Alternative polyadenylation: New insights from global analyses

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Alternative polyadenylation: New insights from global analyses

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ABSTRACT
Recent studies have revealed widespread mRNA alternative polyadenylation (APA) in eukaryotes and its dynamic spatial and temporal regulation. APA not only generates proteomic and functional diversity, but also plays important roles in regulating gene expression. Global deregulation of APA has been demonstrated in a variety of human diseases. Recent exciting advances in the field have been made possible in a large part by high throughput analyses using newly developed experimental tools. Here I review the recent progress in global studies of APA and the insights that have emerged from these and other studies that use more conventional methods.

Keywords: alternative polyadenylation; genomics; pre-mRNA processing

INTRODUCTION
3’ end processing is an essential step of eukaryotic mRNA maturation, which typically involves a cotranscriptional endonucleolytic cleavage and the subsequent addition of a string of adenosines (Colgan and Manley 1997; Zhao et al. 1999; Danckwardt et al. 2008; Millevoi and Vagner 2009; Chan et al. 2011). The poly(A) site (PAS) is defined by multiple cis-elements. For canonical mammalian PASs, these cis-elements include an AAUAAA hexamer, a U- or GU-rich downstream element (DSE), and other auxiliary sequences including the U-rich upstream stimulatory element (USE). PAS recognition relies on several multisubunit protein complexes. The CPSF (cleavage and polyadenylation specificity factor) complex recognizes the AAUAAA hexamer, while the CstF (cleavage stimulation factor) and CFIm (cleavage factor Im) complexes bind the DSE and UGUA motifs, respectively. These factors in turn recruit CFIm (cleavage factor Im) and other proteins to form the 3’ processing complex in which cleavage and polyadenylation take place (Colgan and Manley 1997; Zhao et al. 1999; Shi et al. 2009; Chan et al. 2011). The majority of the core 3’ processing factors are well conserved.

For a large portion of eukaryotic genes, their mRNAs have multiple alternative 3’ ends that are formed by cleavage/polyadenylation at distinct sites, a phenomenon known as alternative polyadenylation (APA) (Di Giammartino et al. 2011; Lutz and Moreira 2011). Various systems and nomenclatures have been used for classification of APA. Here I consider two basic groups based on the distribution of the alternative PASs (Fig. 1). The first group is called same exon APA (SE-APA), in which the alternative PASs are found within the same terminal exon. In the second group, referred to as different exon APA (DE-APA), the alternative PASs are located within distinct exons. DE-APA can be further divided into three subgroups based on the sequence context of the PASs (Fig. 1). In DE-APA3 (nomenclature partially adapted from Zhang et al. 2010), the usage of alternative 3’ splice sites is coupled with cleavage/polyadenylation within mutually exclusive terminal exons. In the second subgroup, called DE-APA5, when an alternative 5’ splice site within an internal exon is skipped, cleavage/polyadenylation occurs at a downstream otherwise intronic PAS. Unlike DE-APA3 and DE-APA5, which are coupled with alternative splicing, the third subgroup involves the usage of a PAS within a constitutive internal exon that results in a truncated mRNA (Yao et al. 2012), and therefore is called DE-APA1 (I for internal). The various types of APA described here may have different regulatory mechanisms and functional consequences, which will be further discussed below. It should be pointed out that although most previous APA studies have focused on mRNAs, APA has also been reported for noncoding RNAs (Liu et al. 2010). The classification system proposed...
here could be applied to the APA of noncoding RNAs as well.

**APA STUDIES: FROM SINGLE GENES TO GLOBAL ANALYSES**

In 1980, the earliest examples of APA were discovered in the mRNAs of the IgM and dihydrofolate reductase (DHFR) genes (Alt et al. 1980; Early et al. 1980; Rogers et al. 1980; Setzer et al. 1980). It was shown that the APA (DE-APA, according to the nomenclature proposed in this article) isoforms of IgM encode distinct proteins, the secreted or the membrane-bound forms of IgM (Alt et al. 1980; Early et al. 1980; Rogers et al. 1980). In contrast, the APA (SE-APA) isoforms of DHFR differ in their 3′ untranslated regions (3′ UTRs) (Setzer et al. 1980). These early studies also suggested that APA is developmentally regulated. Since then, evidence for APA had been slowly accumulating in the literature. By 1997 APA was reported for the mRNAs of ~80 genes (Edwards-Gilbert et al. 1997). The potential prevalence of APA was first revealed by global characterizations of APA through bioinformatic analyses of the Expressed Sequence Tag (EST) data (Tian et al. 2005; Yan and Marr 2005). These studies suggested that over 50% of human genes undergo APA and many APA events display tissue-specific patterns (Zhang et al. 2005). In 2008 and the following year, the APA field was ushered into the “global analyses era” by several pioneering studies that, using microarray-based approaches, detected widespread and coordinated APA changes during T-cell activation (Sandberg et al. 2008), neuronal activity (Flavell et al. 2008), and development (Ji et al. 2009). These studies were quickly followed by the introduction of high-throughput sequencing-based techniques for global APA analyses that dramatically accelerated the pace of discovery in the field (Jan et al. 2010; Mangone et al. 2010; Ozsolak et al. 2010; Shepard et al. 2011). In the past 3 yr alone, the RNA polyadenylation profiles have been characterized for yeast (Ozsolak et al. 2010), *Caenorhabditis elegans* (Jan et al. 2010; Mangone et al. 2010; Haenni et al. 2012), *Drosophila* (Smibert et al. 2012), *Arabidopsis* (Wu et al. 2011; Sherstnev et al. 2012), and mammals (Ozsolak et al. 2010; Fox-Walsh et al. 2011; Fu et al. 2011; Shepard et al. 2011; Derti et al. 2012; Jenal et al. 2012; Lin et al. 2012). In most of these species, the polyadenylation profiles have been characterized for different tissues and developmental stages. Recent global studies have identified new APA regulators and novel regulatory mechanisms and revealed widespread APA changes associated with human diseases. Here, I review the technologies that propelled the recent advances and new insights that were generated by these global analyses as well as other studies using conventional methods.

**TECHNOLOGIES FOR GLOBAL APA ANALYSES**

The earliest global experimental studies of APA used various microarray platforms (Flavell et al. 2008; Sandberg et al. 2008; Ji et al. 2009). In these studies, the APA profile of a gene was deduced by the ratio between the average signal intensities of probes targeting the extended region that are found only in the longer APA isoform and that of other probes targeting the same gene. The success of these studies in detecting widespread APA changes demonstrated that microarrays are effective tools for global APA analyses. Additionally, when splicing-sensitive arrays are used, both alternative splicing and APA information can be obtained from the same analysis (Flavell et al. 2008; Sandberg et al. 2008). However, microarrays also have several drawbacks. For example, microarrays cannot directly detect novel PASs or map the PAS locations precisely, and quantification may be challenging for mRNAs with more than two APA isoforms. Finally, the availability of microarrays may be another limiting factor, especially for less-studied species.

The advent of high-throughput sequencing has dramatically transformed biomedical research in recent years. For example, RNA-seq has become the method of choice for gene expression analyses (Ozsolak and Milos 2011). For mRNA analysis, a cDNA library generated from fragmented poly(A)+ RNAs is sequenced and the resulting reads are mapped to the reference genome to obtain information on mRNA sequence and relative abundance. mRNA-seq is highly quantitative and is particularly powerful for analyzing mRNA isoform expression. mRNA 3′ ends can be mapped using reads that contain a string of untemplated adenosines (Nagalakshmi et al. 2008; Wang et al. 2008). Quantification of APA isoforms with mRNA-seq data is achieved by calculating the ratio of read density in the region found only in the longer APA isoform and that in the common region. An mRNA-seq analysis of the human
transcriptome, including 10 diverse tissues and five mammary epithelial or breast cancer cell lines, has provided a wealth of information on tissue-specific APA and the coordinated regulation of alternative splicing and APA (Wang et al. 2008). However, similar to microarrays, quantification may be difficult for mRNAs with more than two APA isoforms. More importantly, because only a small portion of the reads can be used to distinguish different APA isoforms, and even less reads map to the poly(A) junctions, greater sequencing depths are required for accurately mapping PASs and reliably quantifying APA isoforms.

To harness the power of high-throughput sequencing specifically for RNA polyadenylation analysis, a number of methods have been developed for targeted sequencing of poly(A) junctions. The most commonly used methods rely on oligo(dT)-primed reverse transcription–PCR (RT–PCR) for library construction (Mangone et al. 2010; Fox-Walsh et al. 2011; Fu et al. 2011; Shepard et al. 2011; Derti et al. 2012). Different versions of this method have been described with modifications designed to improve library construction and/or to facilitate sequencing (Shepard et al. 2011; Derti et al. 2012; Martin et al. 2012). The obvious advantage of the approach is its simplicity and quantitative nature. The main disadvantage, however, is the possibility of oligo(dT) priming at internal A-rich sequences at the RT step, resulting in potentially higher false discovery rate. To address these limitations, multiple computational approaches have been developed to filter out the majority of sequencing reads generated by internal priming (Mangone et al. 2010; Derti et al. 2012). Recent studies have demonstrated that these methods are highly effective in detecting global APA changes (Fu et al. 2011; Shepard et al. 2011; Jenal et al. 2012; Martin et al. 2012). A more sophisticated and labor-intensive method, called 3P-seq, uses more enzymatic steps for library construction, but avoids oligo(dT) priming and the associated internal priming issue (Jan et al. 2010). Due to its low false-positive rate, 3P-seq is especially suited for accurately mapping the 3′ ends of mRNAs. However, some of the enzymatic steps in 3P-seq, such as RNA adaptor ligation, are known to introduce bias and could affect its quantitative performance (Hafner et al. 2011). cDNA libraries prepared using either the oligo(dT)-primed RT–PCR or the 3P-seq method are most commonly sequenced on the Illumina platform, which offers high yield, low error rate, and multiplexing capability (Ozsolak and Milos 2011). In contrast, a third method, called direct RNA sequencing (DRS), uses the Helicos single-molecule sequencing platform (Ozsolak et al. 2010). In this method, the poly(A)+ RNAs are captured on oligo(dT) coated slides and the RNAs are directly sequenced by synthesis. DRS is believed to be more quantitative, as individual RNAs are sequenced without RT, PCR, or any other enzymatic reactions associated with typical cDNA library construction (Ozsolak and Milos 2011). Additionally, internal priming is rare in DRS, and attomole levels of RNAs are sufficient for DRS analysis (Ozsolak et al. 2010). When compared with the Illumina system, the disadvantages of DRS include higher error rate, shorter read lengths, lower yield (read counts), and the lack of multiplexing capability (Ozsolak and Milos 2011). Although at present no direct comparison of the three methods is available, all have been successfully applied to global analyses of polyadenylation. It should be emphasized, however, that novel PASs identified using these ultra-sensitive deep-sequencing methods must be carefully validated using independent methods to distinguish between genuine and functionally relevant PASs from sequencing noise and the intrinsic heterogeneity of cleavage/polyadenylation sites (Pauws et al. 2001).

Another key technology that is critical for mechanistic studies of APA regulation is the global mapping of protein–RNA interactions by UV cross-linking immunoprecipitation coupled with high-throughput sequencing (HITS–CLIP or CLIP–seq) (Ule et al. 2003; Licatalosi et al. 2008). Modifications of this method, such as PAR–CLIP (Photoactivatable-Ribonucleoside-Enhanced CLIP) and iCLIP (individual-nucleotide resolution CLIP), were subsequently introduced to improve the cross-linking efficiency and/or mapping resolution (Hafner et al. 2010; Konig et al. 2010). Recently, PAR–CLIP has been applied to globally mapping the RNA-binding sites for many core 3′ processing factors (Martin et al. 2012). Integration of transcriptome-wide protein–RNA interaction mapping with global functional analyses allows for construction of information-rich mechanistic models, called “RNA maps” (Licatalosi et al. 2008; Yeo et al. 2009). This approach has proven highly effective in elucidating the functions of RNA-binding factors in the regulation of pre-mRNA processing.

**INSIGHTS FROM GLOBAL STUDIES OF APA**

Recent global studies have made significant contributions to our understanding of APA. First, recent studies revealed that APA is much more pervasive than previously appreciated. Secondly, global analyses revealed new APA regulatory mechanisms at the transcriptome level. Thirdly, global studies have begun to reveal the functional consequences of APA and its role in a number of human diseases including cancer. These new insights are discussed in detail below.

**APA is a pervasive and evolutionarily conserved mechanism for gene regulation in eukaryotes**

Table 1 summarizes the results of recent global RNA polyadenylation mapping in various eukaryotic species. Thirty to seventy percent of genes from all species studied are estimated to produce alternatively polyadenylated mRNAs. In human, the latest estimate is that ~70% of genes undergo APA (Derti et al. 2012), which is significantly higher than earlier estimates based on EST data (Tian et al. 2005; Yan and Marr 2005). Perhaps most surprisingly, APA was detected for the mRNAs of ~70% of genes in the
budding yeast *Saccharomyces cerevisiae* (Ozsolak et al. 2010), a prevalence level comparable to that observed in the plant and human transcriptomes (Wu et al. 2011; Derti et al. 2012; Sherstnev et al. 2012). Although the large number of novel PASs detected in these high-throughput sequencing analyses need to be rigorously validated and further analyzed, recent global studies provided evidence that APA is more widespread than previously thought. The prevalence of APA in all eukaryotes tested is in sharp contrast to alternative splicing, which is extremely rare in yeast but nearly universal in all eukaryotes tested (Jan et al. 2010; Shepard et al. 2011; Martin et al. 2012; Smibert et al. 2012; Ulitsky et al. 2012). Therefore, the conserved PAS sequence features may have important implications for the evolution of the genomes (Jan et al. 2010).

Secondly, a general pattern for alternative PAS arrangement has been observed for many genes with APA. Overall, the most distal PASs (relative to the 5’ end of the mRNA) tend to have strong consensus features of canonical PAS, including the AAUAAA hexamer and strong DSEs, and are generally well conserved. In contrast, proximal PAS tend to diverge from the canonical sequence features and are generally less well conserved (Fig. 2B). This pattern has been observed in all metazoans studied (Beaudoing et al. 2000; Jan et al. 2010; Shepard et al. 2011; Martin et al. 2012; Smibert et al. 2012; Ulitsky et al. 2012). Interestingly, a similar arrangement was also suggested for alternative PASs in several yeast genes (Sparks et al. 2011). Therefore, the conserved PAS sequence features may have important implications for the evolution of the genomes (Jan et al. 2010).}

**TABLE 1. Summary of alternative polyadenylation in different species**

<table>
<thead>
<tr>
<th>Species</th>
<th>Genes with APA</th>
<th>Samples analyzed</th>
<th>References</th>
</tr>
</thead>
<tbody>
<tr>
<td><em>Saccharomyces cerevisiae</em></td>
<td>72.1%</td>
<td>Poly(A)+ RNAs</td>
<td>Ozsolak et al. (2010)</td>
</tr>
<tr>
<td>Arabidopsis</td>
<td>70%–75%</td>
<td>Seeds and leaves</td>
<td>Wu et al. (2011)</td>
</tr>
<tr>
<td>C. elegans</td>
<td>43%</td>
<td>Embryo, larvae (L1, L2, L3, L4, dauer), adult</td>
<td>Sherstnev et al. (2012)</td>
</tr>
<tr>
<td>Drosophila</td>
<td>54.3%</td>
<td>29 tissues and S2R+ cells</td>
<td>Smibert et al. (2012)</td>
</tr>
<tr>
<td>Zebrafish</td>
<td>55%</td>
<td>Four embryoic stages, mixed gender adults, and three tissues</td>
<td>Ulitsky et al. (2012)</td>
</tr>
<tr>
<td>Mammals</td>
<td>~70%</td>
<td>24 matched tissues</td>
<td>Derti et al. (2012)</td>
</tr>
</tbody>
</table>

**TABLE 2. Conserved poly(A) site sequence features and arrangement.**

(A) Conserved poly(A) site sequence features. U-rich sequences are shown in blue, A-rich sequences in red, UA-rich sequences in pink. Nucleotide positions relative to the cleavage site are labeled. (USE) Upstream sequence element; (ARE) A-rich element; (DSE) downstream sequence element. (B) Conserved arrangement of poly(A) sites. The arrows point to the 3’ end of the transcript. The most distal PAS has strong canonical features and is shown as a blue and red box, similar to A. The proximal PAS tend to diverge from canonical PAS features and are shown as light blue boxes.

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and Dieckmann 1998). These observations indicate that the conserved distal PASs may serve as the default sites, while the proximal PASs may mainly play regulatory roles. Finally, the temporal and spatial regulation of APA appears highly conserved. For example, mRNAs in the brain or neuronal cells tend to use the most distal PAS and therefore have the longest 3' UTRs, while mRNAs in the testis or ovaries tend to favor proximal PAS to generate mRNAs with the shortest 3' UTRs (Smibert et al. 2012; Ulitsky et al. 2012). In mammals, the APA profiles are more similar among the same tissues than among different tissues in the same species (Derti et al. 2012). Together, these observations strongly suggest that APA is a pervasive and conserved mechanism for eukaryotic gene regulation.

Mechanisms for APA regulation: The “polyadenylation code”

APA is a dynamic process and recent global analyses have revealed widespread APA changes in many physiological processes as well as in human diseases. One of the most important observations from these studies is that APA, especially SE-APA, is intimately connected with the cellular proliferation and differentiation status (Sandberg et al. 2008; Ji et al. 2009; Shepard et al. 2011). In proliferating cells, proximal PASs tend to be used to generate mRNA isoforms with shorter 3' UTRs. In contrast, in resting or differentiated cells distal PASs are favored to generate mRNAs with longer 3' UTRs. This type of systematic and coordinated APA regulation has been observed upon activation of neurons and T cells (Flavell et al. 2008; Sandberg et al. 2008) and in development (Ji et al. 2009), stem cell differentiation (Shepard et al. 2011), and the generation of induced pluripotency (iPS) cells (Ji and Tian 2009). In some cases, there is evidence suggesting that APA regulation has significant functional consequences. For example, 3' UTR shortening in the mRNAs of proto-oncogenes have been shown to activate these genes and lead to higher transformation (Mayr and Bartel 2009). However, the mechanisms underlying these APA changes remain poorly understood.

Global studies of RNA polyadenylation have provided a rich resource for deciphering the rules that govern PAS choice, which I refer to as the “polyadenylation code.” To understand the mechanisms for PAS selection, it is helpful to examine the chemical reaction of mRNA 3' processing itself (Fig. 3A). In this reaction, the essential components include CPSF, CstF, CFIm, CFIIIm, PAP, and the PAS-containing pre-mRNA. Two molecules of CstF and CFIm are required as they likely function as dimers (for review, see Chan et al. 2011). The overall reaction rate is a function of the concentrations of all reactants (Fig. 3A). Each alternative PAS has its unique rate constant (k) based on how efficiently it is recognized by the 3' processing machinery and regulators. In addition, proximal PAS have an intrinsic advantage over the distal ones, as they are transcribed earlier and have more time to be recognized. In other words, the concentration of the proximal PASs is higher than that of the distal PASs during transcription. Therefore, both the sequence and the relative position of the alternative PASs are important factors that ultimately determine APA outcome. Below I present a few specific models of APA regulation that have emerged from previous studies (Fig. 3B–E).

FIGURE 3. Regulatory mechanisms for poly(A) site selection. (A) Depiction of the mRNA 3' processing reaction and the reaction rate formula. $k$ is the reaction constant. [X]: Concentration of X. PAS is the poly(A) site-containing pre-mRNA. See text for details. (B–E) Regulatory mechanisms for poly(A) site selection. Poly(A) sites (PAS) are shown as rectangular boxes. Core 3' processing factors are shown as orange boxes. The directionality of the mRNAs is marked with arrows. In B and E, the strengths of the alternative PAS are labeled. Strong PAS are shown as dark blue boxes and weak PAS as light blue boxes.
The ‘survival of the fittest’ model

PASs with higher affinities for the 3’ processing machinery most likely have greater rate constants and therefore tend to be favored in APA. In addition to the intrinsic strengths of alternative PASs, however, PAS selection can be modulated by the levels of the core 3’ processing factors. The Manley group provided the first evidence for such a mechanism (Takagaki et al. 1996). They observed that during B-cell activation elevated protein levels of the core 3’ processing factor CstF64 promoted the usage of a proximal PAS in the IgM mRNAs that contains a weak CstF64 binding site. However, in resting B cells, which have much lower levels of CstF64 (limited resource), the distal PAS with a strong CstF64 binding site was preferentially recognized. I will refer to this mechanism as the “survival of the fittest” model (Fig. 3B). A similar mechanism was proposed to explain the APA changes of the NF-ATc mRNAs during T-cell activation (Chuvpilo et al. 1999). This mechanism may underlie, at least in part, the global APA differences between proliferating and differentiated cells (Di Giammartino et al. 2011). As mentioned earlier, proximal PASs tend to have limited consensus PAS features (likely weaker sites) and are favored in proliferating cells. In differentiated cells, polyadenylation shifts to the distal PASs that have canonical cis-elements (likely stronger sites). Interestingly, both microarray and sequencing analyses suggest that most core 3’ processing factors are expressed at higher levels in proliferating cells than in differentiated or resting cells (Ji et al. 2009; Elkon et al. 2012). Therefore, higher levels of 3’ processing factors in proliferating cells may promote the recognition of weaker proximal PAS, while the limited amounts of 3’ processing factors in differentiated cells may be preferentially recruited to the stronger distal PASs. The expression of the core 3’ processing factors can be regulated transcriptionally, for example, by the transcription factor E2F (Elkon et al. 2012) or by signaling pathways, including the p38 MAP kinase pathway (Danckwardt et al. 2011). In addition to the molecular abundance of core 3’ processing factors, post-translational modifications or interactions with other factors can modulate their activities or availability in the nucleus (Millevoi and Vagner 2009), which may contribute to APA changes as well.

Although the “survival of the fittest” model can explain many APA events, global studies suggest that the relationship between the levels of 3’ processing factors and PAS selection is more complicated. For example, it has been reported recently that CFIm68 promotes the usage of distal PASs in many mRNAs (Martin et al. 2012). Depletion of CFIm68 by RNAi results in a widespread APA shift toward proximal PASs, which, in general, are poorer binders for CFIm68. The observed APA changes are consistent with earlier studies of CFIm-mediated APA regulation (Kubo et al. 2006; Kim et al. 2010). These studies suggest that the protein levels of different core 3’ processing factors may influence the global APA profile differently. Refinement of this model will require deeper understanding of the interactions between the core 3’ processing factors and the PAS sequence and more comprehensive studies on APA regulation mediated by the core 3’ processing factors.

The agonist/antagonist model

Recognition of alternative PASs by the core 3’ processing factors can be modulated, either positively or negatively, by regulatory factors (Fig. 3C). The best-known examples of APA regulators are RNA-binding proteins that function through two mechanisms. First, some APA regulators directly compete with the core 3’ processing factors for binding to the PAS region. For example, it was shown that the mammalian PTB and Droso phila sex lethal proteins can modulate PAS recognition by competing with CstF64 for binding to the DSEs (Castelo-Branco et al. 2004; Gawande et al. 2006). Similarly, recent studies demonstrated that the RNA-binding protein HuR regulates the APA of its own transcripts by competing with CstF64 for binding to one of the alternative PASs, thereby suppressing its usage (Dai et al. 2012; Mansfield and Keene 2012). Global analyses revealed that when Nova2, a brain-specific RNA-binding protein, binds to RNAs within the PAS regions, it tends to inhibit the usage of that PAS, possibly by a steric hindrance mechanism (Licatalosi et al. 2008). The epithelial specific RNA-binding proteins ESRP1/2 may regulate APA through a similar mechanism as they bind to GU-rich motifs, and thus may directly compete with CstF64 for the DSEs (Dittmar et al. 2012). Another example of this type of regulators is PABPN1. In addition to its well-characterized functions in polyadenylation and poly(A) tail length control (Eckmann et al. 2011), PABPN1 has recently been shown to suppress the usage of proximal PAS globally (Jenal et al. 2012). As mentioned earlier, proximal PASs tend to lack consensus PAS sequences, such as the AAUAAA hexamer. It was proposed that PABPN1 competes with CPSF for binding to the A-rich regions at proximal PASs, thereby inhibiting their usage. When PABPN1 is depleted, however, CPSF can recognize these PASs and promote their usage (Jenal et al. 2012).

The second type of APA regulators binds to mRNA sequences outside of the PAS region to regulate their usage. U1 snRNP is one of the best-characterized examples. In vitro studies demonstrated that U1 snRNP bound to a 5’ splice site suppresses cleavage/polyadenylation at downstream PAS, which is mediated by U1-70K and U1A inhibition of PAP (Gunderson et al. 1994). Recent global studies showed that U1 globally suppresses premature cleavage/polyadenylation within introns and that U1 depletion leads to the activation of intronic PASs and global APA changes in a concentration-dependent manner (Kaida et al. 2010; Vorlova et al. 2011; Berg et al. 2012). Although
overall U1 levels are not known to fluctuate, global transcriptional activation has been suggested to cause a transient U1 shortage, which in turn results in changes in PAS selection (Berg et al. 2012). In addition to negative regulators such as U1, some RNA-binding factors can promote PAS recognition. FCA and FPA are RNA-binding proteins that play important roles in flowering time control in Arabidopsis (Simpson et al. 2003; Hornýik et al. 2010). Both proteins regulate the APA of their own transcripts by binding to RNAs near the proximal PASs to promote their usage. For FCA, its effect on PAS selection relies on its interaction with the FY protein, a highly conserved core 3’ processing factor (Simpson et al. 2003).

The “poisedness” model

It is now appreciated that all the steps of mRNA biogenesis are tightly coupled and one step could affect another (Hirose and Manley 2000; Maniatis and Reed 2002; Bentley 2005; Moore and Proudfoot 2009). For example, most core 3’ processing factors have been shown to associate with the transcription machinery and these interactions play an important role for efficient 3’ processing (Chan et al. 2011). Accumulating evidence suggests that transcription factors can regulate the efficiency of cotranscriptional recruitment of 3’ processing factors, which in turn modulates APA. For example, ELL2, a transcription elongation factor, has been suggested to promote the loading of CstF onto the transcription machinery so that CstF is more poised to engage with PAS, thereby promoting the usage of proximal PASs (Martincic et al. 2009). I refer to this mechanism as the “poisedness” model (Fig. 3D). A recent global study provided further evidence for this mechanism by demonstrating that the promoters and transcriptional activities have a significant impact on APA (Ji et al. 2011). Reporter assays and bioinformatic analyses of RNA-seq data from human and mouse tissues suggested a correlation between higher transcription activity and higher usage of the proximal PASs. Additionally, more RNA polymerase II (RNAP II) pausing and differential nucleosome occupancy and histone modification patterns were observed at the proximal PASs for highly expressed genes. These results support a model in which transcriptional activators promote efficient recruitment of the mRNA 3’ processing machinery, and thus promote the recognition of the proximal PASs. The PAF complex may play an important role in this mechanism given its functions in coordinating transcription, histone modifications, and mRNA 3’ processing (Rozenblatt-Rosen et al. 2009). It was shown in vitro that transcriptional activators stimulate cotranscriptional mRNA 3’ processing by directly recruiting the PAF complex, which in turn recruits CPSF and CstF (Nagaike et al. 2011). It will be of interest to test the impact of the PAF complex on global APA profiles. The Mediator, a transcriptional coactivator complex, has also been shown to influence the global APA profile by modulating the occupancy of HnRNP L, a Mediator-associated protein and a regulator of mRNA alternative processing, on its target transcripts (Huang et al. 2012). These studies provided evidence that the efficiency of cotranscriptional recruitment of the 3’ processing machinery and regulatory factors is an important determinant of the APA outcome.

The “first come, first served” model

As mentioned before, proximal PAS have an intrinsic advantage over the distal ones as they are transcribed earlier (“first come, first served”) (Fig. 3E). Conceptually, such an advantage should correlate with the distance between alternative PASs and the transcription elongation rate. Indeed, it has recently been shown that APA is modulated by the transcription elongation rate. Using the Drosophila polo mRNAs as a model, Pinto et al. (2011) showed that a RNAP II mutant with slower elongation rate kinetically favors the recognition of the proximal PAS. In contrast, in cells expressing the wild-type RNAP II, the intrinsic advantage of the proximal PAS becomes less significant, and as a result, the relative usage of the distal PAS increases (Pinto et al. 2011). A similar mechanism has been previously shown to regulate alternative splicing (de la Mata et al. 2003). Although it remains poorly understood how transcription elongation rate is regulated, accumulating evidence suggests that chromatin and chromatin-associated factors may play an important role (Brown et al. 2012). For example, CTCF is a DNA-binding factor that insulates active genomic regions from inactive ones and promotes long-range interactions in the genome (Phillips and Corces 2009). Recently, it has been shown that CTCF, when bound to DNA in the coding regions, causes local RNAP II pausing, which in turn promotes the recognition of upstream splice sites (Shukla et al. 2011). Furthermore, CTCF–DNA interactions are inhibited by DNA methylation, thereby linking DNA methylation to alternative splicing regulation (Shukla et al. 2011). It is likely that similar mechanisms could influence APA as well. In fact, it has been reported that genomic imprinting regulates the APA of the mouse H13 and Herc3/Nap1l5 transcripts (Wood et al. 2008; Cowley et al. 2012). In both cases, the genomic sequences encoding alternative PASs are separated by a CpG island, which is differentially methylated on the paternal and maternal alleles. Transcripts expressed from the unmethylated allele are polyadenylated at the PAS upstream of the CpG island to produce a truncated mRNA, while those expressed from the methylated allele favor the downstream PAS. Similar to the mechanism for CTCF-mediated alternative splicing regulation, a methylation-sensitive DNA-binding factor may bind to the CpG island on the unmethylated allele, cause RNAP II pausing, and promote the usage of the upstream PAS. This model needs to be tested in future studies.
It is important to note that the transcription activities of a gene are not directly correlated with its transcription elongation rate. In fact, as discussed above, high transcription activity promotes the usage of proximal PASs through the “poisedness” model, while higher elongation rate leads to enhanced recognition of the distal PASs (the “first come, first served” model).

PAS selection is most likely under combinatorial control of multiple mechanisms. For example, tissue- or developmental stage-specific APA profiles may be generated by several possible mechanisms. The core 3’ processing factors or APA regulators may be expressed at different levels in different tissues or at different times during development (Ji et al. 2009). For instance, CstF64r, a CstF64 paralog, is highly expressed in testis (Wallace et al. 1999; Dass et al. 2007). As described earlier, the APA regulators, Nova2 and ESRP1/2, are specifically expressed in the brain and epithelial cells, respectively (Ule et al. 2005; Warzecha et al. 2009). Thus, these factors may contribute to tissue-specific PAS selection. Similarly, multiple mechanisms have been proposed to explain the 3’ UTR lengthening during differentiation, including changes in 3’ processing factor abundances and U1 snRNP availability (Ji et al. 2009; Berg et al. 2012). Therefore, a major challenge in deciphering the “polyadenylation code” is to integrate all of the different mechanisms in a quantitative manner.

**Functional impact of APA**

A major contribution of the recent global studies is the realization that APA is a widespread mechanism for eukaryotic gene regulation. APA could impact gene expression through several mechanisms. First, different APA isoforms, especially DE-APA isoforms, may encode distinct proteins. A comprehensive human transcriptome analysis using RNA-seq found that over 5000 human genes produce APA isoforms that have differences in their coding regions, and half of these APA events showed tissue-specific profiles (Wang et al. 2008). Therefore, like alternative splicing, APA significantly expands the proteome diversity. The proteins encoded by APA isoforms may have distinct functional properties. As mentioned earlier, the IgM protein expressed from the shorter APA isoform is secreted, while the product of the longer isoform is membrane bound (Alt et al. 1980; Early et al. 1980; Rogers et al. 1980). In a recently described example, polyadenylation within a constitutively spliced internal exon of the EPRS mRNA converts a Tyr codon into a stop codon (Yao et al. 2012). The shorter EPRS protein isoform expressed from the truncated mRNAs attenuates the function of the full-length protein in translation inhibition. Furthermore, some of the APA isoforms are poorly translated or the expressed proteins are unstable. Therefore, the generation of nonfunctional APA isoforms can be viewed as a mechanism to turn off gene expression post-transcriptionally. For example, both CstF77 and HuR have been shown to autoregulate their own expression by stimulating the usage of an alternative PAS that leads to the production of truncated or unstable mRNAs (Juge et al. 2000; Dai et al. 2012; Mansfield and Keene 2012).

Secondly, APA isoforms differ in their 3’ UTRs, which can modulate the localization of the mRNAs. Many mRNAs are known to localize to specific subcellular loci from yeast to human, and the sequences required for such localization are often found in the 3’ UTRs (Kislauskis and Singer 1992). This mechanism is particularly prevalent in polarized cells, including neurons (Jung et al. 2012). Indeed, we have recently observed widespread 3’ UTR lengthening during the differentiation of mouse embryonic stem cells to neurons (Shepard et al. 2011). Two recent global analyses of APA in *Drosophila* and zebrafish showed that the mRNAs in the brain have the longest average 3’ UTRs (Smibert et al. 2012; Ulitsky et al. 2012). These studies indicate that mRNAs acquire their localization signals through APA changes during neurodevelopment. However, no global analyses of the localization patterns for APA isoforms have been reported.

Thirdly, APA isoforms may have different stabilities and/or translation efficiencies. It is well known that many RNA sequence elements in the 3’ UTR, such as the AU-rich and GU-rich elements (AREs and GREs) and microRNA target sites, can negatively regulate mRNA stability (Di Giammartino et al. 2011). Therefore, it has been suggested that shorter APA isoforms are generally more stable and produce more proteins (Sandberg et al. 2008; Mayr and Bartel 2009). More than half of the putative microRNA target sites are located downstream from the first PASs (Legendre et al. 2006), indicating widespread cross-talk between microRNA- and APA-mediated gene regulation. An EST-based global analysis showed that miR-1 or miR-124-targeted APA isoforms are down-regulated in tissues where these two microRNAs are expressed (Legendre et al. 2006). A number of examples have been reported in which APA changes lead to production of mRNAs with shorter 3’ UTRs, thereby evading microRNA-mediated inhibition (Wirsing et al. 2011; Boutet et al. 2012). Other than these sequencespecific mechanisms, the sheer length of the 3’ UTR may influence the stability of the mRNA through the nonsense-mediated mRNA decay (NMD) pathway. NMD is an RNA quality control mechanism that selectively degrades mRNAs harboring premature stop codons (Hwang and Maquat 2011). Although it is still not fully understood what RNA characteristics define NMD targets (Rebbapragada and Lykke-Andersen 2009), the 3’ UTR lengths have been suggested to be an important determinant of NMD from yeast to human (Amrani et al. 2004; Hogg and Goff 2010). According to this model, the NMD factor UPF1 associates with 3’ UTRs to promote NMD, and the level of UPF1 recruitment is proportional to the 3’ UTR length (Hogg and Goff 2010). Therefore, the APA isoforms with longer 3’ UTRs are expected to be more susceptible to NMD.
Global insights into alternative polyadenylation

Global analyses of APA regulation in human diseases

Aberrant mRNA 3‘ processing has long been implicated in human diseases (Danckwardt et al. 2008). Genetic mutations or single-nucleotide-polymorphisms (SNPs) can lead to loss or gain of PASs in functionally important genes. The resultant alterations in the expression of affected genes have been shown to cause or contribute to a number of human diseases, including thalassemia (Higgs et al. 1983; Orkin et al. 1985), Fabry disease (Bishop et al. 1988), metachromatic leukodystrophy (Gieselmann et al. 1989), amytrophic lateral sclerosis (ALS) (Lin et al. 1998), IPEX (immune dysregulation, polyendocrinopathy, enteropathy, X-linked) (Bennett et al. 2001), and facioscapulohumeral muscular dystrophy (FSHMD) (Lemmers et al. 2010). Even subtle changes to a PAS may have pathological consequences. For example, in many highly proliferative mantle lymphoma (MCL) tumors, mutations at a proximal PAS of the proto-oncogene cyclin D1 lead to higher levels of its mRNAs with shorter 3‘ UTRs, which is more stable and produces more proteins. This shift in cyclin D1 APA profile is associated with an inferior survival rate in MCL patients (Wiestner et al. 2007).

Recent studies have begun to analyze global APA changes in a number of human diseases, including cancer. Mayr and Bartel (2009) observed APA-mediated 3‘ UTR shortening for the mRNAs of a number of proto-oncogenes in cancer cell lines compared with the corresponding normal tissues and immortalized nontransformed cell lines. The APA isoforms of these proto-oncogenes with shorter 3‘ UTRs were shown to be more stable, generate greater amounts of proteins, and most importantly, lead to higher neoplastic transformation rates (Mayr and Bartel 2009). Based on these observations, it was proposed that 3‘ UTR shortening through APA plays an important role in oncogene activation in the absence of genetic mutations. Several recent studies suggest that 3‘ UTR shortening in cancers is not limited to proto-oncogenes. For example, a comparison between the immortalized but nontransformed mammary epithelial cell line MCF10A and the breast cancer cell line MCF7 detected a global APA shift toward proximal PASs in MCF7 cells (Fu et al. 2011). However, a similar shift was not observed in a different breast cancer cell line MB231 (Fu et al. 2011). A comparison of five different cancers and their corresponding normal tissues revealed a widespread increase in the relative levels of the shorter APA isoforms (Lin et al. 2012). Although APA changes seem to occur in many cancers, it remains unclear what role APA regulation plays in cancer development. A recent study suggests that 3‘ UTR shortening is more strongly associated with proliferation than with transformation induced by oncogene activation (Elkon et al. 2012). This may indicate that, among all of the hallmarks of cancer (Hanahan and Weinberg 2011), APA regulation mainly contributes to sustaining proliferation.

Global deregulation of APA was also detected in cardiac hypertrophy (the enlargement of the heart), which could occur under both physiological and pathological conditions (Park et al. 2011). A microarray-based analysis detected higher usage of the proximal PAS and overall 3‘ UTR shortening in hypertrophy compared with normal hearts (Park et al. 2011). Such APA changes are accompanied by the up-regulation of many microRNA target mRNAs, suggesting coupling between APA- and microRNA-mediated gene regulation (Park et al. 2011). 3‘ UTR shortening was also observed in oculopharyngeal muscular dystrophy (OPMD) (de Klerk et al. 2012; Jenal et al. 2012). This neuromuscular disease is caused by ‘prion’-like mutant PABPN1 proteins that have an expanded polyalanine region due to a trinucleotide (GCN) expansion in the PABPN1 gene. Recent studies suggested that the mutant PABPN1 protein, called trePABPN1, acts as a dominant-negative factor by trapping the wild-type protein in insoluble nuclear inclusion bodies (Jenal et al. 2012). Global mapping of RNA polyadenylation in trePABPN1-expressing cells and using an OPMD mouse model revealed a systematic shift of the APA profile toward proximal PASs, similar to the effect of PABPN1 loss-of-function (de Klerk et al. 2012; Jenal et al. 2012). It will be important to understand how the observed aberrant APA pattern contributes to the pathology of OPMD and why only certain muscle cells are affected by this defect.

CONCLUSIONS AND FUTURE PERSPECTIVES

Thanks to the recent wave of global studies, APA has now been recognized as a major mechanism for eukaryotic gene regulation. Regulatory mechanisms and functional impacts of APA are being revealed at an unprecedented pace. But a number of important issues remain to be addressed by future studies. First, major efforts are needed to systematically characterize the APA regulatory mechanisms and to...
compile a "polyadenylation code." For example, it remains
difficult to accurately identify PASs based on the sequence
information and to predict the "strength" of known PASs.
For splicing, the strength of a splice site can be estimated by
its complementarity with the snRNAs (Yeo and Burge
2004). Although the basic cis-elements for PASs are known,
quantification of PAS strength using sequence information
has not been adequately accomplished. Additionally, future
studies are needed to systematically identify regulatory cis-
elements, such as "enhancers" and "silencers" of polyadene-
ylation and trans-acting regulatory factors that recognize
them. A key challenge will be to integrate the various
mechanisms to achieve an understanding of APA regula-
tion at the systems level. Secondly, although widespread
APA changes have been observed under a wide variety of
physiological and pathological conditions, relatively little is
known about the specific functional consequences of these
changes. In most cases, the cause and effect relationship
between APA changes and the involved biological processes
remains to be determined, and the functionally important
APA changes remain to be identified and characterized.
Finally, as aberrant APA associates with many human
diseases, it will be important to explore whether APA can be
specifically and effectively manipulated in vivo to help
treat these diseases. Antisense oligo-based methods have
been successfully used to modulate alternative splicing of
many human genes and are being tested for clinical pur-
poses (Kole et al. 2012). A similar approach has recently
been used to modulate APA (Vorlova et al. 2011). It will be
of great interest to test whether this method can be applied
to treating diseases caused by aberrant mRNA polyadenylation.

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REFERENCES

Alt FW, Bothwell AL, Knapp M, Siden E, Mather E, Koshland M,
immunoglobulin μ heavy chains is directed by mRNAs that differ
at their 3’ ends. Cell 29: 293–301.

2004. A faux 3’-UTR promotes aberrant termination and triggers

Patterns of variant polyadenylation signal usage in human genes.
Genome Res 10: 1001–1010.

Bennett CI, Brunkowski ME, Ramsdell F, O’Briant KC, Zhu Q, Fuleihan
RL, Shigeoka AO, Ochs HD, Chance PF. 2001. A rare polyadenyla-
tion signal mutation of the FOXP3 gene (AAUAAA → AUGAA)

Bentley DL. 2005. Rules of engagement: Co-transcriptional recruit-
ment of pre-mRNA processing factors. Curr Opin Cell Biol 17:
251–256.
Global insights into alternative polyadenylation


Simpson GG, Dijkwel PP, Quesada V, Henderson I, Dean C. 2003. FY is an RNA 3′ end-processing factor that interacts with FCA to control the *Arabidopsis* floral transition. *Cell* 119: 777–787.


