Analysis of *in situ* pre-mRNA targets of human splicing factor SF1 reveals a function in alternative splicing

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ABSTRACT

The conserved pre-mRNA splicing factor SF1 is implicated in 3’ splice site recognition by binding directly to the intron branch site. However, because SF1 is not essential for constitutive splicing, its role in pre-mRNA processing has remained mysterious. Here, we used crosslinking and immunoprecipitation (CLIP) to analyze short RNAs directly bound by human SF1 *in vivo*. SF1 bound mainly pre-mRNAs, with 77% of target sites in introns. Binding to target RNAs *in vitro* was dependent on the newly defined SF1 binding motif ACUNAC, strongly resembling human branch sites. Surprisingly, the majority of SF1 binding sites did not map to the expected position near 3’ splice sites. Instead, target sites were distributed throughout introns, and a smaller but significant fraction occurred in exons within coding and untranslated regions. These data suggest a more complex role for SF1 in splicing regulation. Indeed, SF1 silencing affected alternative splicing of endogenous transcripts, establishing a previously unexpected role for SF1 and branch site-like sequences in splice site selection.

INTRODUCTION

Pre-mRNA splicing is critical for the expression of genetic information in most eukaryotes, and alternative splicing events largely contribute to proteome diversity in metazoans (1,2). On average, human pre-mRNAs contain eight exons separated by introns that vary in length between <100 and >100,000 nt (3), but signals that mark 5’ and 3’ splice sites are short and degenerate (4). Precise juxtaposition of cognate exons for intron removal is accomplished by dynamic interactions between the pre-mRNA, five small nuclear ribonucleoprotein particles (snRNPs) and more than 100 non-snRNP proteins (1). With few exceptions, splice sites are defined at the onset of spliceosome assembly. At this time U1 snRNP binds the 5’ splice site and the 3’ splice site is recognized by three proteins: splicing factor 1 (SF1, or mammalian branch point binding protein, mBBP) and the two subunits of the U2 snRNP auxiliary factor, U2AF65 and U2AF35. SF1 specifically binds the intron branch point sequence (BPS; 5,6), which is degenerate in mammals (YNCURAY; N = any nt, R = A or G, Y = C or U) but almost invariant in yeast (UACUAAC; 4). The underlined adenosine acts as the nucleophile in the first catalytic step of splicing (1). U2AF65 interacts with the polypyrimidine (Py) tract, located downstream of the BPS (7). U2AF35 recognizes the conserved AG dinucleotide that marks the intron 3’-end (8). SF1 and U2AF65 interact *in vitro* and *in vivo* and cooperatively bind the pre-mRNA (9–12). Recruitment of the U2 snRNP, which involves base pairing of the U2 snRNA with the BPS and binding of U2 snRNP proteins at and adjacent to the BPS, displaces SF1 from the spliceosome (13).

A hnRNP K homology/Quaking 2 (KH/QUA2) domain in the N-terminal half of SF1 (Figure 3A) contacts the bases of the BPS and buries the BPS-adenosine in a hydrophobic pocket of the KH-fold, which is thought to facilitate the formation of the BPS-U2
snRNA helix (6). U2AF65 binds to the Py tract through two central RNA recognition motifs (RRMs; Figure 3A; 14,15) and an arginine–serine-rich N-terminal region contacts the BPS in a sequence-independent manner (16,17). A third, non-canonical RRM of U2AF65 (or U2AF homology motif, UHM) interacts with the N terminus of SF1 (9–11). UHMs are also found in other proteins, which engage in networks with ligand proteins and coordinate constitutive and alternative splicing (18–20).

These results implied an important role for SF1 in early spliceosome assembly, emphasized by a requirement of SF1 for embryonic development in mice and Caenorhabditis elegans and viability in human cells and yeast (10,21–24). However, depletion of SF1 from yeast or human splicing extracts slowed the kinetics of early splicing complex formation without compromising splicing outcome, suggesting a kinetic role for SF1 in splicing (13,25). Moreover, splicing defects were not apparent after SF1 silencing (24), suggesting SF1 is only required for the splicing of a subset of pre-mRNAs in human cells, as reported for yeast SF1 (26), or plays another essential role in mammalian cells. SF1 has been implicated in changes in alternative splicing mediated by the β-catenin/TCF4 complex involved in colorectal carcinogenesis, but it is not clear whether this function is direct (27). In addition, an increased susceptibility of Sf1(+/–) mice to colon cancer may relate to a function in alternative splicing (23). Finally, a mutation in SF1 in fission yeast leads to exon skipping (28). Other findings implicated in changes in alternative splicing mediated by the β-catenin/TCF4 complex involved in colorectal carcinogenesis, but it is not clear whether this function is direct (27). In addition, an increased susceptibility of Sf1(+/–) mice to colon cancer may relate to a function in alternative splicing (23). Finally, a mutation in SF1 in fission yeast leads to exon skipping (28). Other findings suggested roles for SF1 in nuclear pre-mRNA retention in yeast (16,29) and as a repressor of transcription activation and elongation in human cells (30,31).

To clarify the role of SF1 in splicing or other aspects of RNA biogenesis we exploited its RNA-binding activity to isolate cognate RNA targets from HeLa cells. We used the crosslinking and immunoprecipitation (CLIP) method, which combines UV crosslinking in live cells with immunoprecipitation of short RNA fragments bound to a protein of interest (32,33). Consistent with a function for SF1 in mRNA maturation, the majority of SF1 target proteins, which engage in networks with ligand proteins and coordinate constitutive and alternative splicing (18–20).

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**MATERIALS AND METHODS**

**Crosslinking and immunoprecipitation**

CLIP was performed as described by Ule et al. (32) with the following modifications. HeLa cells were grown in Dulbecco’s Modified Eagle Medium (DMEM; Invitrogen) containing 10% fetal calf serum (Sigma-Aldrich). Cells (4 × 10⁷) were rinsed in PBS and irradiated at 400 mJ/cm² in a Model 2400 Stratalinker UV Crosslinker (Stratagene) in 10 ml ice-cold Hank’s balanced salt solution. Digestion with RNase T1 (Roche) was performed at a final concentration of 15 U/ml. SF1 and crosslinked RNA were precipitated by incubation of the cell lysate for 1 h at 4°C with mAb24D1 (directed against a region common to all SF1 isoforms; Z. Rafi and A.K., unpublished data) bound to Dynabeads Protein A (Dynal Biotech). Linker RL5 (Supplementary Table S5) was ligated to RNA and adducts were purified from a denaturing 10% polyacrylamide gel followed by ligation of linker RL3. RT–PCR was performed with primers DP5 and DP3. DNA primers for the concatamerization step were DP5EcoR1, DP3EcoR1 and DP3. Concatamers were cloned into pCR2.1-TOPO (Invitrogen) and sequenced by Sanger sequencing.

**Tag annotation**

CLIP tag sequences were mapped to the human genome assembly version that was available for BLAST-like alignment tool (BLAT) searches on the UCSC genome browser (http://genome.ucsc.edu/cgi-bin/hgBlat), hg18, and to the ENSEMBL genome and transcript databases (34).

**Determination of a weight matrix describing the sequence preferences of SF1**

To determine the sequence preference of SF1, we used a formalism that has been initially developed in statistical mechanics. Briefly, it can be shown that if a system can take on a certain set of states, each of them associated with an energy, then at equilibrium, each of the states will be visited in proportion to e⁻βE, where E is the energy of a state, and β is the inverse temperature. In our case, the states correspond to all the ways one or more proteins can be bound to the RNA. In the simplest example there is only a single type of protein and a short RNA fragment that can be bound in one way only. Let S be the concentration of the RNA, P the concentration of the protein and C the concentration of the complex. RNA–protein binding can be described by S + P ⇌ C with kₜ and kᵣ, being the forward and reverse rate constants. Let B and U be the measured intensities of bound and unbound RNA, with a scaling factor βᵣ relating the signal intensity to the RNA concentration and Pᵣ be the total protein. Finally, let us assume that we can write the dissociation rate of the protein from an RNA as K = (kᵣ/kₜ) = k₀ exp(−E(s)), where E(s) is the sequence-dependent energy of binding between the RNA and the protein. After rewriting the rate equations in terms of these quantities, we derive the following relationship:

\[ I_B = \exp[E(s)]/k_0[P_T/B - \beta_0], \]

which relates the intensity of unbound RNA to the total protein concentration and the intensity of bound RNA. With the assumption that the energy of interaction between the RNA and protein is composed of the individual energy contributions of individual positions in the binding site, and that these contributions depend only on the nucleotide that is present at that position in the binding site, the proportionality factor is \( \exp[E(s)]/k_0 \) depends only on the sequence of the RNA. That is, assuming a weight matrix model, \( E(s) \) can be written as \( E(s) = \sum_{i=1}^{L} E_i(s) \), where L is the length of the binding site (and the number of columns in the...
weight matrix), and \( s_i \) is the nt at position \( i \) of the binding site. The probability to observe nt \( s_i \) at position \( i \) of the binding site is proportional to \( e^{-E(s_i)} \) as mentioned above. Because we have three measurements for each individual RNA sequence, corresponding to three different total protein concentrations, we can fit a straight line to the protein concentrations, we can fit a straight line to the RNA sequence, corresponding to three different total nucleotides at that position. That is, the entry \( w_{iB} ^g \) in the weight matrix, which represents the expected frequency of nucleotide \( B \) at position \( i \) in a binding site of the protein is given by

\[
\exp[E(B)] \sum_{B \in \{A,C,G,U\}} \exp[E(B')] = \frac{\exp \sum_{j \neq i} E(s_j) + E(B)}{\sum_{B \in \{A,C,G,U\}} \exp \sum_{j \neq i} E(s_j) + E(B')},
\]

which can be derived from the slopes fitted above.

**Determination of sequence motifs overrepresented in SF1 CLIP tags**

We compared the frequency of tetrameric motifs in isolated SF1 CLIP tags to randomized sequences with the same nucleotide composition, which were obtained by shuffling the tags. Motifs, whose frequency in the real and the randomized sets were significantly different, were identified as follows. Given the frequencies of the motifs and the total number of tetramers in the real and randomized sets, we computed the posterior probability for the model that assumes that the frequency of a motif is different as opposed to being the same between the two datasets, as described (35). Motifs that were found enriched in introns and exons are shown in Supplementary Table S3. We also compared the frequency of tetrameric motifs in exonic/intragenic tags with those found genome-wide in exons or introns as follows. We downloaded mRNAs sequences to Genbank (www.ncbi.nlm.nih.gov) and, based on the Entrez Gene database, we chose the longest mRNA corresponding to each annotated gene in the genome. We mapped the mRNAs to the hg18 release of the human genome assembly, which we obtained from the University of California at Santa Cruz (genome.ucsc.edu) with the GMAP alignment program (36). This mapping defined exonic and intronic genomic regions. We then sampled for each data set (exonic or intronic tags) 100 randomized data sets. Each of these datasets was generated by choosing with uniform probability the same number and length of regions as present in the CLIP sets from the appropriate type of region in the genome. For each of the 100 randomized sets, we computed the frequency of all possible tetramers. Supplementary Table S4 shows the frequency of the tetramers in the CLIP sets and the z-value of this frequency with respect to the frequencies computed in the randomized datasets.

**In vitro transcription and RNA purification**

RNAs were synthesized in the presence of \( [\alpha^{32}P]UTP \) with the T7-MEGAscript kit (Ambion) and gel-purified. Additional G’s were added to the 5’-end of the RNAs to optimize transcription efficiency. The BPS RNAs (GGGGAGUAUACUAACU with the T7-MEGAscript kit (Ambion) and gel-purified. Additional G’s were added to the 5’-end of the RNAs to optimize transcription efficiency. The BPS RNAs (GGGGAGUAUACUAACU) (5) were purifed on Ni-NTA columns (Qiagen) under native conditions. GST-U2AF65Δ1-94 (16) was purified on glutathione-agarose (Sigma). Purified proteins were dialyzed against 20 mM HEPES-KOH pH 7.9, 100 mM KCl, 20% glycerol, 0.2 mM EDTA and 0.5 mM dithiothreitol.

**Electrophoretic mobility shift assays**

SF1-KH/QUA2 domain was expressed and purified as described (6). His6-SF1-C4 and His6-SF1-C4/W22A (10,11) were purified on Ni-NTA columns (Qiagen). GST-U2AF65Δ1-94 (16) was purified on glutathione-agarose (Sigma). Reaction products were resolved in native 8% polyacrylamide gels (acrylamide:bisacrylamide = 80:1) in 1×TBE at 4°C for 3 h at 100 V.

Cooperative binding was performed for 15 min at room temperature in 10-µl reactions containing GST-U2AF65Δ1-94 and His6-SF1-C4 or His6-SF1-C4/ W22A, 50 pmol [\( \alpha^{32}P \)UTP-RNA, 80 mM KCl, 16 mM NaCl, 4 mM imidazole, 2 mM β-mercaptoethanol, 5 μg tRNA and 7 U rRNasin (Promega). Reaction products were resolved in native 8% polyacrylamide gels (acrylamide:bisacrylamide = 80:1) in 1×TBE at 4°C for 3 h at 100 V.

 Autoradiographs were quantified with the Molecular Imager FX (BioRad) and software Quantity One V 4.2.1 (BioRad).

**RNA interference and semiquantitative RT-PCR of pre-mRNA targets of SF1**

HeLa cells were grown in DMEM/high glucose containing 10% fetal calf serum (Sigma-Aldrich), Cells \( 2 \times 10^5 \) were transfected in the presence of 73 nM SF1 or SF3a120 siRNAs (Dharmacon; Supplementary Table S5) and oligofectamine (Invitrogen) according to Dharmacon’s instructions. Control transfections were performed in the presence of siRNA LUC, targeting firefly luciferase (LUC; 37) or in the absence of siRNAs. Duplicate samples were lysed 48 h post-transfection for western blot analysis (24) and isolation of cytoplasmic RNA with the RNeasy Mini kit (Qiagen). RNA was treated with 2 U RQ-DNase (Promega) for 45 min at 37°C and
phenol–chloroform extracted. RT was done for 2 h at 40°C in the presence of 4μg cytoplasmic RNA, 4μg oligo dt(12–18) (Sigma-Aldrich), 40 U rRNasin and 200 U MMLV-RT (Invitrogen). PCR was performed with the Expand High-Fidelity PCR system (Roche) for 28 cycles. Control PCR reactions of 32 cycles indicated amplification in the linear range (not shown). Primers for amplification of SF1, SF3a120, H3F1 mRNAs have been described elsewhere (24). Other primers are listed in Supplementary Table S5. Forward primers were 5′-end-labeled with [γ-32P]-ATP and T4 polynucleotide kinase (Invitrogen). PCR products were separated in native 4% polyacrylamide gels, visualized by autoradiography and quantified by phosphoimaging as above. Alternatively, PCR products were separated in agarose gels and quantified with ImageJ 1.410 (National Institutes of Health). PCR products were gel-purified and sequenced.

RESULTS

Isolation of in situ RNA targets of SF1 from HeLa cells

We isolated SF1–RNA complexes from a whole-cell lysate of UV-irradiated HeLa cells with antibody mAb24D1 (see ‘Materials and Methods’ section). SF1 was efficiently removed from the lysate after immunoprecipitation in the presence, but not absence of mAb24D1 (Figure 1A). Non-specific binding of RNA to beads in the absence of antibody was not observed (I. Bagdiul and A. Kra¨mer, data not shown). Following 5′-end-labeling of RNA, SF1–RNA complexes were resolved by PAGE and transferred to nitrocellulose. Autoradiography revealed a series of bands between 50 and 220 kDa superimposed on a smear of radioactivity (Figure 1B). SF1 isoforms detected by mAb24D1 in HeLa cells migrate between 70 and 85 kDa; SF1 crosslinked to RNA fragments of ≥30 nts are expected to migrate at least 10 kDa higher. Denaturing PAGE of RNA purified from consecutive slices of a membrane comparable to that shown in Figure 1B revealed an increase in RNA length with increasing size of the RNA–protein complexes (Figure 1C), suggesting that RNAs of increasing size are crosslinked to one protein rather than RNAs of similar size distribution being crosslinked to proteins of different molecular mass. As RNAs (CLIP tags) of 50–200 nt are best suited for analysis (32), we further processed SF1–RNA complexes of 80–100 kDa, followed by cloning and conventional sequencing.

Distribution of RNA targets in the human genome

We obtained ~360 SF1 CLIP tags and compared their sequences to the human genome by BLAT searches. Sequences with <90% match to the genome, multiple hits and exact duplicates were removed, resulting in a final set of 227 sequences (Supplementary Table S1). The majority of these (193) are located in protein-coding genes and 14 in genes of non-coding RNAs (Figure 1D). A total of 20 CLIP tags are present in intergenic regions, 7 of which map to the opposite strand of known genes. Intergenic CLIP tags were also found in other studies and may correspond to yet unannotated transcripts (38–41).

SF1 CLIP tags in protein-coding genes are largely confined to introns (Figure 1E and Supplementary Table S1). Despite SF1’s function in BPS binding (5,6), only 19 intronic tags are present at or close to 3′ splice sites, two map to 5′ splice sites and the remainder are located >40 nt from intron ends. Four intronic CLIP tags map to snoRNAs. One of these spans the intron–snoRNA border and is thus derived from the intron, which may also be true in the other cases.

Among 40 exonic CLIP tags, 21 are located in coding sequences, two in 5′-untranslated regions (UTRs) and 17 in 3′-UTRs (Figure 1E). Moreover, of the 43 tags located within exonic regions of known genes, 22 originated in 3′ terminal exons (Supplementary Table S1). Since human genes have on average 8–9 exons (3), this distribution indicates an enrichment of 3′ terminal exons among the exonic tags (P-value 9.9 x 10^{-10} in a binomial test assuming a probability to observe a 3′ terminal exon of 1/8). Additionally, in repeated sampling of 43 human exons from all human genes (dataset described in
‘Materials and Methods’ section), we only obtain an average of four terminal exons, indicating that targets in such exons are indeed overrepresented among SF1 CLIP tags. In addition, three CLIP tags span exon–exon junctions, suggesting that SF1 can bind mature mRNA. Finally, 42% of the CLIP tags in protein-coding genes are located in regions that are subject to alternative splicing (Supplementary Table S1).

Thus, SF1 mainly targets pre-mRNAs and binding sites are associated with constitutive and alternatively spliced regions of transcripts. An unexpectedly high percentage of SF1 target sites are located in 3'-UTRs and SF1 appears to bind mature mRNA as well. Compared with other studies that combined CLIP with high-throughput sequencing (38–41), our dataset only provides a glimpse at possible in situ SF1 binding sites and it is not surprising that, with a few exceptions containing two or more CLIP tags, only one SF1 CLIP tag was found per gene.

RNA binding preferences of SF1

SF1 specifically recognizes the BPS (5,6). Yeast SF1 selected the consensus BPS from a pool of randomized sequences (42), but information on binding preferences of human SF1 is limited. Mutational analyses and the NMR structure of the KH/QUA2 RNA-binding domain of human SF1 in complex with a UACUAAC-containing RNA showed that the URA in the 3’ half of the BPS is specifically recognized, whereas interactions with the 5’ half are less important (5,6,43). About 80% of the SF1 CLIP tags contained at least one copy of the URA motif; however, this sequence was not specifically enriched in our dataset compared to randomized sequences with the same overall length and nucleotide composition.

To precisely define binding preferences of human SF1, we tested 25-nt RNAs containing the consensus BPS or mutations at all BPS positions in electrophoretic mobility shift assays (EMSAs) for binding to recombinant SF1-KH/QUA2 protein (Figure 2A). The results were used to infer a weight matrix describing the sequence specificity of SF1 (Figure 2B and Supplementary Table S2). The derived motif resembles the human BPS consensus, except that SF1 strongly prefers an A at position –4 for optimal binding, whereas no nt bias is found at this position in the BPS consensus (4). The motif also comes close to the yeast BPS but lacks the preference for U at position –5. The information score at individual positions in the weight matrix is generally less than one (Supplementary Table S2), suggesting that human SF1 can bind a large variety of sequences. This is also reflected in only 4.4% of the CLIP tags containing the sequence ACUNAC (which we will refer to as the SF1 binding motif).

In contrast, Py-rich motifs were the most significantly enriched sequences in the CLIP tags relative to unrelated sequences with the same mononucleotide frequency. Approximately half of both intronic and exonic tags contained at least one occurrence of the motifs CCUG or UCCU (Supplementary Tables S3 and S4). CCUG repeats, which form mismatched stems, are known binding sites for MBNL proteins; the repeats sequester

Figure 2. Determination of the optimal SF1–RNA-binding motif. (A) In vitro-transcribed RNAs containing the wild-type or mutant BPS were incubated with buffer or SF1-KH/QUA2 (5, 10 and 20 μM; indicated by triangles above the figure). Reaction products were separated by native PAGE and visualized by autoradiography. The same wild-type images were used for positions –5/–4/–3, –2/+1 and BP/–1. (B) Web logo representing the weight matrix for each position of the SF1 binding site derived from the quantification of the data presented in A.
Cooperative interaction of SF1 and U2AF65 with RNA targets

Association of SF1 with branch sites likely reflects SF1 and U2AF binding to adjacent RNA sequences; in principle, SF1 and U2AF interactions with RNA could be competitive, neutral or cooperative. To address this, we performed EMSA with CLIP tag-derived RNAs and recombinant U2AF65A1-94 and SF1-C4 (Figure 3A), comprising the domains required for binding each other and RNA (11,16). U2AF65 and SF1-C4 alone bound RNA 2-50 (Figure 3D, lanes 2–6 and 7), containing an 11-nt Py tract following the 5’ most SF1 binding motif (Figure 3B). A ternary complex formed in the presence of both proteins, while the SF1-C4/RNA complex decreased (lanes 8–12). Consistent with cooperative RNA binding of SF1 and U2AF65, the ternary complex formed at a lower U2AF65 concentration when SF1 was present (compare lanes 2 and 3 with lanes 8 and 9).
UPF3 regulator of nonsense transcript homolog A (UPF3A) pre-mRNA. The mRNA containing exons 1–5 was strongly reduced compared to the controls, whereas levels of mRNAs lacking exon 4 or exon 4 plus exon 2 or 3 (both of which are 107 nt in length) increased (Figure 5E). When we analyzed RT–PCR products from an independent experiment with SYBR Green, the 318 bp product from control cells was resolved into two closely migrating bands (Supplementary Figure S2B). PCR with primers in exons 2 or 3 and 5, as well as sequencing revealed that the slower migrating band corresponded to exons 1, 3 and 5, whereas the faster migrating band...

Figure 3. Cooperative binding of SF1 and U2AF65 to an endogenous SF1 target. (A) Scheme of SF1 and U2AF65 constructs used for EMSA. The U2AF65 interaction domain (U2AF65-ID), KH/QUA2 domain and the zinc knuckle (Zn) of SF1 are shown, as well as the arginine-serine-rich (RS) domain, RRM1 and 2 and the UHM of U2AF65. The star above the U2AF65-ID of SF1-C4 indicates the location of the W22A mutation. Numbers indicate amino acids of the truncated proteins. Variability in the length of SF1 isoforms is indicated by dashed lines. (B) RNAs corresponding to CLIP tag 2-50 (wild-type, WT, or mutants M1–M6) were transcribed in vitro. SF1 binding motifs are indicated in bold. Mutations are underlined. (C) Wild-type and mutant RNAs were incubated with buffer or SF1-KH/QUA2 (5, 10 and 20 μM; indicated by triangles). Reaction products were separated by native PAGE and visualized by autoradiography. (D) RNA 2–50 WT was incubated with buffer or U2AF65Δ1–94 (0.2, 0.5, 1, 2 and 4 μM; indicated by triangles) in the absence or presence of 6.6 μM SF1-C4 as indicated. Reaction products were separated by native PAGE and visualized by autoradiography. The migration of RNA 2–50 bound to U2AF65Δ1–94 (open arrowheads), SF1-C4 (vertical bar) or both (closed arrowheads) is indicated.
contained exons 1, 2 and 5 (data not shown). SF1 silencing reduced the exon 1-2-5 mRNA to background levels, whereas the exon 1-3-5 mRNA was strongly increased. Together, these results show that SF1 can act as a negative or positive regulator of exon inclusion.

Of note, although depletion of SF3a120 resulted in a general reduction of mRNA levels (as apparent by inspection of the autoradiographs), in some cases the ratio of spliced products was also changed compared to controls (Supplementary Figure 5B–E). This result is not unexpected, since SF3a120 has been shown to be involved in alternative splicing in Drosophila (47).

PDE3A pre-mRNA was identified as a potential SF1 target by three CLIP tags (Figure 3 and Supplementary Table S1). SF1 and SF3a120 knockdown reduced the single endogenous PDE3A mRNA detected in HeLa cells to 51–53 and 19%, respectively, compared to the controls (Supplementary Figure S3A), suggesting that SF1 affects the maturation or stability of PDE3A mRNA. Similarly, the level of a single mRNA of the asparagine-linked glycosylation 8 homolog (ALG8) decreased to 50–54 and 39% after SF1 and SF3a120 depletion, respectively (Supplementary Figure S3B). Thus, SF1 knockdown negatively affects the expression of these pre-mRNAs, although to a lesser extent than depletion of SF3a120.

These data demonstrate that SF1 indeed functions in splicing. However, it does not appear to be a constitutive splicing factor. SF1 may be required for the splicing of all introns of PDE3A and ALG8 pre-mRNAs. In contrast, in other cases tested, SF1 downregulation changed the ratio of mRNA isoforms without decreasing total mRNA levels, as observed after depletion of SF3a120. We conclude that SF1 plays a role in alternative splicing events.

DISCUSSION

We have isolated RNAs bound to SF1 in HeLa cells to clarify SF1's role in splicing and/or other steps of RNA biogenesis. As expected for a protein that functions in mRNA maturation, the majority of SF1 targets mapped to protein-coding genes. We determined the RNA binding preferences of SF1, validated selected CLIP tags as bona fide SF1 ligands and demonstrated a role for SF1 in the splicing of several pre-mRNAs identified as potential targets. Our data indicate that SF1 is not required for the splicing of all introns, but influences alternative splicing decisions.

A role for SF1 in alternative splicing

Silencing of SF1 changed the ratio of exon inclusion/skipping in alternatively spliced products of several endogenous pre-mRNAs identified as potential SF1 targets (Figure 5 and Supplementary Figure S2). Depending on the event analyzed, SF1 activated or repressed exon inclusion, which is not without precedent. Nova, Fox and several hnRNP proteins regulate alternative splicing in both directions (2). Moreover, CLIP combined with high-throughput sequencing (HTS) provided a genome-wide view of splicing factor binding sites and demonstrated that Nova, Fox2 and PTB regulate alternative splicing in both directions (2).}

Figure 4. Cooperative binding of SF1 and U2AF65 to an endogenous SF1 target with a suboptimal SF1 binding site. (A) RNAs corresponding to wild-type (WT) and mutant (M) CLIP tag 1–10 were transcribed in vitro. A SF1 binding motif is indicated in bold. The mutation is underlined. (B) Wild-type and mutant RNAs were assayed as in Figure 3C. (C) Wild-type and mutant RNAs were tested as in Figure 3D in the absence or presence of 6.6 μM SF1-C4 or SF1-C4/W22A as indicated. The migration of RNAs bound to U2AF65Δ1–94 (open arrowheads), SF1-C4 (vertical bar) or both (closed arrowheads) is indicated.
in splicing (49). Third, an increased susceptibility to colon cancer of Sf1 (+/-) mice could be caused by aberrant splicing due to reduced SF1 levels (23). Finally, a mutation in SF1 in fission yeast leads to exon skipping (28) and Saccharomyces cerevisiae SF1 is required only for the splicing of pre-mRNAs with suboptimal splice sites (26).

SF1 silencing also reduced the levels of single endogenous mRNAs (Supplementary Figure S3). Whether this is due to defects in splicing one or more introns is not clear. Lower mRNA levels rather than aberrantly spliced products may be due to faulty splicing upon SF1 depletion, which can introduce premature termination codons into mRNAs and cause mRNA degradation by nonsense-mediated decay (50).

SF1 is not an isolated case of a splicing factor thought to merely function in constitutive splicing but involved in alternative splice site choice. Other examples are U2AF65 and SF3a120 (47,51). It would be interesting to see whether these and possibly other proteins are required for all constitutive splicing events or, similar to SF1, also participate in the splicing of a subset of introns.

The SF1–BPS interaction is thought to facilitate base pairing of U2 snRNA with the BPS by pre-bulging the branch point adenosine (6). If SF1 is involved in the splicing of only certain introns, does another protein take over SF1’s role? The arginine–serine-rich domains of U2AF65 and SR proteins contact the phosphates of the BPS and could aid U2 snRNP in binding the BPS in the absence of SF1 (16,17,52). Moreover, the U2 snRNP-associated p14/SF3b14a, which specifically binds the branch point adenosine during A complex assembly (53,54), could facilitate base pairing.

SF1 binding preferences

An advantage of CLIP over other methods to identify targets of RNA-binding proteins is the possibility to locate approximate binding sites within targets and define consensus binding motifs (32,38–41,55). SF1 specifically recognizes the BPS (5,6,43). However, BPS-like sequences were not enriched in our dataset compared to random sequences with the same mononucleotide composition. Analysis of SF1 binding to a comprehensive set of BPS mutants defined ACUNAC as the consensus SF1 binding motif (Figure 2), which agrees in large part with previous limited mutational studies and the human

Figure 5. Effects of RNAi-mediated depletion of SF1 on the splicing of CLIP tag-containing pre-mRNAs. (A) cDNA from HeLa cells transfected in the absence (mock) or presence of siRNAs targeting SF1 (SF1 #1 and #2), luciferase (LUC) or SF3a120, as indicated on top of each panel, was PCR-amplified with primers specific for SF1, SF3a120 and H3F1 mRNAs, products were separated by PAGE and visualized by autoradiography. The numbers below the ‘mRNA’ panels indicate the percentage of mRNA normalized to the LUC control. Data represent the average of two experiments; the standard deviations were <1. HeLa cell lysates prepared in parallel were separated by SDS-PAGE and proteins revealed by western blotting. (B, C, D and E) The effect of SF1 depletion on the splicing of endogenous FGFR1OP (B), TNIP1 (C), PLOD2 (D) and UPF3A (E) pre-mRNAs was analyzed as in panel A. Sizes of DNA markers and spliced products are shown to the left and right of the images, respectively. Results of at least two experiments were quantified and are expressed as percent exon inclusion below the panels in addition to the standard deviation (std. dev.). The asterisk (panel B) indicates a PCR product that could not be identified by sequencing. The schemes depict pre-mRNA regions subject to alternative splicing (not to scale) and observed splicing patterns. Forward (F) and reverse (R) PCR primers including the exon number (E#) are shown above the schemes. Filled circles indicate the approximate location of CLIP tags. TNIP exon 2 is transcribed from an alternative promoter and not included in mRNAs transcribed from exon 1.
consensus BPS. The ACUNAC also resembles a BPS-like sequence identified downstream of exons specifically included in muscle (56). An exception is position −4, where SF1 favors an A, but no preference is evident in the consensus BPS (4), nor is the A specifically recognized in the SF1–BPS solution structure (6). In addition, a −4 A to G mutation did not affect SF1 function in previous studies (5,43). Another exception is position −1, where our data reveal the lowest information score for SF1 binding, although a −1 C can compromise SF1 binding (Figures 2 and 4, Supplementary Table S2). The SF1–BPS solution structure indicated coordination of a purine at this position (6); however, a −1 A to G mutation of an in vivo splicing reporter reduced SF1 function (43).

An information score of <1 at individual positions of the weight matrix (Supplementary Table S2) and <5% of the SF1 CLIP tags containing an exact match to the ACUNAC motif suggest a more relaxed RNA binding specificity of human SF1 compared to other splicing factors, which mirrors the variability of the mammalian BPS (4). This agrees with an RNA binding affinity of SF1, even for the consensus BPS, in the micro-molar range (5,6). In addition, the efficiency of SF1 binding to CLIP tag-derived RNAs did not always correlate with a perfect or suboptimal fit to the SF1 binding motif, suggesting that other parameters, such as neighboring sequences or RNA secondary structure, may influence the interaction. This may explain the variability in SF1 binding preferences observed in different experimental systems (see above).

The enrichment of Py-rich motifs in SF1 CLIP tags, many of which resemble U2AF65 binding sites (46; Supplementary Tables S1 and S3), is not unexpected, since SF1 binds RNA cooperatively with U2AF65 (9,43). Importantly, we not only observed cooperativity when SF1 alone bound RNA, but also when SF1–RNA binding was not evident (Figures 3D and 4C; data not shown). Thus, the interaction with U2AF65 increases the binding repertoire of SF1. Domains similar to the U2AF65 UHM, which contacts SF1, are present in other splicing factors engaging in networks with UHM ligands, including SF1 (19,20; G. Gregorovic and A. Krämer, unpublished data). We therefore speculate that not only U2AF65, but also other UHM proteins direct SF1 to appropriate binding sites in a pre-mRNA and regulate its function in the selection of specific splice sites.

Potential additional functions of SF1
Yeast SF1 dissociates from the spliceosome upon U2 snRNP recruitment (13) and human SF1 is most likely displaced from the BPS by binding of SF3b14a and base paring of the U2 snRNA (53,54,57), consistent with the absence of SF1 from late spliceosomes or spliced mRNPs containing the U2 snRNA (53,54,57), consistent with the ACUNAC motif. This agreement with a more relaxed RNA binding specificity of human SF1 compared to other splicing factors, which mirrors the variability of the mammalian BPS (4). The SF1–BPS interaction plays an important role in alternative splicing decisions. Furthermore, the identification of an unexpectedly high number of SF1 target sites in 3′ terminal exons hints to additional functions of SF1 in other steps of mRNA biogenesis.

SUPPLEMENTARY DATA
Supplementary Data are available at NAR Online.

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