

## SURVEY AND SUMMARY

# Downstream elements of mammalian pre-mRNA polyadenylation signals: primary, secondary and higher-order structures

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### ABSTRACT

**Primary, secondary and higher-order structures of downstream elements of mammalian pre-mRNA polyadenylation signals [poly(A) signals] are reviewed. We have carried out a detailed analysis on our database of 244 human pre-mRNA poly(A) signals in order to characterize elements in their downstream regions. We suggest that the downstream region of the mammalian pre-mRNA poly(A) signal consists of various simple elements located at different distances from each other. Thus, the downstream region is not described by any precise consensus. Searching our database, we found that ~80% of pre-mRNAs with the AAUAAA or AUUAAA core upstream elements contain simple downstream elements, consisting of U-rich and/or 2GU/U tracts, the former occurring ~2-fold more often than the latter. Approximately one-third of the pre-mRNAs analyzed here contain sequences that may form G-quadruplexes. A substantial number of these sequences are located immediately downstream of the poly(A) signal. A possible role of G-rich sequences in the polyadenylation process is discussed. A model of the secondary structure of the SV40 late pre-mRNA poly(A) signal downstream region is presented.**

### INTRODUCTION

Poly(A) tails are essential structural and functional elements of eukaryotic mRNAs. They are important for the regulation of mRNA stability, mRNA export from the nucleus to the cytoplasm, translation initiation and, possibly, for other cellular events (1–5). Moreover, use of alternative polyadenylation sites has been shown to be important for the regulation of gene expression (6). It has been suggested that polyadenylation signals [poly(A) signals] are required for

effective splicing, transcription termination and, possibly, even for translation termination (7–10). Thus, further study of polyadenylation mechanisms is of great importance.

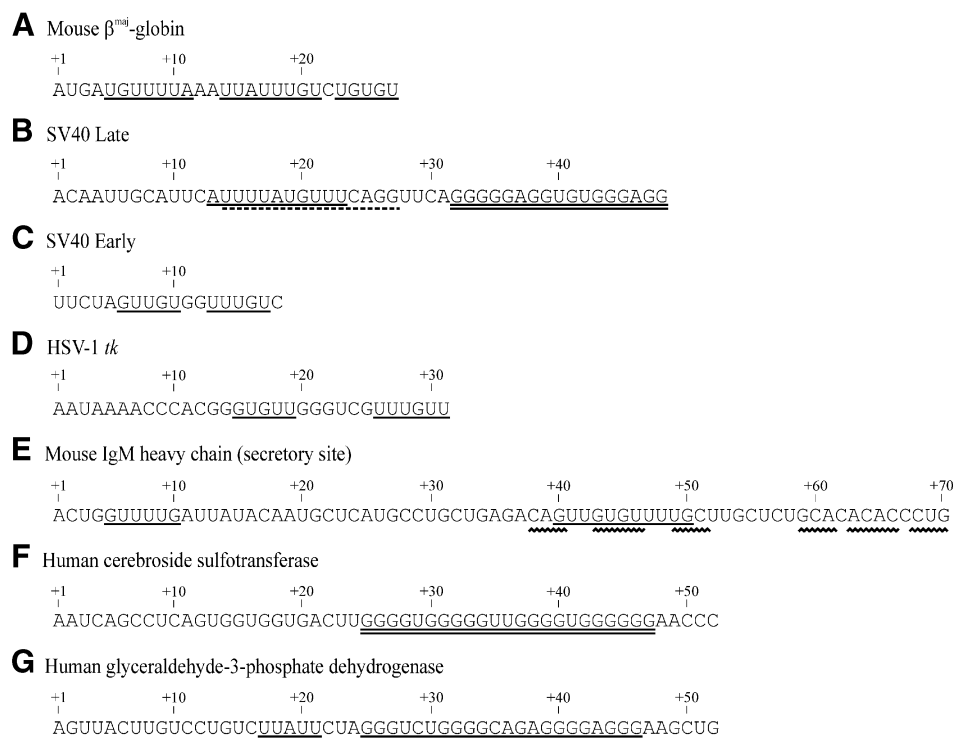
Polyadenylation of pre-mRNA is a universal modification, but different organisms use different mechanisms to carry it out (11–14). The polyadenylation mechanism of mammalian pre-mRNAs has been well studied (12,14–17). The reaction is performed by a large set of proteins and is dependent on a minimum of two elements in the pre-mRNA molecule. There is an upstream element consisting of a highly conserved AAUAAA hexamer, and a downstream element often described as a poorly conserved GU- or U-rich sequence (for example, see 18–20).

This review is devoted to analyzing the downstream elements of higher eukaryotic pre-mRNA poly(A) signals. After analyzing data from the literature, we conclude that these elements usually consist of collections of short U-rich and/or GU-rich sequences.

### PRIMARY STRUCTURE OF DOWNSTREAM ELEMENTS

The polyadenylation reaction of mammalian pre-mRNAs proceeds in two stages: the endonucleolytic cleavage of pre-mRNA and the subsequent addition of poly(A) sequence to the newly formed 3' end (12,14–17). The cleavage reaction requires the cleavage and polyadenylation specificity factor (CPSF), the cleavage stimulation factor (CstF), the cleavage factors I and II (CF I and CF II), and, in most cases, poly(A) polymerase (PAP). The addition of poly(A) requires CPSF, PAP and a third protein, poly(A) binding protein 2. The assembly of the cleavage complex, which contains most or all of the processing factors and the substrate RNA, occurs cooperatively (16). CPSF binds to the AAUAAA hexamer, CstF interacts with the U/GU-rich element located downstream of the hexamer, and these two proteins interact with each other. Pre-mRNA is cleaved at a site located between the hexamer and the U/GU-rich element. The interaction of CPSF and CF I with pre-mRNA is apparently one of the earliest events of the cleavage complex assembly. CF II has been only

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**Figure 1.** The sequences of downstream regions of pre-mRNA poly(A) signals. The four out of five base UREs and 2GU/U elements are indicated by a single line, G-rich tracts are indicated by a double line. The binding site for CstF is indicated by a dotted line. The segments which form the double-helical structure are indicated by a wavy line.

partially purified so far, and its role is not completely clear (21).

The characteristic feature of the AAUAAA element is that most single base substitutions significantly reduce the cleavage and polyadenylation efficiencies. The only exception is AUUAAA, which directs cleavage *in vitro* with an efficiency of 66%, relative to the wild-type level (22). Polyadenylation of precleaved pre-mRNAs proceeds with an efficiency of >10% in the presence of eight out of the possible 18 singly substituted variants of the AAUAAA hexamer. Recent statistical analysis of higher eukaryotic DNA databases (23–25) has revealed that the AAUAAA element is not as universal as it has been considered to be previously (15–17). According to recent data (23–25), only ~70% of human 3' expressed sequence tags (ESTs) contain one of the two optimal sequences (AAUAAA or AUUAAA). By *in silico* detection of poly(A) signals, Graber *et al.* (23) showed that the occurrence of AAUAAA variants with single base substitutions correlates with their respective processing efficiencies measured *in vitro*. The most efficiently processed variants appeared to be elements of poly(A) signals of eukaryotic pre-mRNAs (24). They are described by an NNUANA consensus, where N is any nucleotide. Thus, nucleotides at positions 3, 4 and 6 are the most conserved.

The use of suboptimal AAUAAA-like elements *in vivo* could be explained by the cooperative mode of pre-mRNA sequences recognition by the polyadenylation factors, when a weak interaction of one protein factor with the binding site can be compensated by a stronger interaction of another factor (20,23,26,27).

An exact consensus sequence for the downstream element of the poly(A) signal has not been determined. Here we suggest that no general consensus exists, but rather the downstream element consists of various simple elements located at different distances from each other.

Results of Chen and Nordstrom (28) provide a good illustration of some special features of downstream elements. The efficiency of polyadenylation of chimeric constructs containing the poly(A) signal of the mouse  $\beta^{\text{maj}}$ -globin pre-mRNA was examined. Some parts of the downstream element (Fig. 1A) were replaced with random CA-containing tracts in this signal. This replacement reduced the efficiency of 3' end processing. In particular, replacements of +5/+10, +11/+16, +17/+22 and +23/+27 tracts (Fig. 1A) decreased the processing efficiency approximately by 27, 7, 50 and 10%, respectively. This led to a conclusion that the downstream element of the poly(A) signal of the mouse  $\beta^{\text{maj}}$ -globin pre-mRNA has two major U/G-rich functional components (+5/+10 and +17/+22), and that the portions of the element are functionally redundant. The signals of other pre-mRNAs examined had similar features (26).

U-rich downstream elements have been investigated in more depth than G/U-rich tracts. Wilusz and co-workers (29–32) have shown that an oligo(U) tract composed of 5 nt could functionally substitute for the downstream region of the poly(A) signal of SV40 late (SV40 L), SV40 early (SV40 E) and Ad5 L3 pre-mRNAs (29). Moreover, pentamers consisting of four U residues and any other residue within this pentamer [a four out of five base U-rich element (URE)], were proven to be sufficient for polyadenylation of SV40 L

pre-mRNA (31,32). The authors also characterized the positional requirements for both the AAUAAA and URE motifs in the polyadenylation reaction. The cleavage was found to occur within a region 11–23 nt downstream of the AAUAAA element (with a few exceptions). The optimal position for the URE is 10–30 nt downstream of the cleavage site, but this element can be located closer than 10 nt to the cleavage site. The authors suggested that the spacing requirements for the AAUAAA and URE motifs reflect the spatial requirements for a stable interaction between CPSF and CstF (32).

Chen *et al.* (32) analyzed 131 poly(A) signals for mammalian pre-mRNAs available in the GenBank database. The four out of five base UREs, located within 30 nt downstream of the cleavage site, were present in 74% of natural signals. A UV cross-linking study showed that such elements serve as the binding sites for the 64 kDa subunit of CstF (30) (as an illustration, the CstF binding site in the SV40 L pre-mRNA is shown in Fig. 1B). Proceeding from these facts, Chen *et al.* (32) concluded that the URE is a major downstream element of poly(A) signals.

In certain instances, G/U-rich tracts are important components of the downstream element of the poly(A) signal. McDevitt *et al.* (33) showed that the GUUGUGGU sequence, which is a fragment of a natural poly(A) signal of SV40 E pre-mRNA (Fig. 1C), could substitute for the intact downstream element of this mRNA. In this case, the efficiency of the cleavage reaction was ~35% of wild type. Different single base substitutions in this fragment led both to increases and decreases in processing efficiency. In particular, an increase in efficiency occurred when either G residue in the GG dimer was substituted by U. The GUUGUUGU and GUUGUGUU variants were approximately three times more effective than the original fragment.

These results are interesting in the light of data of Takagaki and Manley (20). Using a SELEX method, they established that the isolated RNA binding domain of the 64 kDa subunit of CstF selected GU-rich sequences containing GU- and U-repeats, but not G-repeats. They also demonstrated that such sequences were specifically recognized by full-length CstF and served as downstream elements in pre-mRNA cleavage assays. These experimental data (20) allow to assume that the functional part of the GUUGUGGU fragment of the downstream element of the SV40 E pre-mRNA poly(A) signal (33) is the GUUGU portion, which does not contain G-repeats.

As mentioned above, the UGUGU fragment of mouse  $\beta^{\text{maj}}$ -globin pre-mRNA downstream element (+23/+27 tract, Fig. 1A), although not essential for polyadenylation, contributes to maximal efficiency (28). Since the replacement of this fragment with a CA tract does not affect the +5/+10 and +17/+22 fragments containing the four out of five base UREs (Fig. 1A), a true role of UGUGU as a downstream element may not be adequately reflected in this experiment. Moreover, this element is more essential in another assay where a partially mutated poly(A) signal of  $\beta^{\text{maj}}$ -globin pre-mRNA was used. In this context, deletion of the fragments containing the UGUGU element (+22/+27) or URE (+16/+22) (Fig. 1A) led to a ~20% decrease in the cleavage efficiency (28).

Other short G/U-rich sequences in the downstream region of the poly(A) signal were found in Herpes Simplex Virus type 1 thymidine kinase (HSV-1tk) pre-mRNA (34). The

replacement of a CGGGUGUU tract (Fig. 1D) in the downstream region of this pre-mRNA with a linker sequence led to a 69% reduction in polyadenylation efficiency. Based on results with other pre-mRNAs, we suggest that the functional portion of this tract may be the sequence GUGUU. The three above-mentioned tracts (GUUGU, UGUGU and GUGUU) are all possible variants of a pentamer consisting of GU dimers and a U residue. Since all these sequences play some role in the polyadenylation reaction, we propose that another simple downstream element is present along with the well known element four out of five base URE. We term it the 'two GU and one U' (2GU/U) element (12).

McLauchlan *et al.* (35) studied sequences at the 3' termini of 95 pre-mRNAs from higher eukaryotes and their viruses. They proposed that a YGUGUUY sequence (where Y is pyrimidine), located ~30 nt downstream of AAUAAA, is a consensus for the downstream element of the poly(A) signal, since 67% of the examined pre-mRNAs contained this or similar sequences. Other consensus sequences were also proposed, for example, UUGANNNUUUUU (36). Nevertheless, a convincing consensus for the downstream element was not achieved (16). It may be that there is no general consensus sequence for the downstream region of the pre-mRNA poly(A) signal, but rather this region contains a different number of various simple elements (each 5 nt long, as a minimum) located at different distances from each other.

To analyze further the structure of poly(A) signals, we collected a database of human poly(A) signals (37), which includes 244 DNA sequences randomly selected from GenBank (National Center for Biotechnology Information, USA). The sequences correspond to regions of pre-mRNAs extending 200 nt upstream of the cleavage site (or up to the 5' end of the last exon) and 200 nt downstream. The sequences downstream of the cleavage site were identified by comparing the genomic DNAs with mRNA sequences from GenBank. The sequences in our database belong to different chromosomes and are expressed in various tissues. A portion of the sequences in the database, representing 70 nt upstream and 70 nt downstream of the poly(A) site, is presented in the Supplementary Material. The full database containing the 400 nt long sequences is available from the authors upon request.

Results of our analysis are presented in Tables 1–3. We found that 69% of the pre-mRNAs analyzed contained the canonical AAUAAA hexamer. This percentage is higher than that obtained using 3' EST analysis (50–60%), but lower than the percentage obtained in earlier studies using cDNA (80–90%) (25 and references therein). Our value is similar to that obtained by Tabaska and Zhang (~74%) who used both 3' ESTs and sequences from GenBank to build a database of poly(A) signals that contains 280 mRNA and 136 DNA sequences (38). MacDonald and Redondo (25) provided some explanations why 3' EST data differ from conventional cDNA data. In particular, they noted that when a cDNA is submitted to GenBank, the most common variant of the 3' end sequence is reported, whereas the EST approach gives equal weight to every instance of a 3' end sequence, including both rarely expressed and more common variant forms of mRNAs. Thus, our database is probably biased toward more common variants of poly(A) signals. Even so, 17% of our pre-mRNAs contained neither AAUAAA nor AUUAAA hexamers. We therefore classified pre-mRNAs in our database by the type of core

**Table 1.** Core upstream elements of 244 human pre-mRNA poly(A) signals studied<sup>a</sup>

Element type	Element	Portion of pre-mRNAs containing the element (%)
I	AAUAAA	69
II	AUUAAA	14
III	NAUAAA	9
	ANUAAA	
	AAUANA	
IV	AANAAA	2
	AAUNAA	
	AAUAAN	
V	Substitutions of any two bases in AAUAAA	3
VI	Other than I–V	3

<sup>a</sup>The database of human poly(A) signals is reported in Zarudnaya *et al.* (37) and available as Supplementary Material (Table S1).

upstream element (Table 1). We defined the AAUAAA and AUUAAA hexamers as type I and II, respectively. The occurrence of the AUUAAA element is ~12–15% in both our database (Table 1) and in other studies (25,38). Some pre-mRNAs in our database contain the AAUAAA hexamer with a single base substitution at the 1st, 2nd and 5th positions (type III). As was already mentioned, these variants function in natural pre-mRNAs.

Hexamers of types I–III were found to occur as far as 16–35 nt upstream of the cleavage site (counting from the first nucleotide of the element). The distribution of distances between the hexamer and the cleavage site is skewed, with a maximum at +21. In contrast to the core upstream elements, the downstream elements were found over a much wider region, throughout the downstream region (+1/+70) (see Table S1). However, they occurred 3-fold more frequently in the +2/+22 than in the +36/+70 region. Thus, the distance between the upstream and the corresponding downstream elements observed in the most pre-mRNAs studied covers 25–50 nt.

A very small portion of the pre-mRNAs in our database (2%) contained AAUAAA hexamers with single base substitutions at the 3rd, 4th or 6th positions (type IV, Table 1). Some of these elements, for example AAGAAA or AAUGAA, are functional in some pre-mRNAs and deleterious in others (24 and references therein). The portion of pre-mRNAs containing the AAUAAA hexamer with single base substitutions other than AUUAAA (elements of types III and IV) made up 11% of the transcripts analyzed (Table 1), which is close to the value reported by Tabaska and Zhang (9%) (38).

Elements with substitutions at any two bases in the AAUAAA consensus were grouped as type V (Table 1). Information on elements of this type is very sparse in the literature. Only UAUUA has been shown to be functional. This element functions weakly in papilloma virus pre-mRNA (24 and references therein). The remaining 3% of pre-mRNAs were found to contain neither the canonical hexamer nor its variants in the –16/–35 region (type VI, Table 1), but the majority of these pre-mRNAs contain hexamers of types I–III (true signal elements) in the –38/–200 region. It could be that hexamers located in the –36/–54 region can be functional elements if the corresponding downstream elements are located very close to the cleavage site, so that the distance

between the core upstream and downstream elements does not exceed 50–55 nt.

Some hexamers located far from the cleavage site may be functional if they are brought together with the downstream element by formation of a stem-loop structure, as was shown to occur in case of the Human T-Cell Leukemia Virus Type I (HTLV-1) poly(A) signal (39). In addition, hexamers of types I–III located outside the –16/–35 region may be upstream elements of alternative poly(A) signals.

What is the possible role of non-canonical upstream elements? Hexamers of types III–V or other undetermined elements may be used preferentially in specific tissues or cell types. Wallace *et al.* (40) and Dass *et al.* (41) discovered a new form of CstF-64,  $\tau$ CstF-64, which is highly expressed in male germ cells, and to a smaller extent in brain. The authors proposed that  $\tau$ CstF-64 plays a significant role in polyadenylation of pre-mRNAs with non-AAUAAA core upstream elements in germ cells. As a pilot study, MacDonald and Redondo (25) sequenced 3' ends of 122 cDNAs from a library prepared using mouse male germ cell types and showed that 59% of these RNAs contained AAUAAA and 12% contained AUUAAA, which is similar to the frequencies obtained in other 3' EST studies (25 and references therein). Thus, the question remains open whether cell types expressing  $\tau$ CstF-64 will show a higher incidence of non-AAUAAA poly(A) signals.

We compared the occurrence of non-AAUAAA core upstream elements of poly(A) signals in 78 human pre-mRNAs from different cells (37) and 39 mammalian germ cell-expressed pre-mRNAs earlier reported by Wallace *et al.* (40). Surprisingly, the percentage of pre-mRNAs with core upstream elements of types III and V in germ cells appeared to be ~2-fold higher than in other cells, and the percentage of germ cell pre-mRNAs with hexamers of type II is ~4-fold lower than in other cells (Table 2). The portions of pre-mRNAs with hexamers IV and VI appeared to be approximately equal. These results suggest that the hexamers III and V are major non-AAUAAA upstream elements of poly(A) signals in male germ cells, which is consistent with the hypothesis of Wallace *et al.* about the significance of non-canonical upstream elements of poly(A) signals in germ cells. A more extended database of germ cell pre-mRNAs should be analyzed to support this hypothesis.



**Table 2.** Comparison of non-AAUAAA core upstream elements of poly(A) signals in 78 human pre-mRNAs from different cells (37) and 39 mammalian germ cell-expressed pre-mRNAs (40)

Element type	Portion of pre-mRNAs containing the element (%)	
	Human cells	Mammalian germ cells
II	45	10
III	23	41
IV	6	8
V	12	26
VI	14	15

The elements are located within the –16/–35 region.

**Table 3.** Downstream regions of human poly(A) signals of pre-mRNAs studied<sup>a</sup>

Element	Variant	Portion of pre-mRNAs containing downstream elements (%) <sup>b</sup>
URE	NUUUU	66–70
	UNUUU	
	UUNUU	
	UUUNU	
	UUUUN	
2GU/U	GUUGU	27–31
	UGUGU	
	GUGUU	
Two or more UREs and 2GU/U elements	–	46
Core downstream elements other than URE and 2GU/U	–	20

<sup>a</sup>The database of human poly(A) signals is reported in Zarudnaya *et al.* (37) and available as Supplementary Material (Table S1).

<sup>b</sup>204 pre-mRNAs with core upstream elements of types I and II were surveyed. Core downstream elements are located within the +1/+32 region downstream of the cleavage site.

Our findings on the structure of the downstream region of poly(A) signals of pre-mRNAs with core upstream elements I and II are presented in Table 3. As shown here, 66–70% of pre-mRNAs contain UREs, in agreement with reported data (32). A substantial portion of pre-mRNAs (27–31%) contain the 2GU/U element. The percentages of transcripts containing UREs and 2GU/U elements are given as a range because some pre-mRNAs contain tracts that cannot be unambiguously interpreted. For example, the UUUGUGU sequence in mouse cystatin 9-like (CST9L) pre-mRNA includes a signal pentamer that may be classified as both a URE (UUUGU) and a 2GU/U element (UGUGU).

Notably, Graber *et al.* (23), using computer analysis of ~900 *Drosophila* 3' ESTs, revealed that 6 nt 'words', such as UGUUUU, UGUGUU and UUUUUU, were the most common words in the region of 10–20 nt downstream of the cleavage/polyadenylation site. Though the authors studied six-letter 'words', the UGUGU and GUGUU pentamers are present. We suppose that 2GU/U elements can be significant not only in *Drosophila* pre-mRNAs but also in mammalian transcripts. The following findings support this suggestion. The AAUAAA hexamer was shown to occur in *Drosophila* pre-mRNAs approximately as often as in mammalian pre-mRNAs (23). It is therefore likely that the downstream elements of poly(A) signals are also identical in insects and

mammals. According to Hatton *et al.* (42), *Drosophila* CstF and CPSF recognize mammalian poly(A) signals, since these signals have been successfully used in vectors for expressing proteins in *Drosophila* cells (42 and references therein).

As many as 20% of analyzed pre-mRNAs with hexamers I and II contain neither UREs nor 2GU/U elements in the region between the cleavage site and the nucleotide +32. However, the majority of these contain tracts that are similar to one of these elements. For example, a sequence in the ficolin 3 pre-mRNA (UCUUC) could be a suboptimal URE. Our database analysis showed that 46% of the transcripts with AAUAAA or AUUAAA hexamer (types I and II) contained two or more UREs or 2GU/U elements. In most of these transcripts the downstream elements are separated by 2 nt or more, as seen in Figure 1D, but in some cases the elements are present in one long tandem sequence, as can be seen in Figure 1B and E. It is reasonable to suppose that the presence of multiple downstream elements increases the probability of CstF binding to pre-mRNA in the vicinity of the upstream element of the poly(A) signal, and thus the overall efficiency of cleavage complex assembly.

The frequency of UREs and 2GU/U elements in pre-mRNAs with upstream elements of types III–VI is comparable with that in pre-mRNAs with hexamers of types I and II (see Table S1 in the Supplementary Material). However, these data

are excluded from Table 3, because the pools of these pre-mRNAs in our database are not sufficient for a reliable estimation.

Earlier in our review (12) we discussed the analysis of downstream poly(A) signals performed by McLauchlan *et al.* (35) using different eukaryotic species and we also evaluated the frequency of UREs and 2GU/U elements in database of McLauchlan. Remarkably, the YGUGUUY consensus proposed by McLauchlan contains some variants of UREs and 2GU/U elements. The frequency of these elements in the database of McLauchlan is similar to that which we obtained with human poly(A) signals, which provides additional evidence that the structures of the downstream region of poly(A) signals in different species are not drastically different, at least in the case of pre-mRNAs with the canonical AAUAAA hexamer. Further investigations will be required to elucidate the sequences of the downstream elements in non-human mammalian pre-mRNAs with core upstream elements of different types.

Using a SELEX method to determine CstF-binding sites, Beyer *et al.* (43) showed that purified CstF from calf thymus and HeLa cells selected RNA ligands with one of three conserved elements: element 1 (AUGCGUCCUCGUCC), element 2a (YGUGUN<sub>0-4</sub>UUYAYUGYGU) or element 2b (UGYUN<sub>0-4</sub>AUUUACU(U/G)N<sub>0-2</sub>YCU). These tracts and their fragments could serve as downstream elements of poly(A) signals in chimeric constructs. Computer-assisted analysis of the EMBL library data showed the majority of the element 2a-like sequences to be located downstream of the coding region, and the authors postulate that this element may represent a novel consensus sequence for downstream elements. The authors also showed that CstF factors purified from different organisms preferentially selected different RNA ligands (43). The factors extracted from HeLa cells and a calf thymus predominantly selected element 1 and elements 2a/2b, respectively.

As seen from an analysis of the novel consensus 2a, it contains simple elements, four out of five base UREs or 2GU/U tracts, the number of which, as a rule, is not less than two. This lends support to our model where the downstream poly(A) signal consists of different numbers of various simple elements located at different distances from one another. The majority of variants of element 1 reported in Beyer *et al.* (43) do not contain UREs or 2GU/U pentamers, although most of them contain the GCGUU or UUCCU tracts which are homologous to GUGUU and UREs, respectively.

To conclude this section, we would like to emphasize that the precise sequence of the downstream element is, apparently, not crucial for the polyadenylation reaction. In some cases (for example, see 20,28,43) the cleavage reaction proceeds in the absence of any known downstream elements, though less efficiently. This could be explained by the properties of the RNA-binding domain of the 64 kDa subunit of CstF, which is of the RNP2/RNP1 type (44). Proteins with this domain are known to specifically recognize many different RNA sequences (44–47). Wang and Hall (47) studied the crystal structures of the first two RBDs (of the RNP2/RNP1 type) of the HuD protein in complex with fragments of AU-rich elements (AREs) which control the stability of short-lived mRNAs. The authors show that mutations at some positions in ARE severely reduce HuD

binding to the ARE, while mutations at other positions are better tolerated. By analogy, such position dependence is expected upon CstF binding to the U/GU-rich downstream elements. We suppose that in this case the key contacts occur between RBD of the 64 kDa subunit of CstF and two or three U residues located in the downstream element. The conformation of these U residues is probably optimal when they are located in the most frequently occurring elements: UGUUUU, UGUGUU and UUUUUU. In the absence of UREs or 2GU/U-pentamers, CstF, which cooperatively interacts with other polyadenylation factors, is likely to bind to other U-rich tracts, where the arrangement of U residues is less optimal.

## SECONDARY STRUCTURE OF PRE-MRNA POLY(A) SIGNALS

Studies of the role of RNA secondary structure in the recognition of the AAUAAA hexamer reveal that this element is recognized in a single-stranded form (16,48,49). Effective binding of CPSF to pre-mRNA (in nuclear extracts or *in vivo*) is also possible when a part of the hexamer is involved in a double-helical structure, provided the stability is not very high (18,50).

Not many studies on the secondary structure of poly(A) signal downstream elements are known in the literature, and the conclusions drawn from them are ambiguous. Chen and Wilusz (48) showed that chimeric RNAs containing any of the main regions of poly(A) signal (the AAUAAA hexamer, the cleavage site or the URE) in a secondary structure were cleaved very inefficiently as compared to pre-mRNAs containing a wild-type signal.

The structure of the poly(A) signal of adenovirus-2 L4 pre-mRNA, the downstream element of which contains neither UREs nor 2GU/U pentamers, was studied by Sittler *et al.* (49). The authors failed to find any clear correlation between the secondary structure of this element and its function.

Results obtained by Phillips *et al.* (19) support the conclusion that one of two downstream elements of mouse IgM secretory poly(A) signal participates in a stem-loop structure with two asymmetric internal loops. Both the stem and the internal loops are important for efficient polyadenylation. In Figure 1E we show the sequences involved in the formation of the double-stranded segments of this hairpin. As seen here, the element consisting of GU/U repeats is partially involved in the double-stranded structure, although four U residues of this element are single stranded. The GU/U element is located 39 nt, and the second element (URE) only 4 nt downstream of the cleavage site (51,52). Thus, both elements are located suboptimally: the first is rather far from the cleavage site, and the second is rather close. The authors suggest that the participation of the distal downstream element in a hairpin structure may result in better recognition of this element by polyadenylation factors, compensating for its suboptimal location.

How can CstF interact with downstream elements that participate in secondary structures? Does it interact, for example, at the moment when double-helical segments of the hairpin are opened in the process of RNA 'breathing', and one of the signal pentamers (UREs or 2GU/U tracts) becomes accessible for the polyadenylation factor? Are downstream

hairpins a target for structure modifying enzymes (RNA helicases etc.) as Phillips *et al.* (19) have suggested? The answers to these questions are still unknown. In this connection, we refer to recent results of Bléoo *et al.* (53), which suggest a functional and a spatial relationship between CstF-64 and human DEAD box protein DDX1, a putative RNA helicase.

Phillips *et al.* (19) suggest that involvement of the downstream element in a hairpin structure is not peculiar to the mouse IgM poly(A) signal. They demonstrate that the downstream regions of the hamster and the human IgM secretory poly(A) signals can also form hairpin structures with internal loops. The sequences forming the descending arms of the hairpins are highly conserved in all three cases [fig. 8 in Phillips *et al.* (19)]. The proximal segments are similar in the mouse and the hamster hairpins, while the proximal segment of the human hairpin is G-rich. The role of the highly conserved sequence in the distal segment of the hairpins is unknown. The authors noted (19) that there is another obvious candidate, besides CstF, for the role of protein binding specifically to the hairpin. This is a protein with a molecular mass of 30 kDa. The binding of this protein to mouse IgM secretory pre-mRNA depends on a 55 nt sequence encompassing the distal downstream element (51). Induction of this 30 kDa protein correlates with the increase of secretory poly(A) site usage.

The poly(A) signal of SV40 L pre-mRNA contains an auxiliary downstream element (AUX DSE) in addition to the core element (URE) (54). This element is a G-rich sequence (GRS) GGGGGAGGUGUGGG (containing neither URE nor 2GU/U). The GRS serves as the binding site for hnRNP H/H' protein (55). Interaction between this protein and the GRS stimulates polyadenylation of SV40 L pre-mRNA. It has to be noted that the full GRS in the downstream region of SV40 L pre-mRNA (Fig. 1B) is 3 nt longer than the element GRS which binds hnRNP H/H' protein.

Hans and Alwine (56) used nuclease sensitivity structure analysis techniques to determine the secondary structure of the SV40 L poly(A) signal. They found that a large part of the U-rich downstream element and a part of the AAUAAA hexamer are involved in a secondary structure, while the GRS is primarily in a single-stranded form. By replacing portions of the downstream region with unrelated sequences, they showed that the ability of this region (particularly nucleotides 2696–2700) to form double-stranded structures correlates with cleavage efficiency. This indicates that the secondary structure of the region downstream of AAUAAA has a functional significance. The mechanism by which pre-mRNA secondary structure influences the polyadenylation process is still unknown. The authors propose that the secondary structure may aid CstF interaction with the downstream element. They also make an intriguing suggestion that the RNA structure in the downstream region may have a catalytic role in the cleavage process.

Though the latter suggestion cannot be excluded, it is generally assumed that the 30 kDa subunit of CPSF is the endonuclease that cleaves pre-mRNA in the polyadenylation process, since this protein is a homolog of *Drosophila* CLP protein, which possesses endoribonucleolytic activity (17,57–60). Others suggest that the cleavage is performed by a predicted nuclease in the metallo- $\beta$ -lactamase fold of the

73 kDa subunit of CPSF (61). In both these cases catalytic function is attributed to a protein, not to RNA.

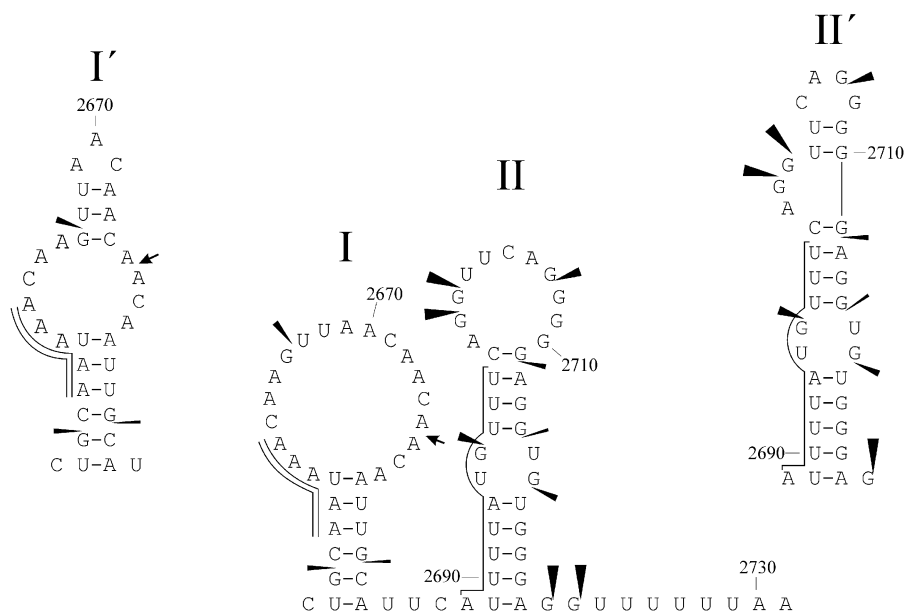
A secondary-structure scheme of the SV40 L poly(A) signal was not presented in Hans and Alwine (56) because of discrepancy between the computer-predicted structures based on programs available at that time and the nuclease sensitivity data. Inasmuch as computer programs to predict RNA secondary structure are constantly improving, we used a more recent version of the mfold program, together with the latest free energy rules (62,63) (<http://www.bioinfo.rpi.edu/applications/mfold/>) to predict the folding of the SV40 L pre-mRNA fragments. We then compared the predicted structures with the experimental results of Hans and Alwine (56).

Figure 2 shows a computer-predicted secondary structure of the SV40 L pre-mRNA fragment from nucleotides 2653 to 2731, which includes the AAUAAA hexamer, URE and GRS. This structure has two hairpins. Hairpin I includes AAUAAA and hairpin II includes the URE and the GRS. Variants of hairpins I and II (hairpins I' and II'), which have a less favorable free energy of folding, are also shown in Figure 2.

We also predicted the folding of longer fragment of the SV40 L pre-mRNA (nucleotides 2634–2731), which along with AAUAAA, the URE and the GRS contained one of three auxiliary upstream elements of the SV40 L poly(A) signal. The auxiliary upstream elements specifically interact with U1A protein (64). Hairpins I and II are also formed in the 2634–2731 nucleotide fragment. However, the formation of hairpin I was not predicted on folding the 2603–2731 nucleotide fragment containing all three auxiliary upstream elements and also the full substrate RNA (nucleotides 2531–2731) studied in Hans and Alwine (56). In both cases the AAUAAA hexamer was involved in other double-stranded structures. Hairpin II was predicted to form in one of the thermodynamically favorable variants of the secondary structure of the full substrate RNA. We suppose hairpin II may also form in much longer RNA sequences, including the full-length SV40 L pre-mRNA.

Hans and Alwine (56) used the following nucleases: RNase V1, which cleaves double-helical RNAs non-specifically; RNase T1, which cleaves at single-stranded G residues, and RNase PhyM, which cleaves at single-stranded A and U residues. The reported data (56) are consistent with the secondary structure model presented in Figure 2, provided that the variants of RNA hairpins coexist in dynamic equilibrium. As an example of nuclease sensitivity data, all the sites cleaved by RNase T1 more or less efficiently (56) are marked in Figure 2 with triangles, the sizes of which approximately correlate with the intensity of the RNA digestion.

The only discrepancy between the experimental data and the secondary structure presented in Figure 2 is the upper part of the descending arm of the hairpin II stem (nucleotides 2711–2714), which is insensitive to RNase V1. But it should be noted that insensitivity of some RNA segments to this nuclease does not necessarily mean that they are in a single-stranded form. According to Lowman and Draper (65), RNase V1 recognizes a 4–6 nt segment of sugar–phosphate backbone with an approximately helical conformation and does not require paired bases. The conformation of the GAGG segment in the upper part of the hairpin II may be suboptimal for the nuclease V1 in contrast to the conformation of the complementary UUUC segment. Consistent with our model, the



**Figure 2.** The secondary structure scheme of the core poly(A) signal of the SV40 L pre-mRNA. The secondary structures of the pre-mRNA fragments were determined using the mfold version 3.1 program for RNA folding by Zuker and Turner (<http://www.bioinfo.rpi.edu/applications/mfold/>). The U-rich downstream element and the AAUAAA hexamer are marked by single and double lines, respectively. The cleavage site is marked by an arrow. The sites attacked by RNase T1 (56) are marked by triangles. Thermodynamic stabilities of the hairpins I, I', II and II' are -3.1, -2.5, -4.3 and -2.4 kcal/mol, respectively.

GAGG tract is poorly cleaved by both single-stranded-specific nucleases.

Interestingly, some previous computer programs predicted the formation of the hairpin II without the internal loop, i.e. with all the nucleotides in the stem being paired, but as can be seen in Figure 2, this contradicts the experimental results.

Thus, our computer predictions of SV40 L poly(A) signal secondary structure and the experimental results of Hans and Alwine (56) enable us to propose that URE and the auxiliary downstream element (GRS) may be present in the ascending and descending arms of hairpin II structure, respectively. Remarkably, such functional arrangement of hairpin II of the SV40 L poly(A) signal is similar to that of the hairpin with the distant downstream element of the IgM poly(A) signal, where the GU/U-rich downstream element and the highly conserved sequence (putative auxiliary downstream element) are located in ascending and descending arms of the hairpin, respectively. It would be interesting to determine how widespread this pattern may be.

Recent studies of Arhin *et al.* (66) showed that G-rich auxiliary elements are not limited to SV40 L pre-mRNA. According to their data, ~34% of mammalian poly(A) signals contain short G-tract(s) in the region downstream of the core elements. All the tested G-rich elements bound hnRNP H/H' protein, which resulted in stimulation of polyadenylation. The affinity of hnRNP H/H' binding for various poly(A) signals was significantly different, which affected the abilities to stimulate 3'-end processing. These authors suggested that downstream G-rich tracts are common auxiliary elements of poly(A) signals (66).

Protein hnRNP H/H', which binds to GRSs, belongs to the H subfamily of hnRNP proteins, which also includes hnRNP F

and hnRNP 2H9 (67,68). HnRNP H and H', which are 96% identical to each other (67), and hnRNP F, contain three RNA-binding domains of the quasi-RNP2/RNP1 type, while hnRNP 2H9 contains two such domains. Caputi and Zahler (69) showed that all the members of the H group specifically interact with RNA sequences containing a GGGGA tract, whereas only hnRNP H/H' recognizes a GGGGGC sequence. HnRNP H/H' is likely to recognize the GGGU sequence as well (70).

Considering these facts, it can be assumed that the other members of the H group of hnRNP proteins may also bind to G-rich auxiliary downstream elements, since most of auxiliary downstream elements, which are reported in Arhin *et al.* (66), contain the GGGA tract(s) and the rest contain the GGGU or GGGGGC tracts. The role of hnRNP 2H9 binding to the G-tracts in the polyadenylation process is currently unclear. hnRNP F was shown recently to be able to diminish pre-mRNA 3'-end processing (71). The authors supposed that the binding of hnRNP F or H'/F heterodimer to a poly(A) signal near the core downstream element may inhibit the association of CstF with pre-mRNA and/or the cleavage reaction.

All the members of the H group of hnRNP proteins also participate in splicing regulation (68–70,72–75), the mechanism of their functioning in this process is not yet determined. In some cases, hnRNP H may sterically hinder the binding of SR proteins to splicing regulatory sequences, thereby inhibiting their functions (70,73). In other cases (72,74,75) the interaction of hnRNP H protein with its binding site was required for assembly of other proteins onto splicing regulatory elements, and the protein was essential for function of these elements. HnRNP H/H' was suggested to play a similar role in the polyadenylation process. Arhin *et al.* (66) suggested that it



directly interacts with CstF to stimulate assembly of general polyadenylation factors on the core poly(A) signal. Alternatively, the authors suggested that the protein may alter the structure of the nascent transcript to present the core signal elements to the general polyadenylation factors in a productive fashion. In this connection, interestingly, the RNA-binding sites for hnRNP H/H'/F proteins may form G-quadruplexes, which we discuss in the next section.

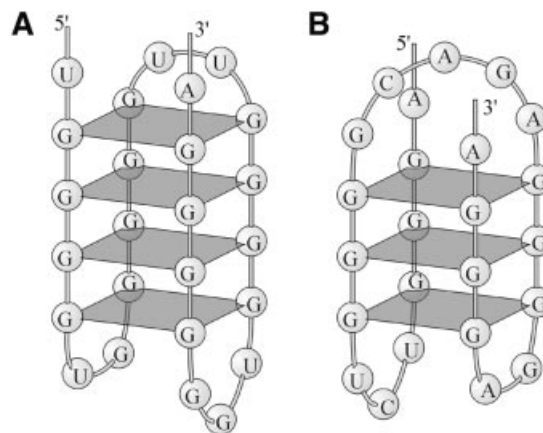
### G-QUADRUPLLEXES AS POSSIBLE AUXILIARY ELEMENTS OF POLY(A) SIGNALS

DNA or RNA sequences consisting of clusters of G-repeats may form four-stranded structures composed of stacked G-tetrads, so called G-quadruplexes (76). The thermodynamic stability of G-quadruplexes depends on the number of G-tetrads, the sequence of the loops connecting G-tetrads, and the size of the loops (77–79). Based on these data and other experimental findings on the formation of G-quadruplexes, we previously reported that ~27% of pre-mRNA sequences in our database of human poly(A) signals have a potential to form such structures (37). A portion of the article (37) is presented as Supplementary Material. In particular, this article discusses G-quadruplex formation and shows that the G-rich region of the SV40 L poly(A) signal is likely to be capable of forming G-quadruplexes. The schematic structures of G-quadruplexes formed by the full GRS of SV40 L pre-mRNA and possible quadruplexes adopted by GRSs of several human pre-mRNAs available in our database are also presented as Supplementary Material (Fig. S1). Here we present the schematic illustrations of two other putative quadruplexes of human pre-mRNAs. The first is formed by a GRS located 24 nt downstream of the cleavage site of human cerebroside sulfotransferase pre-mRNA (Figs 1F and 3A). The second quadruplex is formed by the fragment located 3 nt downstream of URE of human glyceraldehyde-3-phosphate dehydrogenase pre-mRNA (Figs 1G and 3B).

Interestingly, the sequences with a potential to form G-quadruplexes occur in pre-mRNAs of our database ~5-fold more often in the +1/+70 region (11% of pre-mRNAs) than in the –1/–70 region (2%), while in the +71/+200 and –71/–200 regions they occur approximately equally (8 and 7%, respectively). The frequency of the clusters of G-repeats in the region located immediately downstream of the cleavage site is unlikely to be fortuitous and may be evidence that G-quadruplexes play some role in the polyadenylation process.

It is also notable that hnRNP H, H' and F, besides binding to different G-rich tracts, are capable of binding to poly(G) at 2 M NaCl (80). At this ionic strength (under equilibrium conditions), poly(G) is a completely four-stranded macromolecule (81). Certain proteins are known to recognize G-quadruplexes, facilitate their formation, or disintegrate them (82). Some of them can bind specifically both to four-stranded and single-stranded forms of polynucleotides (83). It is possible that H, H' and F proteins can also recognize both single-stranded G-rich tracts and G-quadruplexes and/or promote the formation of four-stranded structures.

What functions could G-quadruplexes fulfill in the polyadenylation process? Consideration of how auxiliary elements may function helps to answer this question. Chen and Wilusz



**Figure 3.** Patterns of pre-mRNA fragments folding into G-quadruplexes. (A) Fragment of human cerebroside sulfotransferase pre-mRNA. (B) Fragment of human glyceraldehyde-3-phosphate dehydrogenase pre-mRNA.

(48) have proposed three possible mechanisms of stimulation of the polyadenylation reaction by auxiliary DSEs. First, they suggest that some of these elements may promote processing efficiency by maintaining the elements of the core poly(A) signal in an unstructured form, which enables the general polyadenylation factors to assemble efficiently on the pre-mRNA molecule. Secondly, the authors suggest that auxiliary DSEs may form stable structures which prevent CstF from sliding along the pre-mRNA, and thereby limit the region of interaction with the transcript to the downstream region of the poly(A) signal only. It was shown that a pseudoknot from a viral RNA can functionally substitute for the undetermined auxiliary DSE of the Ad5 L3 pre-mRNA poly(A) signal. A mutated pseudoknot sequence had no effect on polyadenylation efficiency. Thirdly, the authors propose that the interaction of specific proteins with auxiliary DSEs, as in the case of hnRNP H/H', may enhance the efficiency of 3'-end processing, probably by stimulating an assembly of the general polyadenylation factors on the pre-mRNA.

Four-stranded structures stabilized by G-tetrads could promote all three above-mentioned functions of auxiliary DSEs. First, a G-quadruplex, like a pseudoknot, could prevent CstF migration from the U/GU-rich downstream element. This could stimulate the polyadenylation process. In addition, G-quadruplexes could maintain the core poly(A) signal in an accessible conformation. Some GRSs form highly stable intramolecular quadruplexes that can limit a number of possible conformations adopted by RNA and thereby maintain the poly(A) signal in an optimal conformation. Finally, G-quadruplexes could serve as the binding sites for proteins that influence the efficiency of the polyadenylation process. The GRSs in regions of pre-mRNAs upstream of the cleavage site may function in the same manner as GRS located downstream, in particular, by interacting with auxiliary proteins.

It should be noted that the formation of the G-quadruplex in the SV40 L poly(A) signal may ensure opening of the hairpin with the URE (Fig. 2) and access of the CstF to the downstream element.

In conclusion, we would like to mention other possible functions of GRSs. They may function as specific signals not only in pre-mRNAs, but also in the corresponding DNAs that encode these GRS-containing pre-mRNAs. It has been shown that tandemly arranged GRSs ...CTGGCCTTGGGGG-AGGGGGAGGC... (which are the binding sites for the transcription factor MAZ), located downstream of the poly(A) signal in chimeric constructs, specifically paused RNA polymerase II and stimulated polyadenylation *in vitro* (84). We note that the binding site for MAZ being in the single-stranded form (for example, in the transcription bubble) might form quadruplex structures with two G-tetrads and this structure could induce pausing of RNA polymerase.

## SUPPLEMENTARY MATERIAL

With permission of the journal *Biopolymery i Kletka*, a portion of the text of the article (37), one table and one figure are available as Supplementary Material at NAR Online. The portion of the article reprinted here concerns some trends in G-quadruplex formation. Table S1 contains the database of poly(A) signals of 244 human pre-mRNAs. Figure S1 displays the schematic structures of possible quadruplexes for several human pre-mRNAs.

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