Recognition of GU-rich polyadenylation regulatory elements by human CstF-64 protein

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Vertebrate polyadenylation sites are identified by the AAUAAA signal and by GU-rich sequences downstream of the cleavage site. These are recognized by a heterotrimeric protein complex (CstF) through its 64 kDa subunit (CstF-64); the strength of this interaction affects the efficiency of poly(A) site utilization. We present the structure of the RNA-binding domain of CstF-64 containing an RNA recognition motif (RRM) augmented by N- and C-terminal helices. The C-terminal helix unfolds upon RNA binding and extends into the hinge domain where interactions with factors responsible for assembly of the polyadenylation complex occur. We propose that this conformational change initiates assembly. Consecutive Us are required for a strong CstF–GU interaction and we show how UU dinucleotides are recognized. Contacts outside the UU pocket fine tune the protein–RNA interaction and provide different affinities for distinct GU-rich elements. The protein–RNA interface remains mobile, most likely a requirement to bind many GU-rich sequences and yet discriminate against other RNAs. The structural distinction between sequences that form stable and unstable complexes provides an operational distinction between weakly and strongly processed poly(A) sites.

Keywords: nuclear magnetic resonance/polyadenylation/RNA-binding proteins/RNA processing/RNA recognition

Introduction

Maturation of mRNA precursors is a constitutive process for most transcripts but is also a major regulatory event in eukaryotic cells, producing multiple isoforms of the same mRNA by alternative processing. Alternative splicing has long been recognized as a major source of diversity in gene expression, while polyadenylation has been considered largely constitutive. However, ~30% of all human mRNAs contain alternative polyadenylation signals (Beaudoin et al., 2000). Furthermore, an increasing number of developmental and differentiation decisions are now known to be executed by alternative polyadenylation of the same mRNA (Barabino and Keller, 1999; Zhao et al., 1999). Differential usage of alternative poly(A) sites produces mature transcripts with distinctive 3′-UTRs and gene expression profiles and, sometimes, even different coding sequences.

In polyadenylation, as in splicing, the efficiency with which different processing sites are utilized is controlled by regulatory cis-acting RNA elements and their interactions with trans-acting protein factors. In some cases, of which UIA is the best understood example (Boelens et al., 1993), the timing and efficiency of polyadenylation decisions are controlled by gene-specific factors and their interaction with specific RNA elements and with the polyadenylation machinery. More puzzling are situations where poly(A) site selection is controlled instead by altering the intracellular levels of general processing factors such as CstF-64 (Takagaki et al., 1996; Colgan and Manley, 1997; Takagaki and Manley, 1998). Despite its obvious importance, structural and mechanistic insight on how alternative mRNA processing decisions are executed remains sparse.

In higher eukaryotes, the core polyadenylation signal comprises two elements in addition to the site where the mRNA is cleaved [often a CpA dinucleotide (Figure 1)]; the highly conserved AAUAAA hexamer 10–30 nucleotides upstream of the cleavage site, and a more diffuse GU-rich downstream sequence element (DSE) (Wahle and Keller, 1996; Colgan and Manley, 1997). These RNA sequences are specifically recognized by two protein complexes that assemble onto the pre-mRNA to form the cleavage complex (Figure 1, top). The poly(A) signal (AAUAAA) is recognized by the cleavage and polyadenylation specificity factor (CPSF) through its 160 kDa component (CPSF-160). The DSE is recognized by the heterotrimeric cleavage and stimulation factor (CstF) through its 64 kDa subunit CstF-64 (Beyer et al., 1997; Takagaki and Manley, 1997). Both CPSF and CstF bind RNA weakly, but form a strong cooperative complex when bound together to the same pre-mRNA (Bienroth et al., 1993). Enzymatic factors responsible for the phosphodiesterase and polymerase activities (CF I, II, PAP etc.) are then recruited to the CstF–CPSF–pre-mRNA complex. Because the AAUAAA sequence is very highly conserved, the relative strength of competing poly(A) sites is defined primarily by the distance between cis-acting sequences (AAUAAA and DSE) and the affinity of the DSE–CstF interaction. However, how the affinity and positioning of the CstF–pre-mRNA interaction regulate assembly of the 3′-end processing machinery remains unclear.

In addition to recognizing the pre-mRNA and each other, CPSF and CstF also link polyadenylation with other RNA processing events and with transcription. For example, CstF-64 binds PC4, a factor involved in transcriptional termination, and symplekin, a probable architectural component of the polyadenylation apparatus (Takagaki and Manley, 2000; Calvo and Manley, 2001). CstF-77 mediates the interaction between CstF and CPSF.
and also interacts with CstF-50, which in turn binds to the CTD of RNA polymerase II and to the DNA repair factor BARD1 (Kleiman and Manley, 1999, 2001). At least some of the polyadenylation factors are recruited to the transcription machinery on the promoter (Dantone et al., 1997) and a functional poly(A) signal is necessary for transcriptional termination (Proudfoot et al., 2002).

At present there is little structural information on any of the components of the polyadenylation apparatus or their RNA and protein interactions, the exceptions being the catalytic domain of poly(A) polymerase (Bard et al., 2000; Martin et al., 2000) and the U1A system (Varani et al., 2000). As a first step towards characterizing the polyadenylation reaction and its regulation at the structural level, we have studied how CstF-64 binds G/U RNA DSEs. CstF-64 is a multidomain protein (Figure 1, bottom) that is both necessary and sufficient for DSE recognition by CstF (Beyer et al., 1997; Takagaki and Manley, 1997). Immediately following the RNA-binding region at the N-terminus of the protein is the hinge domain responsible for the interaction with CstF-77 (Hatton et al., 2000) and with symplekin (Takagaki and Manley, 2000). The C-terminal half of the protein contains two Pro/Gly-rich domains separated by 11 repeats of the MEARA sequence proposed to form a long α-helix (Takagaki et al., 1992). The final 100 amino acids bind the transcriptional coactivator PC4 (Calvo and Manley, 2001). Thus, recognizable protein domains are separated by regions of low sequence complexity; CstF-64 very likely possesses a ‘bead-on-a-string’ structure. Because of this structural characteristic and because different interactions of CstF-64 map to specific regions of the protein, it is possible to study different domains of CstF-64 and their interactions in isolation. Therefore we have characterized the structure and RNA interaction of the RNA-binding domain of CstF-64 using NMR. Because recognition of G/U-rich elements by CstF-64 is an early event during assembly of the cleavage and polyadenylation complex and its regulation, the data presented here also provide insight into how polyadenylation is executed and regulated.

### Results

#### Structure of CstF-64 N-terminal RNA recognition motif

The solution structure of the N-terminal RNA-binding domain human CstF-64 was determined by NMR as described in Materials and methods. Experimental and structural statistics are summarized in Table I and a superposition of converged structures is displayed in Figure 2A. The domain adopts the classical RNA recognition motif (RRM) fold (Nagai et al., 1990; Varani and Nagai, 1998) augmented by two additional features. A long C-terminal α-helix (three turns) lies across the β-sheet while a shorter N-terminal helix (two turns) precedes the first β-strand. The first nine and last six residues of the construct are unstructured. The protein is highly stable. For example, the hydroxyl proton of Tyr36 is observable even at pH 6.0 and 300 K, providing a large number of experimental restraints.

The short α-helix at the N-terminus of the protein is reminiscent of the shorter 310 helix observed at the N-terminus of the first RRM of human hnRNP A1 (Shamo et al., 1997; Xu et al., 1997). The CstF-64 N-terminal helix has few contacts with the rest of the domain and its formation is probably driven by intrinsic residue propensities. In contrast, helix C (beginning at Ser94 and ending at Gly105) runs nearly perpendicular to the main β-sheet, masking the RNA recognition surface (Figure 2A). Hydrophobic residues within this helix (Leu101 and Leu104) and the side-chain of Glu100 form a small hydrophobic core with the aromatic residues within the RNP1 and RNP2 motifs (Phe19 and Phe61) (Figure 2B and C). Helix C is further stabilized by a hydrogen bond between the side-chain amide of Asn97, in the first turn of the helix, and the backbone carbonyl

<table>
<thead>
<tr>
<th>Table I. Experimental NMR and structural statistics for CstF-64(1–111)</th>
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*aObtained from Cα, Cβ and Cγ chemical shifts using the program TALOS.

*bMinimum and maximum values within 20 converged structures.

**Pairwise among the 20 converged structures.**
oxygen of Asn91. Each of these residues is conserved in vertebrates at least and conservatively substituted in Drosophila melanogaster as well (although not in plants and yeast), suggesting very strongly that helix C is a conserved structural feature at least for vertebrate CstF-64 proteins (Figure 3). These positions within CstF-64 have not been probed by site-directed mutagenesis with respect to the RNA binding activity of the protein or its function; our structure identifies these residues as interesting targets for future studies.

The presence of a C-terminal helix in an RRM was first reported for U1A (Avis et al., 1996) and the C-terminus of several RRM proteins folds into a helical conformation upon RNA/DNA binding: Pab (Deo et al., 1999), Sxl (Handa et al., 1999), HuD (Wang and Tanaka-Hall, 2001), nucleolin (Allain et al., 2000) and hnRNP A1 (Ding et al., 1999). CstF-64 resembles U1A and not these other proteins, but the interaction between the C-terminal helices and the β-sheet differs between CstF-64 and U1A (Figure 4). In U1A, helix C mainly contacts the edges of the β-sheet; in CstF-64, it is perpendicular to the RRM. Most importantly, aromatic residues of RNP1 and RNP2 are considerably more buried in CstF-64 than in U1A. The implications of this result for RNA binding are discussed later.

RNA recognition
CstF-64 plays a key role in polyadenylation by recognizing the DSEs. However, these sequences lack strong sequence conservation, being simply characterized by an abundance of Gs and Us. Therefore CstF-64 must somehow bind a class of sequences without strong preference for any particular sequence, and yet must also discriminate effectively against other RNAs. Thus, CstF-64 appears to provide an example of RNA recognition distinct from that of other well studied members of the superfamiliy (U1A, Pab or Sxl) that bind a distinctive sequence with high selectivity. Consistent with the functional requirements, in vitro selection experiments have demonstrated that CstF-64 binds a wide range of GU-rich sequences with comparable affinity, but not G/A or C repeats for example.
yet no unique highly specific target sequence could be identified (Beyer et al., 1997; Takagaki and Manley, 1997). A similar study of the complete CstF complex also

identified U-rich stretches ubiquitously among sequences selected to bind with high affinity, and GU dinucleotides as lower-efficiency ligands (Beyer et al., 1997; Takagaki and Manley, 1997). How does CstF-64 recognize sequences that are rich in Us and Gs without requiring a strict consensus, and yet discriminate so effectively against other RNAs? To address this question, we elected to compare a number of GU-rich sequences based on the results of the selection experiments. A compilation of all sequences studied is provided in Table II.

The NMR experiments confirmed that the RNA recognition surface of CstF-64 is the β-sheet (Figure 5) and allowed us to divide the sequences studied into two groups. The first group includes sequences containing exclusively GU repeats which tend to bind CstF-64 with lower affinity (μM) (Takagaki and Manley, 1997), i.e.

<p>| Table II. GU-rich RNA sequences studied in the present work |</p>
<table>
<thead>
<tr>
<th>No.</th>
<th>Sequence</th>
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<th>Affinity</th>
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<td>7</td>
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</tr>
<tr>
<td>2</td>
<td>GUGUGUGUGU</td>
<td>8</td>
<td>Weak</td>
</tr>
<tr>
<td>3</td>
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<tr>
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<td>GUGUGUGUGUGUGU</td>
<td>10</td>
<td>Strong</td>
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<td>5</td>
<td>GUGUGUGUGUGUGUGU</td>
<td>11</td>
<td>Strong</td>
</tr>
<tr>
<td>6</td>
<td>GUGUGUGUGUGUGUGUGU</td>
<td>12</td>
<td>Strong</td>
</tr>
<tr>
<td>7</td>
<td>GUGUGUGUGUGUGUGUGUGU</td>
<td>13</td>
<td>Strong</td>
</tr>
</tbody>
</table>

The annotation on binding affinity is based on the NMR observations reported here.

![Fig. 5](https://example.com/figure5.png)

**Fig. 5.** [1H-15N]HSQC spectra of free CstF-64 (black) superimposed on spectra of 1:1 complexes of the same domain bound to (A) (GU)_4 (red) and (B) (GU)_4UG (red). Chemical shift changes upon complex formation are shown for the two complexes in (C) and (D), respectively. (E) Low-affinity sequences form complexes with fast off rates on the NMR chemical shift timescale; only one set of signals migrating from free to complex chemical shift is observed during titrations (highlighted in green). (F) High-affinity complexes with much slower off rates display instead two sets of signals at substoichiometric protein-to-RNA ratios.
(GU)_nG and (GU)_n. The second group contains at least a pair of consecutive Us within the GU-rich repeat and bind RNA tighter. The presence or absence of two or more consecutive uracils has a remarkable effect on CstF-64 binding. For example, GUGUGUGU belongs to the ‘weak’ class and titrates in the NMR fast-exchange regime, resulting in dissociation that is faster than \(10^{-3}\) \(s\). In contrast, UGUUGUUU (very similar to the previous sequence except for the presence of two consecutive Us in the 3’ end) forms a much more stable complex with off rates slow enough to allow the observation of two sets of signals for free and bound protein during the titration (Figure 5). The NMR spectra of the first class of complexes display minor changes scattered across the entire RNA-binding surface, suggesting diffuse and perhaps non-specific binding. Members of the second class of sequences induce much larger changes in the position of the NMR signals as well as in their linewidth upon binding (Figure 6).

When RNA sequences from the second class bind CstF-64, the entire helix C (residues 94 onwards) displays sharper linewidths as it unfolds (see below); at the same time most of the \(\beta\)-sheet signals exhibit broad lines in the complex (Figure 6B). Broadening is particularly severe for clusters of residues in strands \(\beta_2\) and \(\beta_4\) (e.g. 45–48, 54, 57–59) and in the loop between \(\beta_4\) and the C-terminal helix (88, 91, 93 and 94). It undoubtedly reflects local motion in the microsecond to millisecond timescale that is only present in the complex and not in the free protein. Neither line broadening nor helix C unfolding was observed in the complexes with the first two sequences.

The dynamic behaviour of CstF-64 is unusual. For example, in U1A quenching of motion occurred upon binding (Mittermaier et al., 1999). However, it may be a functionally important characteristic; it may be critical for the ability of CstF-64 to recognize a wide range of sequences rich in Us and Gs without specifically binding a single well defined sequence or consensus. Slow binding kinetics indicative of tight and specific binding are also uncommon in single-RRM proteins, except for U1A (Allain et al., 1996). The RNA complexes of RRMs derived from hnRNP D0 (Nagata et al., 1999), U2AF65 (Ito et al., 1999) and HuC (Inoue et al., 2000) all show fast exchange behaviour, consistent with complexes with short lifetimes. Stable complexes with longer lifetimes are achieved in many cases by cooperative binding of two RRMs to single-stranded RNA–DNA (e.g. Sxl, Pabp, HuD and HnRNP A1). The single RRM from CstF-64 binds G/ U-rich sequences tightly enough to display slow kinetics, unlike all other known examples of single RRMs binding to single-stranded RNA.

Taken together, our results strongly suggest that CstF-64 has a high-affinity binding pocket that specifically recognizes a pair of consecutive uracils. Nucleotides flanking the UU anchor point contact other regions of the protein surface while retaining significant mobility on the microsecond–millisecond timescale. Those other contacts fine tune the protein–RNA interaction and provide different affinities for distinct GU-rich elements. These observations have several implications for the analysis of the intrinsic strength of different poly(A) sites (see below).

**Recognition of UU dinucleotides by the CstF-64 RRM**

The NMR and binding data discussed above highlight the importance of UU recognition by CstF-64. Therefore dissecting how the dinucleotide is recognized by CstF-64 is important to understanding how the protein binds RNA. However, the broadened NMR spectra, a reflection of the diffuse specificity of this protein, have so far made it impossible to observe many intermolecular nuclear Overhauser effects (NOEs) at the protein–RNA interface and thereby to determine the structure of the complex, despite the exploration of a wide range of conditions and temperatures. Fortunately, we were able to use the remarkable sequence and structural homology between CstF-64 and HuD to define the UU binding pocket. HUDs are neuronal expressed proteins that bind AU-rich elements within the 3’-UTR of certain mRNAs using the first two of three highly conserved RRMs, and affect the stability and half-lives of such mRNAs (Chen and Shyu, 1995).

The superposition of CstF-64 (1–111) on RRM2 of HuD (Wang and Tanaka-Hall, 2001) reveals how similar these proteins are (Figure 7A), even in the orientation of the variable loops, with one significant exception. The C-terminal helix of CstF-64 (green) is located exactly where the uracil bases are placed in the HuD–c-fos complex. In the HuD complex, uracil bases make several base-selective intermolecular hydrogen bonds with the

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**Fig. 6.** Binding of GU-rich RNAs containing at least two consecutive uracils produces large changes in the CstF-64 NMR spectra. (A) Chemical shift changes mapped onto the CstF-64 structure: amino acids showing the largest changes (>200 Hz) are in red, while less strongly perturbed sites (>100 Hz) are in green. (B) RNA binding leads to changes in the linewidth of certain regions of the CstF-64 NMR spectrum: amides whose signals become sharper are coloured green, while amino acids whose signals broaden upon RNA binding are identified in red (larger effects) and orange.
side-chains of Asn126, Arg155 and Lys201 and the backbone of Ala203 (Figure 7B). Arg46 in CstF-64 is equivalent to Arg155 in HuD and is likely to play a similar role in the recognition of the 3’-uracil, while Ser17 is analogous to Asn126 in HuD. The aromatic side-chains from RNP-1 and RNP-2 that position helix C in CstF-64 (Phe19 and Phe61) have the same conformation as Tyr128 and Phe170 in the HuD complex (Figure 7B). Since these two amino acids in HuD form intermolecular stacking interactions with the two uracils, Phe19 and Phe61 appear to be prepositioned in CstF-64 for RNA binding by the long-range interactions with helix C. Asn91 in CstF-64 makes a very important hydrogen bond with the Asn97 side-chain within the helix. This interaction is lost upon RNA binding because of the unfolding of the helix, allowing base-specific contacts analogous to those observed in the HuD complex to be formed instead.

Having identified the UU binding pocket, we used the homology between HuD and CstF-64 to build a model of the complex between CstF-64 and a UU dinucleotide. The structure of the model was calculated (see Materials and methods) by combining our NMR data on the CstF-64 complex with putative intermolecular interactions identified from the sequence and structural homology between HuD and CstF-64. A superposition of 10 calculated model structures is shown in Figure 8. According to our model, the 5’-uracil is recognized specifically by base-selective hydrogen bonds between its O4 carbonyl and the Ser17 side-chain and between its O2 carbonyl and the Arg46 side-chain. Together, these two interactions discriminate against cytosine; G and A are also discriminated against on the basis of their different size in addition to those hydrogen bonds. In the HuD–c-fos structure, the NH of the equivalent uracil base forms a water-mediated hydrogen bond; the model would accommodate this feature as well. Concerning the second position, the equivalent uracil in HuD–c-fos is specifically recognized by hydrogen bonds between the Lys201 side-chain and the O4 carbonyl group and by a main-chain Ala203 carbonyl to base NH contact. In our model, Asn91 in CstF-64 (once its side-chain is released following unfolding of helix C) plays the same role as Lys201 in HuD by recognizing the O4 carbonyl, while an equivalent main-chain contact is provided to the uracil NH. Cytosine and adenine bases would not be able to form equivalent hydrogen bonds; guanine could form the same hydrogen bonds but would be discriminated on the basis of its different size. A reorganization of the protein–RNA interface and a subsequent loss in affinity would be required to accommodate G instead of U at this position.

Discussion

We have determined the structure of the CstF-64 RNA-binding domain and studied its interaction with GU-rich sequences mimicking downstream polyadenylation regulatory elements. Binding of CstF-64 to these GU-rich elements defines a distinct example of recognition from all other well studied cases where RRM proteins bind a well defined sequence. However, it is also distinct from non-specific binding, since the protein discriminates effectively between GU-rich and random sequence RNA. The observation of a conformational change at the C-terminus of the domain and the different binding and kinetic
characteristics of complexes involving different GU-rich sequences are most informative of the mechanism by which polyadenylation is regulated.

**CstF-64 binding to GU-rich sequences represents a new example of RNA recognition**
The function of CstF-64 requires it to bind sequences characterized only by a preponderance of Gs and Us, and yet to discriminate effectively against all other RNAs. By comparing a number of RNA sequences characterized by different binding affinity, we have shown that the protein achieves this distinct specificity profile through a binding pocket for GU dinucleotides highly homologous to the HuD–UU interface (Wang and Tanaka-Hall, 2001). Contacts outside this pocket fine tune the protein–RNA interaction and provide different affinities for distinct GU-rich elements, and therefore are likely to be critical for the regulatory activity of this protein. However, unlike HuD, CstF-64 lacks a second RRM that helps define a tight and specific complex. Unlike all other single RRMs studied so far (of which U1A is an exception), CstF-64 binding to G/U-rich sequences leads to the formation of stable complexes with slow off rates. Remarkably, the protein retains extensive mobility on the microsecond–millisecond time-scale at its RNA interface, though not in its RNA-free state. This is the opposite of what we observed for U1A where slow motional processes at the RNA interface disappeared upon RNA binding, leading to a rigid (and highly specific) interface (Mittermaier et al., 1999). Motion at the interface is very likely to be functionally important by providing CstF-64 with the ability to recognize a wide range of sequences rich in Us and Gs without specifically binding a single well defined sequence or consensus. Therefore it is intrinsic to its functional requirement to bind all DSEs and yet to discriminate against non-GU-rich sequences.

**Does the conformational change at the C-terminus of CstF-64 provide a signal for assembly of the polyadenylation apparatus?**
The location of helix C within CstF-64 suggests that unfolding on GU-rich RNA binding may play a role in controlling formation of the precleavage complex. Both CPSF and CstF are preassembled and recruited to the CTD early during transcription (Dantontel et al., 1997). These two protein complexes must somehow be ‘activated’ to a conformation capable of interacting with each other and then to recruit cleavage factors and poly(A) polymerase once the poly(A) site is transcribed. The hinge domain of CstF-64 immediately follows the N-terminal RRM and includes helix C; it interacts with symplekin, an architectural component of the polyadenylation machinery, and with CstF-77 (Hatton et al., 2000). This last protein interacts with CPSF (Takagaki and Manley, 2000). If helix C is also present within the assembled CstF complex, as is very likely given the high stability of the structure presented here, then it must unfold for G/U binding to occur and before the precleavage complex is assembled. The conformational change could affect the CstF-64 interaction with CstF-77 and symplekin, and potentially the CstF–CPSF interaction as well. In this model, the C-terminal helix would act as a molecular switch in gating assembly of the polyadenylation complex by communi-
cating the signal that RNA is bound to CPSF and to the transcription machinery. Whether the signal is transmitted by a rigid-body-like motion or the maturation of the cleavage complex is promoted by an early destabilization event, helix C unfolding upon RNA binding is a good candidate to trigger assembly.

Consistent with the essential functional role suggested by our model, helix C is absolutely conserved in vertebrates, and residues positioning helix C with respect to the RRM (Asn91, Asn97, Glu100, Leu101 and Leu104) are also conserved or conservatively substituted in _D.melanogaster_ (Figure 3). Although sequence conservation does not extend to plants and fungi, it is worth remarking that the mechanism of polyadenylation differs somewhat between yeast and vertebrates and involves differences in both cis-acting sequences and trans-acting factors (Wahle and Keller, 1996; Zhao et al., 1999).

The presence of a helical appendix to the RRM was first considered an anomaly (Howe et al., 1994; Avis et al., 1996), but every RRM studied so far is augmented by an α-helix at its C-terminus, usually in the presence of RNA. However, the helices play different structural and functional roles in different proteins. The C-terminal helix of CstF-64 does not participate directly in RNA recognition but rather prevents it by occluding the RRM surface. In contrast, the C-terminal α-helix of U1A does not unfold and is repositioned upon RNA binding through a small number of changes in the protein backbone to form protein–RNA and protein–protein contacts (Allan et al., 1996; Varani et al., 2000). The behaviour of helix C is also completely different from what is observed in Pab, Sxl, HuD and hnRNP A1, where folding of unstructured linkers between neighbouring RRMs to form α-helices, rather than unfolding, is observed upon RNA binding. In these proteins, the α-helical linker regions participate in base-specific contacts and therefore contribute to specific binding both directly and by controlling the topological arrangement of the two RRMs (Sachs and Varani, 2000). In CstF-64, helix C is stably associated with the domain but unfolds upon RNA binding and therefore does not contribute directly to RNA recognition.

**CstF-64–DSE interaction and alternative polyadenylation**
A substantial number of RNA transcripts in vertebrates (30%) present alternative polyadenylation sites (Beaudouin et al., 2000). In the IgM-G heavy chain, perhaps the best understood example of regulated alternative polyadenylation (reviewed in Zhao et al., 1999), and in several other well characterized examples, poly(A) site usage is correlated with intracellular levels of CstF-64. The ‘strength’ of a poly(A) site appears to be determined at least in part by the affinity of the CstF-64–DSE interaction. Our data show that sequences containing at least two consecutive Us form more stable complexes with longer lifetimes and presented a model of how recognition of GU dinucleotides occurs. Analysis of sequences selected to bind CstF-64 confirms that sequences containing multiple successive Us bind more tightly to both CstF and to isolated CstF-64, and tend to be processed more efficiently (Beyer et al., 1997; Takagaki and Manley, 1997; Chao et al., 1999).
In several well studied examples of regulated polyadenylation, weak sites are utilized only at high cellular levels of CstF-64 (Takagaki and Manley, 1998; Chuvilin et al., 1999). Normally, the majority of CstF-64 is associated into the heterotrimeric CstF complex and recruited to the CTD during transcription. When cellular levels of CstF-64 are increased, free CstF-64 (or free CstF, i.e. the fraction not associated with the elongating polymerase) may saturate even weak GU-rich sequences found in upstream polyadenylation sites, perhaps increasing the chance of the formation of a complex committed to cleavage and transcription termination.

**Kinetic competition with transcription may affect polyadenylation site definition**

The affinity of the CstF–DSE interaction may control not just the efficiency of poly(A) site usage but also the kinetics of cleavage complex assembly. Surprisingly, 3'-end processing requires 10 s or more (Chao et al., 1999) and is faster for ‘stronger’ than for ‘weaker’ poly(A) signals. According to this result, selection of the weak sites may be inefficient because of the slow assembly rate of the cleavage complex compared with transcriptional elongation rates (Chao et al., 1999). We notice that the organization of transcripts undergoing alternative polyadenylation follows a common pattern: proximal polyadenylation sites are utilized with low efficiency while more distal sites are utilized by default. If assembly is fast enough, the transcriptional machinery might be directed to termination, while assembly at inefficient sites may occur so slowly that transcription has progressed far enough to allow skipping of the polyadenylation site. According to this ‘kinetic model’, weak sites are skipped because the reorganization of the 3'-end processing machinery into a complex ‘committed’ to cleavage is slow compared with transcriptional elongation; in other words, weak sites are kinetically silenced. We suggested that the unfolding transition in helix C might be an early step in the assembly of the cleavage complex, introducing a further kinetic element to regulation. Perhaps the structural change in CstF-64 observed with the high-affinity sequences might speed up the assembly of a functional cleavage complex. Other mechanisms may influence processing efficiency: for example, the presence of transcription pause sites, competition between CstF-64 and other factors binding the pre-mRNA (e.g. hKRNPs) and competition between splicing and polyadenylation. However, kinetic competition between RNA processing and transcription is an attractive model that may be relevant not just to polyadenylation but also to other situations, such as pre-mRNA splicing, where general processing factors necessary for basal processing reactions also play distinct regulatory roles on different sets of mRNAs.

**Materials and methods**

**Protein expression and purification**

Full-length human CstF-64 was amplified from a vector kindly provided by Professor J. Manley (Columbia University) and subcloned into the pET-28a vector (Novagen) between the NdeI and NotI restriction sites. The construct corresponding to the N-terminal RPM (amino acids 1–111) was generated by introducing a stop codon in the pET-28a–CstF64 full-length construct using site-directed mutagenesis. Cell cultures [BL21(DE3) *Escherichia coli*] were grown in M9 minimal media supplemented with appropriately isothe-labelled NH4Cl and glucose. Cultures were induced with isopropyl-β-D-thiogalactopyranoside (IPTG) during mid-logarithmic phase and harvested 4 h post-induction. Protein purification was by metal chelate affinity chromatography (Amersham-Pharmacia Hi-trap), followed by thrombin (Sigma) cleavage to remove the His tag. The sample was reloaded onto the Ni2⁺ affinity column following thrombin digestion and the flowthrough was passed through a benzamidine minicolumn (Amersham-Pharmacia Hi-trap) for thrombin removal. The protein was then further purified by anion exchange (Q-Sepharose) and size exclusion chromatography (Amersham-Pharmacia Superdex™ 75). No impurities were detected by SDS gel electrophoresis or mass spectrometry.

**RNA synthesis**

All RNA sequences studied (Table II) were prepared by in vitro run-off transcription using T7 RNA polymerase and subsequently purified essentially as described (Price et al., 1998), with the exception of sequence number 3, which was chemically synthesized (Dharmacon). After de-protection following the manufacturer’s instructions, the RNA was purified by gel electrophoresis as for the RNAs prepared in vitro.

**NMR spectroscopy**

NMR experiments were conducted on Bruker AX500, DMX600 and AV800 MHz spectrometers equipped with triple resonance probes and gradient units. All the experiments were performed under the same buffer conditions (10 mM phosphate pH 6.0) and at a temperature of 300 K. Backbone Cα, Cβ, C and N assignments of CstF-64 (1–111) free and in complex with the RNA (GU),GUUG sequence were obtained using triple resonance HNCA, HNCO, HN(CO)CA and CBCA(CO)NH experiments (Sattler et al., 1999). Side-chain assignments were completed with an HCH-NOESY experiment. Titration experiments were performed by adding increasing amounts of unlabelled oligonucleotides to 15N-labelled protein and by following the changes in [15N]HSQC protein spectra. Two-dimensional NOESY experiments (in H2O and D2O) and heteronuclear edited three-dimensional (3D) [15N- and 13C-]HSQC-NOESYs recorded at 600 MHz at mixing times of 80 ms were used to obtain distance constraints for free CstF-64 (1–111) protein. Two-dimensional NOESY spectra were recorded at 800 MHz for the RNA complex.

**Structure calculation**

The 3D structure of CstF-64 (1–111) protein was calculated using the program DYANA (Gürtler et al., 1997). A total of 1486 relevant NOE-derived distance restraints (upper limits only), corresponding to an average of 14 constraints per residue, were used in the calculation. A total of 1486 relevant NOE-derived distance restraints (upper limits only), corresponding to an average of 14 constraints per residue, were used in the calculation. The analysis of the Cα, Cβ and Cγ chemical shifts with the program TALOS (Cornilescu et al., 1999) provided 123 φ and ψ angle restraints that were used in the later stages of the structural refinement. Calculations were performed starting from 40 randomly generated conformers, which were then subjected to a simulated annealing protocol of 5000 torsion angle dynamics (TAD) cycles. Hydrogen atoms were included later, followed by a second low-temperature simulated annealing step of 800 TAD cycles. The protocol showed very good convergence; typically, only two or three of 40 starting structures failed to converge and showed large values of the target function representing deviations from experimental constraints. The 20 structures with the lowest target function were selected to represent the ensemble of converged CstF-64 (1–111) structures. Experimental data and structural statistics are summarized in Table 1. The quality of the structure was analysed with the suite of programs PROCHECK-NMR (Laskowski et al., 1996): 83.1% of residues have backbone angles within the most populated regions of the Ramachandran plot, 14.8% are in additional allowed regions, 1.9% are in generously allowed regions and only 0.2% are in disallowed regions.

**CstF–UU complex**

The ≥30% identity between CstF-64 and HuD was used in combination with the experimental data to model the CstF–UU interaction based on the HuD–RNA complex structure (Wang and Tanaka-Hall, 2001). Modelling was initiated from the constraint set obtained for RNA-free CstF-64, since the data recorded on protein–RNA complexes demonstrated conclusively that there are no structural changes except at the C-terminal helix. Since the NMR data clearly show that the helix unfolds upon RNA binding, all distances involving the C-terminal helix were removed from the distance constraint set. A set of intermolecular short distances (all 153 restraints corresponding to distances ≤6 Å) involving protons from U3 and U4 in the HuD complex was generated from the HuD coordinate set. These non-experimental constraints were arbitrarily given more generous upper bounds (30% larger) to account for possible structural differences
between HuD and Csf1R. We then recalculated the structure of Csf1R (1–111) bound to a DNA dimer using the same simulated annealing protocol used for the free Csf1R. No intramolecular distance restraints were used to define the DNA conformation.

Accession codes
Coordinates, experimental restraints and other NMR parameters have been deposited under PDB code 1PIT and BioMagRes accession number 5774.

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