron splicing.

Since it is known that the 65K-C-RRM contains a well characterized domain binding with Stem III of U12 snRNA, we checked and confirmed that the 65K-C-RRM binds to 3’ element of U6atac snRNA. The human U11/U12-65K protein contains one N- and one C-terminal RRM (RNA Recognition Motif) and a
central proline-rich region (Figure 3.12). It has been established by electrophoretic mobility shift assays (EMSA), that the GST-65K-C-RRM interacts with U12 snRNA (Benecke et al, 2005).

In order to check if the predicted interaction of GST-65K protein exists with U6atac snRNA, we used purified GST-65K-C-RRM and GST-65K-N-RRM and preformed EMSA with $^{32}$P* radiolabelled-U6atac WT snRNA. Our EMSA results validated that only 65K-C-RRM and not 65K-N-RRM could bind to U6atac snRNA. We can therefore predict that, the 3’ RNA element of U6atac snRNA with the stem-loop structure, as well as any or multiple loop nucleotides and the loop-closing base-pair, play a critical role in the 65K/U6atac 3’SL interaction, and therefore can consequently affect splicing.

**Figure 3.11: Proposed secondary structure of U6atac snRNA:**

3.11(A): U6atac snRNA secondary structure with nucleotides (in red circle) in the 3’ element identical to that in U12 snRNA Stem Loop III.

3.11(B): Proposed secondary structure of U12 snRNA showing nucleotides (109 to 125) identical to U6atac snRNA secondary structure from nucleotides circled in red.

3.11(C): U6atac snRNA has nucleotides from 91 to 109 in the 3’ stem loop element with sequence alignment identical to U12 snRNA Stem Loop III nucleotides 109 to 125.
3.11. Mutations Generated in U6atac snRNA 3’ SL between Nucleotides 90 to 110 Can Detect Correct 65K-C-RRM Binding Sites

Once our EMSA data validated that the 65K-C-RRM binds to U6atac 3’S-L, hence we decided to detect which nucleotide(s) or their alignment in U6atac snRNA are critical for binding to 65K-C-RRM protein. After making mutations within the nucleotide region 90 to 110 in U6atac snRNA 3’ element, further EMSA analysis will help us detect which nucleotide or nucleotides comprising the loop or stem of the U6atac snRNA between the 90 to 110 positions are recognized by the 65K-C-RRM, or which nucleotides are crucial to contribute to 65K binding affinity.

To check the relative importance of the nucleotide region 90 to 110 in U6atac snRNA 3’ element, we generated a pool of mutants of U6atac
snRNA, namely L1, L2, L3 and S1, S2, S3, S4, S5 and S6 (Figure 3.13).

L1, L2, L3 are the loop mutants and S1, S2, S3, S4, S5 and S6 are the stem mutants. Mutations of conserved loop nucleotides U (98), A(99) and C (100) were done to G (98), C(99) and G(100) respectively, by exchanging a pyrimidine with a purine and *vice versa* and three loop mutants U98G, A99C and C(100)G so generated, were named L1, L2 and L3, respectively.

The conserved C-G loop closing base-pair was mutated to U-A, and named as S1 (Stem mutant 1). By disrupting the stem region from nucleotides 104 to 109, Stem mutant 2 or S2 was generated to check if a stem-loop structure and not a single-stranded RNA alone are required for binding. The entire sequence of the stem was altered in Stem mutant 3 (S3). The orientation of the loop-closing C-G base pair was changed to G-C base pair to generate Stem mutant 4 or S4. All the stem nucleotides with the exception of the C-G loop-closing base pair were mutated to generate Stem mutant 5 or S5. The Stem mutant 6 was generated by disrupting the stem region from nucleotides 91 to 96, to check if a stem-loop structure and not a single-stranded RNA alone are required for binding (Figure 3.13).

**3.12. Expected Outcomes from The U6atac snRNA (90-110) Nucleotides Mutations On 65K-C-RRM Protein Binding**

The purpose of making the mutations in the U6atac snRNA is to investigate if the stem, loop, or stem-loop structure, as well as, the identity of the several of the loop nucleotides and the loop-closing base-pair, plays a critical role in U12-dependent splicing. Mutations of conserved loop nucleotides, U (98), A(99) and C (100) in loop mutants, L1, L2 and L3, respectively are expected to abolish 65K-C-RRM-
RNA complex formation.

Conversion of the conserved C-G loop closing base-pair was mutated to U-A, in Stem mutant 1 is expected to show severe reduction in complex formation. Disruption of the stem in Stem mutant 2 and Stem mutant 6 will abolish any complex formation, because a defined stem loop structure and just not a single stranded RNA alone is probably not binding to 65K-C-RRM. Changing the orientation of the loop closing base pair, as done with Stem mutant 4, is presumed to have a moderate effect on protein binding, because the other structures remain unaltered. Finally, altering the entire sequence of the stem, which is done to generate Stem mutant 3, will leave the RNA with no protein binding sites. 65K-C-RRM will fail to recognize any sequence for RNA recognition and so, we expect that S5 will completely abolish 65K-C-RRM binding.

In summary, the mutations were designed to test the requirement of loops as well as stem of the predicted U6atac-65K-C-RRM interacting region in U12-dependent splicing.

3.13. **U6atac snRNA Stem and Loop Nucleotides (90-110) Are Important In U12-dependent in vivo Splicing**

Under normal splicing conditions, minor class intron splicing will be unchallenged with involvement of WT U6atac snRNA interaction during the spliceosomal assembly reactions. We have identified the involvement of U6atac snRNA during *in vivo* genetic suppression assay for minor class intron splicing. However, the predicted critical substructure at 3’ end of U6atac snRNA ISL region, between nucleotides 90 to 110, whose sequence alignment corresponds to U12 snRNA
SLIII (nts. 109-125), might have a crucial role in splicing. At this point, we also know that of various unique, novel proteins of minor spliceosome, protein 65K-C-RRM contains RRM (RNA Recognition Motif), a well-characterized domain binding with Stem III of U12 snRNA. Focusing on these two parameters, we generated stem and loop mutations in U6atac snRNA 3’ element (nts. 91-109). Then, we wanted to analyze the effect of U6atac snRNA stem and loop mutants by \textit{in vivo} genetic suppression assays.

We wanted to check the U6atac mutation effect on U12-dependent splicing on a background of P120 mutant at 5’ splice site (ss) only. We know that, U12snRNA is base-paired to BPS and U11 snRNA is based to 5’ss, but simultaneous recognition of 5’ss and BPs is required for minor class intron splicing. Hence, recruitment of U11 and U12 occurs as a di-snRNP complex, when 65K protein is a potent regulator of minor class intron splicing for its recognition of this di-snRNP
complex. We would like to investigate if mutants within the 3’ U6atac snRNA element (nts.91-109) determine the levels of U12-dependent splicing. *In vivo* genetic suppression assay was performed using loop and stem mutants constructed in GG14/15CC background and the U11 suppressor and U12 suppressors were added accordingly as compensatory mutants to rescue splicing.

### 3.14. Analyses of *in vivo* Splicing Using U6atac Stem and Loop Mutants with P120 Mutant At 5’ Splice Site

In background of P120 5’splice site mutation, the stem and loop mutations in 3’ U6atac snRNA element (nts.91-109) showed differential effects on splicing (*Figure 3.14*).

The 0 lane was loaded with 100bp DNA ladder to check the spliced and unspliced product sizes. As expected, no band was observed in case of mock transfection in lane 1 and in empty vector in lane 2. Wild type product in lane 3 had the expected band size as completely spliced product at 100 bp. P120 with 5’ splice site mutation generated unspliced product in Lane 4. Co-expression of U11 snRNA carrying the GG6/7 CC compensatory mutation and P120 with 5’ splice site mutation generated unspliced product in Lane 5. Co-expression of U6 snRNA carrying the GG14/15CC compensatory mutation and P120 with 5’ splice site mutation generated less unspliced product and more of spliced product in Lane 6. Lane 7 showed that co-expression of all three suppressors, activated splicing *in vivo* and restored the spliced band to almost WT level, as in Lane 7.

Mutations of conserved loop nucleotides, U (98), A(99) and C (100) in loop mutants, L1, L2 and L3, did not abolish splicing in Lanes 8,9 and 10 respectively.
In Stem mutant 1, where conserved C-G loop closing base-pair was mutated to U-A, there was reduction in spliced product in Lane 11. Disruption of the stem in Stem mutant 2 abolished splicing, as in Lane 12, probably because a defined stem loop structure and just not a single stranded RNA alone is required for splicing. Stem mutant 3 with entire sequence of the stem altered, show both spliced and unspliced products in Lane 13. Changing orientation of the loop closing base pair, as done with Stem mutant 4, showed more unspliced product in Lane 14. With all mutated stem nucleotides with the exception of C-G loop-closing base pair in Stem mutant 5, showed both spliced and unspliced products. Disruption of the stem in Stem mutant 6 abolished splicing, as in Lane 16.
CHAPTER IV

CONCLUSIONS AND FUTURE DIRECTIONS

4.1. Conclusions

Using the single and double nucleotide mutations in snRNAs, we were able to utilize CHO cell-based transient transfection system to study pre-mRNA splicing and the U12-dependent minor class intron splicing, in specific. Our results showed that U12/U6atac helix Ia interaction ensured splicing, when adjacently two nucleotides complementary mutants of U12 snRNA and U6atac snRNA were co-expressed. This result could find resemblance with the U2/U6 helix Ia interaction, that exists in major class or U2-dependent intron splicing. U12/U6atac Helix Ib is predicted to have a structurally conserved AGC sequence situated before the U6atac Intra-molecular stem loop (ISL) region, almost identical to U6 ISL. However, single nucleotide mutations of U12, U4atac and U6atac snRNAs in helix region were not sufficient to prevent U12-dependent minor splicing. But, it could be well inferred from our data that, U12/ U6atac interaction at helix region was severely affected with double nucleotide mutants. Furthermore, 3’ element of U6atac snRNA from nucleotides 91 to 109 has sequence similarity with Stem Loop III of U12snRNA from nucleotides 109 to
125. Of various unique novel proteins of minor spliceosome machinery, protein 65K-C-RRM contains a well-characterized domain binding to U12 snRNA SL III nts (109-125). We performed EMSA with 65K-N-RRM, 59K-N-RRM and 59K-C-RRM with U6atac snRNA WT, and, the gel shift assays validated that, only 65K-C-RRM binds to U6atac snRNA WT. Our data confirmed that U6atac 3’ element nts. (91-109) binds to protein 65K-C-RRM.

Each nucleotide between positions 91 to 109 was mutated to generate single nucleotide mutants of U6atac snRNA. U12-dependent splicing was detected in varying levels for all mutants, suggesting that U6atac snRNA 3’ element nts. (91-109) region is an essentially required component for U12-dependent intron splicing.

In conclusion, minor class or U12-dependent intron splicing is regulated by interplay of snRNPs following several unexplored mechanisms. In our work, regulatory mechanisms of U12, U11, U6atac and U4atac snRNAs were outlined. Further analyses of structure-function aspect of minor class U12-dependent intron splicing could significantly improve our understanding about their regulation in gene expression, coupled with their clinical significance, linked to the fact that, U12-dependent intron splicing co-exist with U2-dependent intron splicing.

4.2. Future Directions

We want to determine splicing intermediates and detect free 5’ exon, lariat intermediate with 3’ exon, free lariat intermediate and spliced exons on the gel. In vitro assay will also ensure us to study the kinetics of splicing reaction, by manipulating the in vitro reactions at different time points, during complex formation steps. Furthermore, in vitro approach will also lead us to differentiate whether splicing
occurs by U2-dependent- or U12-dependent path, and, could provide insights about the coupling between the two. Inactivation of the snRNAs will be done using 2’ O-methyl oligos. For blocking U2-dependent splicing, snRNAs in U2-system will be inactivated in the nuclear extracts. Since U2-system is manifold abundant than U12-system, we have to inactivate snRNAs of both U2- and U12- systems, in order to study what happens on blocking the U12 splicing components.

The only drawback with in vitro method is that, when almost every construct transfected in cells splices upon transfection in vivo, splicing ability is often limited for in vitro assays.

For assay validation and quantitation of gene expression, we will approach Quantitative Real-Time PCR (qRT-PCR) method for quantification and for studying kinetics of the reaction in early phases of PCR. Whereas, ethidium bromide staining is not precisely quantitative, the qRT-PCR will be highly sensitive to detect PCR amplicons at their exponential phase of the reaction. Hence, an accurate measurement of RNA will be obtained in an easier and more precise way. Once the expression levels and splicing efficiency can be quantitated, the SPSS package can be used for statistical analyses of each snRNA interaction and relative splicing efficiency can be calculated.

In vitro reconstitution study can be another approach to monitor the effects of reconstitution of functional snRNPs involved in U12-dependent intron splicing. By using a synthetic snRNA for in vitro run off transcription and subsequent incubation of the synthetic snRNA in HeLa cell extracts inactivated by pre-treatment, will give us information about the functional significance of a particular snRNA during U12-dependent intron splicing, which can be calculated from restoration of the splicing efficiency during in vitro assembly, in presence of ATP.
For improving the RNA-protein interaction, our GST-FP (65K-C-RRM) produced from pGEX-6P-1 vector with a PreScission Protease recognition site, can be cleaved from the GST tag either while bound to Glutathione sepharose chromatography media or in solution after elution. This will ensure stability of the target protein while simultaneously maximizing its interaction with its substrate RNA.

The interactions of stem and loop mutants of U6atac 3’ RNA element with the U12 intron on a background of P120 DM bearing mutations, both at 5’ss and branch site, could also be studied along with their interactions with 65K-C-RRM protein.


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APPENDIX
List of U11 Mutants generated: All primers are in 5’ to 3’ direction:

U11-1: (DNA)-Phos-5’ – AAATCGGGAACTGTCGTGAG-3’

U11-2: (DNA)-Phos-5’ –GAGATTTCCGTTCCATAATTTTT-3’

U11-3: (DNA)-Phos-5’ –TTCGGCTTCTCGTGGAGTGGCAC-3’

U11-4: (DNA)-Phos-5’ –CTGTCGTGAGGCTGACCGTACGAGG-3’