Regulated pre-mRNA splicing: the ghostwriter of the eukaryotic genome

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Abstract
Intron removal is at the heart of mRNA synthesis. It is mediated by one of the cell's largest complexes, the spliceosome. Yet, the fundamental chemistry involved is simple. In this review we will address how the spliceosome acts in diverse ways to optimize gene expression in order to meet the cell's needs. This is done largely by regulating the splicing of key transcripts encoding products that control gene expression pathways. This widespread role is evident even in the yeast Saccharomyces cerevisiae, where introns appear to have been lost; yet how this control is being achieved is known only in a few cases. Here we explore the relevant examples and posit hypotheses whereby regulated splicing fine-tunes gene expression pathways to maintain cell homeostasis.

Keywords
spliceosome; Yra1; Sus1; L30; Mer1; export; histone modification

Introduction
In general terms, prokaryotic genes are compact, can be grouped based on their functions, and are translated while still being transcribed. Gene expression in prokaryotes has evolved towards speed and efficiency. In contrast, most eukaryotic genes are isolated and, moreover, have their sequences interrupted, such that the coding pieces must be spliced together to produce a meaningful transcript or mRNA. Transcripts undergo further processing before being transported to the cytoplasm. Accordingly, mRNAs exist in the cell as RNPs (ribonucleoprotein) particles that act as both executors and bearers of their fates. In light of this complex mRNA processing, it is clear that eukaryotic gene expression is under a set of selection rules whereby speed and efficiency are only part of the story.

This review will focus on the diverse roles that pre-mRNA splicing plays in the control of the processes underlying gene expression. Elegant genetic, biochemical, and cell biological studies in the yeast Saccharomyces cerevisiae have made it possible to glean mechanistic insights into these processes and their interplay. As such, these will be the basis of this review. Nonetheless, many of the guiding principles learned from S. cerevisiae are likely to
inform our general understanding of how regulation of splicing and the integrated gene expression networks that involve splicing contribute to the cellular homeostasis.

pre-mRNA splicing

Pre-mRNA splicing is the process by which the coding segments of a gene (exons) are spliced together, and the interrupting fragments (introns) are removed and discarded. Splicing is the task of the spliceosome, one of the most complex macromolecular machines in the cell [1]. This evolutionarily conserved assembly starts anew in each round of splicing (reviewed in [2]). First, once the intronic 5’ end (5’ splice site or 5’ss) is transcribed, it is recognized by the U1 snRNP (small nuclear RNP). This interaction is thought to be driven by the complementarity between the U1 snRNA and the 5’ss, although its protein components play important roles as well [3]. Subsequently, the 3’end of the intron, including the branch-site (BS) and the 3’ splice site (3’ss) are identified by the factors BBP and Mud2 in yeast or their homologs SF1 and U2AF in other systems (including human). These factors play a role in the initial recognition of the BS by the U2 snRNP. Early interactions between the U1 and U2 snRNPs on the same intron are thought to be essential to define the intron to be removed and, therefore, the exons to be joined. There is an alternative way to define the substrate for the spliceosome, based on “exon definition,” which posits that interactions between an upstream U2 snRNP and a downstream U1 snRNP are critical for the further assembly of the nascent spliceosome, marking the pre-mRNA in between as an exon. This appears to be particularly important for genes that contain multiple, large introns, as is usually the case in metazoans [4] As we will further examine, recent studies in *Saccharomyces cerevisiae* support the notion that cross-exon interactions are also important in yeast.

In a subsequent step, which remains somewhat obscure, molecular rearrangements lead to the interaction between U1 and U2 of the same intron (as in the previous “intron definition”), and spliceosome assembly proceeds with recruitment of the U4/U6.U5 tri snRNP. Subsequent remodeling steps, catalyzed by RNA-driven ATPases, displace U1 and U4 and facilitate the formation of the catalytically active spliceosome, which will form a lariat structure via a 2’-5’ phosphodiester linkage between the adenosine at the BS and the 5’ end of the intron, followed by exon joining and intron removal. Thus, a sophisticated molecular ballet culminates in two chemically simple transesterifications. That this rather straightforward chemistry involves such a complex cellular machinery is a testament to the relevance of spliceosome dynamics. By modulating spliceosomal function to produce a variety of splicing outcomes from a single pre-mRNA, evolution has developed a formidable tool to enlarge the coding potential of genomes. The initial recognition of the pre-mRNA by U1 and U2 is an important target for regulation. Regulatory proteins can interfere with either the recognition of intronic sequences (leading to intron splicing or retention); or the interactions that occur during exon definition (promoting exon inclusion or skipping). Switching the 3’ss after spliceosome assembly has been shown in metazoans [5], indicating that regulation of the spliceosome goes beyond controlling initial splice site recognition. In some instances the pre-mRNA structure plays a role in intron recognition, and in many yeast introns it is critical for proper 3’ss selection [6, 7]. Pre-mRNA intronic and exonic sequences, known as splicing enhancers or silencers, participate in the recruitment of splicing factors, such as SR proteins and hnRNPs. As is the case with many of the pre-mRNA sequences involved in splicing, the information content of silencers and enhancers is rather low, and the splicing outcome will also depend on the relative concentration of the splicing regulatory proteins. Thus, alternative splicing is a function of the pre-mRNA sequence (including exonic structure and RNA folding) and the concentration of splicing factors, which together make up a putative “splicing code”. Although regulated alternative splicing has been best characterized in metazoans, studies of the yeast *S. cerevisiae,* in
which regulated splicing primarily is seen in the form of intron retention, offers an opportunity to glean basic insights into the reaction.

**Chromatin, transcription, and splicing converge to modulate gene expression**

About 20 years ago, evidence emerged indicating that intron removal could occur on nascent transcripts. Elegant imaging studies by Beyer and co-workers revealed nascent pre-mRNA shortening caused by splicing [8, 9]. Subsequently, a number of important studies using *Saccharomyces cerevisiae* also showed that spliceosome assembly (beginning with intron recognition) occurred on the nascent transcript, although it was less clear whether the actual chemistry of splicing occurred cotranscriptionally [10-14]. Recently, RNA polymerase II has been reported to pause in a manner that facilitates both co-transcriptional assembly of the spliceosome as well as splicing [15, 16], supporting the notion that most splicing occurs co-transcriptionally [16].

In light of this close spatial and temporal relationship between transcription and splicing, it is not surprising that polymerase activity can have important effects on splicing. For example, in mammals as in yeast, the speed of the polymerase influences recognition of splice sites, such that an exon that is normally excluded (exon skipping) is recognized and included when the polymerase rate is slowed [17, 18]. Additionally, in mammals RNA polymerase pausing has been shown to affect exon recognition [19]. A more extensive survey of co-transcriptional splicing mechanisms can be found in a number of excellent recent reviews (see for example [20-22]). Nonetheless, these studies demonstrate a growing appreciation that in order to understand splicing and its regulation, it is important to view splicing within its co-transcriptional context.

More recently, it has become clear that one of the important mechanistic consequences of transcription-coupled splicing is that the state of the chromatin influences splicing outcomes. Analysis of genome-wide surveys from *Drosophila* and mammals have revealed that nucleosomes and, according to several of these studies, specific histone modifications are enriched in exon sequences, suggesting that there may be specific histone “marks” that are associated with splicing [23-29]. Additionally, proteins that bind to histones have been shown to facilitate the recruitment of snRNPs to the nascent transcript and influence efficiency of splicing and alternative splicing (Reviewed in [30, 31]). The ability of chromatin to influence pre-mRNA splicing appears to be a conserved feature of co-transcriptional splicing. For example, Gcn5, the histone acetyltransferase that is part of the SAGA complex, has strong genetic interactions with components of the U2 snRNP, and its catalytic activity is required for co-transcriptional spliceosome assembly [32, 33].

In light of the integral relationship between chromatin and splicing, one of the outstanding questions is whether the reverse relationship exists. Namely, can the splicing reaction or specific splicing factors influence transcriptional regulation through effects on chromatin—either by direct coupling of histone modification with intron recognition or via regulated splicing of components of the histone modifying machinery? Several recent mammalian studies report that histone H3K36 methylation is mediated through splicing signals, although the mechanism for this is unclear [34, 35], and it is not clear whether other histone marks may also be influenced by splicing. In yeast, splicing has been shown to affect histone H2B ubiquitination by regulating the expression of a component of the histone H2B deubiquitinase machinery *SUS1*. However, thus far, no evidence of a direct relationship between histone H2B ubiquitination and splicing has been reported.
Yeast as a model to dissect regulated splicing

When facing complex scientific problems, a reductionist approach can often be a valuable tactic. The budding yeast *Saccharomyces cerevisiae* has provided such a tool since the discovery of the ACT1 intron [36]. With its streamlined genome and high functional conservation of the basic machinery [37] it has been particularly useful to uncover the intricacies of spliceosome assembly and catalysis. However, the scarcity of introns (~95% of yeast genes are intronless), and the lack of clear homologues of SR-proteins (ubiquitous splicing modulators in metazoan cells), has encouraged the view that the yeast cell does not use this important instrument to regulate gene output. While it is true that there are quantitatively fewer alternate mRNAs that arise from splicing changes in yeast compared to metazoans, regulation of splicing in yeast is essential to cellular fitness and meiotic differentiation (Figure 1). Given that the fundamental aspects of the spliceosome are conserved in evolution, the study of these regulatory strategies is relevant to understanding splicing across eukaryotes, and likely it will reveal possibilities that may be harnessed by other organisms. In fact, its reduced intronic set simplifies the study of splicing at the genomic level. This, combined with the manageability of yeast as a model system, has helped to advance our understanding of some molecular mechanisms behind regulated splicing.

As a single-celled organism, yeast has been subjected to strong selection for rapid and adaptive growth. A growing yeast cell is under the Sisyphean task of constantly adjusting to an ever-changing environment and one might imagine that regulation of splicing is an important part of this business. Indeed, the cell uses splicing to fine-tune RNA biogenesis, as genes encoding proteins involved in a number of critical steps in RNA biogenesis can undergo regulated splicing—including translation, RNA export, RNA splicing, and transcription. This provides a powerful tool for regulating gene expression as changing the capacity to translate mRNAs from the assembled and transported mRNPs can fundamentally alter gene expression. We will now focus on examples of how splicing affects these critical gene expression reactions.

Regulated splicing to control ribosome biosynthesis and function

Ribosome biosynthesis constitutes the yeast cell’s largest energy investment [38]. Hence, it is not surprising that expression of ribosome components is subject to tight regulation at multiple levels, including splicing. Ribosomal proteins are obvious candidates for splicing regulation, as over one third of yeast introns are located in their genes [39]. Furthermore, nearly a third of all mRNAs are produced from intron-containing genes—most of which are ribosomal protein genes (RPGs). This all leads to the obvious prediction that there are likely to be instances of regulation at the level of RPG splicing that may have so far escaped description.

Recent studies that systematically deleted the introns from intron-containing RP genes demonstrated that at least some introns were required for optimal fitness or adaptation to stress conditions, indicating that proper expression is tied to intron-dependent gene expression changes [40, 41]. In light of genome-wide studies indicating that inefficient splicing may be an important and widely used mechanism for downregulating gene expression in yeast [42], it is likely that at least one of the functions of the introns in RP genes is to confer regulatory flexibility and, perhaps, allow the rapid downregulation of ribosome biogenesis.

At a mechanistic level, there are undoubtedly a number of ways by which RPG intron removal is regulated. This includes autoregulation via the binding of the RP to its own pre-mRNA to affect its own splicing [43, 44]. Remarkably most introns are found within...
duplicated ribosomal protein genes, and not only does deletion of the intron in a duplicated RP gene affect the host gene (intragenic regulation), but also the expression of the paralog. Interestingly, in about a third of the cases, this intron-dependent extragenic regulation is dependent upon the intron in the affected paralog. These studies support models in which autoregulation and cross-paralog regulation at the level of intron removal play critical roles in controlled expression of ribosomal subunits [41]. In agreement with this, a new mechanism of nuclear RNA degradation, which is driven by introns that are bound by regulatory factors and the poly(A) binding protein, has been described in fission yeast [45]. Some \textit{S. cerevisiae} RPG's (notably \textit{RPS14} [46]) could be regulated in a similar manner, although it is apparent that cytoplasmic decay also plays a role in intron-dependent RNA degradation [47].

Of course an outstanding question raised by these studies is what role there is for differential expression of individual ribosomal proteins, including RPG paralogs? While the data above are suggestive of a link between splicing and ribosome function, the direct mechanism remains to be determined. Intriguingly, studies by Komili et al. demonstrated a differential requirement for paralogs for expression of localized RNAs. Furthermore, these studies also demonstrated that ribosomal protein paralogs have differential requirements for their assembly and localization, suggesting the existence of “specialized ribosomes” [48] [49]. Future experiments will undoubtedly shed light on this provocative possibility.

An example of how a specific RPG is regulated (and autoregulated) at the level of splicing to control ribosomal function is \textit{RPL30} (Figure 2). Expression of this gene includes an autoregulatory feedback loop, in which excess L30 binds to its own premRNA to stall spliceosome assembly. Thus, L30 can act as either a ribosomal protein or as a splicing factor. Therefore, a newly made L30 molecule is offered two choices: to either participate in ribosome function (ribosomal protein role); or to stop its own synthesis by binding to the pre-mRNA (splicing factor role). This strategy is based on evolving a binding site for the protein in the pre-mRNA plus a delicate balance of the relative affinities of L30 for the RNA and/or proteins it interacts with. Additionally, this binding site must be in a precise location in the pre-mRNA in order for the protein bound to it to block spliceosome assembly [50]. L30 (a mostly cytoplasmic protein, with a fraction in the nucleolus) binds co-transcriptionally to the \textit{RPL30}pre-mRNA to prevent full spliceosome assembly [50] and ultimately promote its export to the cytoplasm, where it gets degraded [51].

This scheme is essential for the cellular fitness [47] and it must be sensitive and fast to be efficient. In a growing yeast cell there are $10^5$ L30 molecules. Thus, it is likely that regulation of splicing must be responsive to small variations, since the starting L30 levels are already high. How the cell achieves this remains mostly unknown, yet it must involve rapid relocalization of L30, as L30 plays roles in distinct nuclear compartments. Presumably after failing to be recruited to the nascent nucleolar ribosome, L30 is able to rapidly find the site of \textit{RPL30} transcription, because every minute, more than one thousand L30 molecules must go through the same ritual (a growing yeast cell makes $10^5$ ribosomes in 1.5 hours). Consistent with the importance of the relative affinity of L30 for its two alternate destinations, unrestricted expression of an extra copy of an L30 version that fails to assemble into the ribosome is lethal. This is a consequence of repression of splicing of the endogenous \textit{RPL30} by this additional L30, thus depriving the cell of this essential protein [52].

From the point of view of splicing mechanisms, the fact that a ribosomal protein can block spliceosome assembly is intriguing. A model for this regulation is emerging based on the multiple conformational changes that occur during spliceosome assembly. According to this model, L30 binding to its pre-mRNA plays its inhibitory role by steric hindrance, blocking...
an early rearrangement. This supposes a novel way of splicing regulation, one that seems well-suited to RNA-binding proteins, and other examples of this strategy will surely follow.

**Regulated splicing to control RNP Biosynthesis and mRNA export: SUS1 and YRA1**

Just as is the case with ribosomal RNAs, most mRNA molecules spend their lives in a complex of proteins and RNA (RNP). During transcription, factors involved in capping, splicing, termination, and polyadenylation interact with the nascent RNA. This protein-bound RNA is delivered to the correct destination, which requires its interaction with specific factors. Once in the cytoplasm (for the RNAs that go there), the RNP is further remodeled to prepare the RNA for translation, storage, or further transport. Finally, many of these proteins bound to the RNA will need to be removed, and others recruited, before the RNA is degraded. These molecular rearrangements are each a potential target for regulation, and splicing appears to play a “behind the scenes” role in a number of these.

*SUS1* encodes a small, evolutionarily conserved protein that participates in both transcription and RNA export. As part of the SAGA complex, it is involved in histone H2B deubiquitination [53, 54]. During transcription, Sus1 interacts with the export factors Yra1 and Mex67. Sus1 is also at the nuclear pore, participating in the TREX2 complex (TRanscription EXport complex 2). In this capacity it plays a central role in RNA export.

Compared to other yeast genes, *SUS1* is atypical because it bears two introns, a feature shared by only a handful of yeast genes, and this gene architecture has been shown to be relevant to its function. Moreover, intron 1 of *SUS1* has a non-consensus 5’ end (A at position 6 instead of U) and BS (UACUGAC, as opposed to the canonical UACUAAC), and both of these features influence *SUS1* splicing. Analysis of *SUS1* splicing in WT cells indicates that some retention of the first intron occurs and, in fact, experiments using NMD-strains (deleted of the nonsense-mediated decay machinery) show that intron retention increases under different environmental conditions such as change in carbon source and upon temperature shift [55, 56]. Furthermore, recent results support the notion that the ability to produce different *SUS1* isoforms is essential for proper RNA export. It is not yet clear whether the change in *SUS1* splicing simply achieves a necessary change in the level of mature *SUS1* mRNA or if *SUS1* splicing changes mediate some other effect on RNA export. However, it is notable that a small peptide can be generated from the intron 1 containing transcript [55]. Further experiments are necessary to determine precisely how this peptide may affect the export machinery.

A second RNA export factor, Yra1, also has features that target it for regulated splicing. Although *YRA1* has a more typical structure, with only one intron, this intron is comparatively large and, more importantly, has a non-consensus BS sequence (GACUAAC). As with *SUS1*, *YRA1* splicing is suboptimal, and this decreased splicing efficiency contributes to its regulated intron retention and degradation to maintain the appropriate levels of Yra1 protein. Yra1 and the RNA helicase Sub2 are recruited cotranscriptionally to the nascent RNA, participating in the assembly of the TREX complex involved in RNA transport. When Yra1 levels are high, the Yra1 protein facilitates the export of the unspliced *YRA1* message such that it can be targeted to the Xrn1 (but not NMD)-dependent decay pathway in the cytoplasm [57-59], thus forming a tight regulatory loop.

While it is clear that splicing of particular transcripts is necessary for RNA export, an intriguing model is that splicing and transport are, in fact spatially coordinated to allow feedback regulation of the two processes. For example, in light of evidence that export factor recruitment and spliceosome assembly occur co-transcriptionally, it is likely that there is competition between RNA transport and splicing, allowing that poor splicing substrates
will tend to be exported as pre-mRNAs; conversely, their splicing will be favored under conditions of poor export (for instance, in export mutants). With this in mind, a simple model is that regulated splicing of export factors provides a mechanism by which the balance between export and splicing can be achieved by a feedback loop between the two pathways. If splicing cannot keep up with the export pipeline, the pipeline must be slowed down, achievable by inefficient production of YRA1 or SUS1. The reverse may also be true. If the export machinery must accommodate high levels of newly made transcripts, then export should be promoted. This could be achieved by more efficient production of export factors via splicing of pre-mRNAs encoding export factors such as Sus1 and Yra1 and a concomitant increase in the capacity for export of newly-synthesized mRNAs. In summary, this model posits that the export machinery is tightly coupled to splicing via the use of splicing sensitized pre-mRNAs. Consistent with the prediction of an important and delicate balance of these factors, overexpression of either Yra1 or Sus1 is toxic to the cell [55, 58].

Overall this model predicts that regulated splicing of specific transcripts could act as a “sensor” of robust RNA processing and a “control switch” to downregulate or upregulate the RNA export machinery. It is intriguing to envision that YRA1 and SUS1 (perhaps through its role in transcription) could play slightly different roles in the delicate fine-tuning of RNA metabolism.

**Regulated splicing to fine tune transcription: Regulation of histone modification and the meiotic transcription program**

A picture is emerging of how regulated splicing in *S. cerevisiae* plays an important role in regulating RNA export and fine-tuning ribosome biogenesis and function. Next we will explore how regulated splicing of specific transcripts influences transcriptional programs in response to environmental change. In addition to its role in RNA export via its interactions with the TREX-2 complex, Sus1 is a component of the transcriptional coactivating complex, SAGA. This complex is comprised of two catalytic modules: one that mediates histone acetylation and gene activation, and another module comprised of the histone H2B deubiquitinase Ubp8. Sus1 (along with Sgf11 and Sgf73) is required for the Ubp8-dependent deubiquitination [60, 61].

Since Sus1 contains two introns, the first of which has features that reinforce its retention, it is worth looking more closely at the details underlying removal of the Sus1 introns. Expression of SUS1 pre-mRNA in which the first intron is absent reveals that removal of the downstream intron is facilitated by recognition of the first [56]. While the mechanisms underlying this “cross-exon” effect are not clear, these results suggest that, as is the case in metazoans, removal of the introns from a multi-intron pre-mRNA can be coordinated [62], and it is possible that the cross-exon effects that are observed involve similar interactions as metazoan “exon-definition.” While the number of yeast genes with multiple introns in *S. cerevisiae* is small (10), non-canonical sequences in the first intron are overrepresented (half of the two-intron genes), suggesting that similar mechanisms observed for SUS1 splicing might influence processing of these transcripts.

As described above, stress conditions (e.g. temperature) lead to preferential retention of the first intron and changes in Sus1 protein levels. Furthermore, at higher temperature cells show increased sensitivity to deletion of *SUS1* [55, 56]. Only those cells that express a construct that is able to undergo splicing (and not the cDNA with the introns removed) can restore WT levels of H2B ubiquitination and WT growth in response to temperature change. These data suggest that production of SUS1 isoforms—mature and intron 1-retained RNA—allow proper levels of H2B ubiquitination in response to stress, perhaps by maintaining the delicate balance of Sus1 protein necessary for H2B deubiquitination within genes involved in stress response. Since the retained intron product appears to have two possible fates—
degradation through NMD and production of a truncated peptide [55], it remains to be
determined precisely how H2B levels are maintained by the production of the retained intron
isoform. Nevertheless, these data suggest that cellular response to stress involves control of
H2B ubiquitination via SUS1 splicing. Consistent with these observations, we have
observed that a number of genes involved in temperature stress response in S. cerevisiae are
upregulated under the very conditions under which the first intron of SUS1 is retained
(TJohnson, unpublished results). The role of SUS1 splicing regulation in RNA export and
histone H2B deubiquitination are summarized in Figure 3.

Both YRA1 and RPL30 provide mechanisms whereby the protein product regulates splicing
of its own transcript. While it is unclear whether Sus1 contributes to its own regulation, it
has previously been shown that the acetylation module of the SAGA complex (catalyzed by
Gcn5) is able to influence pre-messenger RNA splicing via its acetylation activity. It is
appealing to speculate that Sus1-mediated H2B deubiquitination or one of the Sus1 products
could mediate changes in the chromatin of the SUS1 gene that affect its own splicing,
adding an additional layer of regulation of SUS1 mRNA levels.

In the context of RNA export, SUS1 appears to be a “sensor” of robust RNA processing
(indicated by levels of the retained intron), and its gene product can control the fate of the
mRNPs through its export activity. Hence the coupling of splicing with export through one
protein ensures the fine tuned coordination of multiple processes. A parallel “fine tuning”
may occur via Sus1’s role in histone modification. In this case, the SUS1 transcript provides
a readout for splicing efficiency, with production of the retained intron product indicating a
decrease in splicing efficiency. The Sus1 gene product(s) could thereby modulate
transcription, perhaps of a critical subset of genes, via effects on H2B-ub. Such a feedback
mechanism would ensure a balance between the activity of the processing machinery and the
RNA load generated through transcription. These models of feedback between the Sus1
splicing and transcription to maintain homeostasis remain to be tested.

In addition to SUS1, it appears that the entire meiotic regulatory network is under tight
control via regulated splicing. Under conditions of nutrient starvation diploid yeast cells
enter meiosis and sporulate. As with ribosome biosynthesis, this process is tightly regulated
at the transcriptional level [63] as well as at the level of splicing [64, 65]. Consistent with
this, a number of intron-containing meiotic transcripts can be detected in vegetative cells,
but these remain unspliced [42]. Correct processing of four key meiotic messages requires
the meiosis-specific splicing factor Mer1. This factor promotes intron recognition in poor
substrates through its ability to interact with a conserved intronic enhancer sequence [66],
and the meiotic program cannot proceed without it.

Mer1 dependent splicing bridges two transcriptional regulatory networks in the meiotic gene
expression program. First, Mer1, like 3 of the 4 other meiotic genes that it regulates is under
the control of Ume6, the activator of early meiotic genes, thus ensuring that the regulation of
these transcripts is temporally constrained until there is sufficient accumulation of Mer1p
after Ume6-mediated activation. Furthermore, splicing of two Mer1 specific genes MER3
and SPO22 is necessary for expression of NDT80, a meiosis specific transcription factor
[64]. Hence, regulated splicing ensures control of a transcriptional program essential for
cells to induce meiotic gene expression.

Environment-dependent changes in splicing fine-tune the cellular mRNA landscape

While the focus of this review has been specific examples of regulated splicing, it is likely
that these examples, while important for their roles in coordinating multiple gene expression
reactions may only be the tip of the iceberg. Recent data indicate that the spliceosome can
affect the composition of the transcriptome (understood here as the cellular mRNA

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landscape) in response to environmental stimuli [67]. For example, changes in amino acid starvation or ethanol stress provoke alterations in the splicing of many genes. These changes are too rapid to be easily explained by the synthesis of new splicing factor(s) to modify the spliceosome activity, and no obvious cis-elements have been identified in the pre-mRNAs that would explain this behavior [67]. Nonetheless, the spliceosome can identify which, and how, pre-mRNAs are to be changed when the yeast cells encounter certain situations. It is possible that the rapid response to environmental change may be mediated by rapid and regulated changes in the core splicing machinery through, for example, post-translational modifications of components of the splicing machinery. Along these lines, large-scale proteomics studies reveal widespread modification of splicing factors [68]. Furthermore, splicing-sensitive microarrays confirm that intron-containing genes are differentially sensitive to the activities of specific splicing factors [69, 70], raising the possibility that modification of individual components of the splicing machinery could “fine-tune” it to target specific introns (or groups of introns).

Intriguingly, detailed studies of meiotic regulation of splicing in fission yeast, support previous mammalian studies that demonstrate that the promoter can dictate splicing regulation [71]. These studies suggest that some RNA-binding proteins, recruited at the promoter level, may tag the transcript in a way that can affect the spliceosome or that the elongation properties of the polymerase, which can be set up during early stages of transcription, affect splicing. Nonetheless, the ability of the cell to to coordinate changes in splicing of the transcripts from many genes is remarkable and illustrates the likely role for regulated splicing as a cellular strategy for adaptation to environment. Transcriptome analyses in upf1Δ and xrn1Δ mutants (where many RNAs that would otherwise be turned over are stabilized) indicate that there is a small population of “aberrant” transcripts, which includes unspliced, partially spliced, and misspliced RNAs [72]. Whether this is part of a regulatory mechanism or simply evidence that the cell efficiently “cleans up” sloppy splicing remains to be determined. However, the examples above of SUS1, RPL30, and YRA1 all illustrate that regulated degradation of differentially spliced transcripts provide a powerful mechanism whereby the cell can respond to changes in environmental demands.

Conclusions

As a critical player during early mRNA synthesis, pre-mRNA splicing can have a profound effect on gene expression. Akin to a limited molecular ghostwriter, the spliceosome modifies a first draft --the nascent transcripts-- to produce a message tailored to the cell’s needs and that can affect fundamental cellular pathways. Therefore, the cell must communicate to the spliceosome a wide variety of information, and the spliceosome must be able to respond quickly enough to elicit the appropriate response. Surely this underlies the remarkable complexity of the spliceosome; after all, group II introns, the likely ancestors of spliceosomal introns, can perform the same basic chemistry mostly by themselves [73]. This raises a number of outstanding questions. How is this communication carried out? Who are the key players? How do changes in chromatin mediate the communication between environment and splicing? And can splicing in fact, contribute to establish epigenetic changes that mark the “experience” of the cell? While we still know frustratingly little, recent data on co-transcriptional splicing indicate that the coupling between transcription and splicing can be modulated by transcript features, from secondary structures to the recruitment of trans acting splicing factors [74].

Regulation of splicing acts in two ways: working constantly to keep gene expression fine-tuned (e.g. maintaining proper levels of RNA export); and occasionally, when a new cellular program needs to be launched and maintained (e.g. meiosis). Whether there is crosstalk between these modes of regulation remains to be elucidated. For example, it is not known
whether regulated splicing may completely reprogram particular cellular processes when such changes are extremely relevant to the cell (e.g. protein synthesis). Notably, as we have seen more than a third of yeast introns are located in genes related to the ribosome, making ribosomal biogenesis and function highly subject to splicing control. This has been connected to the generation of “specialized” ribosomes [41]. It also may have particular relevance to understanding alterations in ribosome biogenesis, detrimental to health, as variations in the concentration of ribosomal proteins have been linked to disease [75] and development defects [76]. Along these lines, and given the relevance that splicing has to health, it is of great interest to develop strategies to alter the splicing outcome. Some initial successes in this area have been very encouraging [77, 78].

The U12 spliceosome constitutes the obvious example supporting the notion, analogous to that in ribosomes, of “specialized” spliceosomes with different versions of the core components. However, it seems likely that the cell should use less radical strategies to generate different spliceosome variants that could be the basis for some alternative splicing. Some spliceosomal components are encoded by several genes (see [79], for example) and they may assemble into spliceosomes with different properties. Similarly, other components are not essential, or their properties could be post-translationally modified. There is much to be learned as to how homogeneous spliceosomes are, and how this relates to the many cellular factors that control splicing (SR proteins and hnRNPs). Even in yeast, with a simplified splicing network, how the core spliceosome elicits specific splicing outcomes (like the nutrient response in yeast [67]) remains to be explained. Furthermore, many metazoan transcripts encoding regulatory splicing factors undergo regulated splicing themselves. Thus, the splicing code involving regulatory factors and pre-mRNA features [80] may need to be expanded to include the inherent properties of the spliceosome. This may seem a daunting task, but analyses of the increasing amount of transcriptome data are improving our ability to predict which will be the final mRNA synthesized from a given pre-mRNA and set of conditions.

In yeast, with just a relative handful of introns, splicing has a central role in the control of gene expression. Moreover, the possibility that its streamlined spliceosome can be subjected to the same regulatory strategies, including those involving co-transcriptional splicing through a chromatin template and cross exon interactions, is particularly exciting. Thus, future studies that take advantage of the yeast system are likely to yield more fundamental insights into complex aspects of splicing regulation.

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Highlights

- Two simple and consecutive chemical reactions are catalyzed cotranscriptionally by a large macromolecular complex.

- The spliceosome is essential for generating mature mRNA.

- The cell takes advantage of this to generate different amounts and isoforms of mRNAs, according to its needs.

- In yeast, control of pre-mRNA splicing of critical genes allows the cell to affect all the pathways in gene expression.

- The spliceosome can be both the sensor and controller to fine tune genetic output.
Pre-mRNA splicing can fine-tune several different gene expression reactions via splicing sensitive regulators, including RNP biogenesis and transport, ribosome biogenesis and function, or transcription itself. The factors shown in the ovals undergo regulated splicing to affect the processes shown in the boxes. It is likely that others are still unknown. Nevertheless, those indicated provide a paradigm for regulatory strategies centered on the control of pre-mRNA splicing.
Figure 2.
Splicing regulation of the RPL30 transcript. The RPL30 gene encodes the essential protein L30. Pre-mRNA splicing contributes to maintaining the protein levels by an autoregulatory feedback loop. The newly made L30 (shown as a ball) must be transported to the nucleolus to join ribosome biosynthesis (top). However, if that is not possible (conditions of excess L30), the protein binds its own pre-mRNA. This prevents RPL30 splicing and triggers the transport and decay of the pre-mRNA in the cytoplasm. The choice for L30 between either ribosome biosynthesis or splicing control (orange square) has been depicted in the nucleus, but its mechanism and location is unknown (for simplicity reasons the binding site of L30 is shown entirely in exon 1, although it includes the 5' ss).
Figure 3.
Regulated splicing of *SUS1* fine-tunes RNA export and histone H2B ubiquitination. Under unstressed conditions (Left side of figure) *SUS1* pre-mRNA undergoes splicing and is exported to allow production of Sus1 protein that can either participate in histone H2B deubiquitination as part of the SAGA complex (top) or can associate with the TREX-2 proteins at the nuclear pore (Shown in dark blue). Intron 1 is retained in a small fraction of the RNAs and is either targeted to NMD or may make a truncated product. Under stress conditions, intron 1 retention is enhanced. However, it is unclear how much of this product is targeted to NMD vs. translation of a truncated product. It is possible that the truncated protein can associate with either the TREX-2 complex or with SAGA (indicated by “?”).
Sgf11 is destabilized by the lack of Sus1 [81], so it has been left off out of the SAGA complex that contains less SUS1. It is also possible that Sus1, via its H2B ubiquitination activity or by interacting with the splicing (not shown), may influence its own splicing (Dotted arrow).