Genetic analysis of the role of human U1 snRNA in mRNA splicing: I. Effect of mutations in the highly conserved stem-loop I of U1

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The 5' splice site mutation known as hr440 can be suppressed efficiently in vivo by a compensatory base change in U1 small nuclear RNA (snRNA). We have now begun a second-site reversion analysis of this suppressor U1-4u snRNA (which has a C → U change at position 4) to identify U1 nucleotides that are essential for mRNA splicing. Point mutations in U1-4u that disrupt the structure of stem-loop I or alter phylogenetically conserved nucleotides within the loop cause loss of suppression. The level of suppressor activity observed for most mutants correlated with the abundance of the corresponding suppressor RNA, suggesting that mutations in stem-loop I cause loss of suppression by destabilizing U1 snRNA or the U1 snRNP (small nuclear ribonucleoprotein particle). We favor the interpretation that incompletely or improperly assembled U1 snRNPs are unstable, because two severe point mutations in stem-loop I were found to decrease the binding of U1 snRNP-specific proteins in vitro. In a separate set of experiments, we found that increasing the distance between stem-loop I and the 5' end of U1 snRNA also inhibited suppression but did not affect assembly or stability of the U1 snRNP. This suggests that the relationship between the 5' splice site and the body of the U1 snRNP is important for mRNA splicing.

[Key Words: U1 snRNA; mRNA splicing; snRNP assembly]

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U1 small nuclear RNA (snRNA) is the most abundant of the U-snRNAs found in all eukaryotic cells (Reddy and Busch 1988). Except for U6, the U-snRNAs are transcribed by RNA polymerase II and possess a unique tri-methylguanosine cap structure (for review, see Dahlberg and Lund 1988). U-snRNAs exist in the nucleus as components of small nuclear ribonucleoprotein particles (snRNPs; for review, see Lührmann 1988). The protein components of the U1 snRNP can be divided into two classes: the common U-snRNP proteins (B', B, D, D', E, F, and G) and the U1 snRNP-specific proteins (70K, A, and C). Autoimmune sera with anti-Sm specificity recognize several of the common U-snRNP proteins, whereas autoimmune sera with anti-[U1]RNP specificity recognize proteins unique to the U1 snRNP.

The splicing of mRNA precursors requires the U1, U2, U4/U6, and U5 snRNPs (for review, see Steitz et al. 1988). The snRNPs assemble onto the mRNA precursor to form a large multicomponent ribonucleoprotein complex called a spliceosome (Brody and Abelson 1985; Frendewey and Keller 1985; Grabowski et al. 1985). Splicing is then accomplished by two consecutive phosphoester bond transfers: First, the phosphoester bond at the 5' splice site is attacked by the 2' hydroxyl group of the branchpoint residue, located 17–45 nucleotides upstream from the 3' splice site. This cleaves the 5' splice site and generates a lariat structure containing a 2'–5' phosphoester bond between the 5' end of the intron and the branchpoint. Second, the 3' hydroxyl of the 5' exon attacks the phosphoester bond at the 3' splice site. This ligates the two exons and releases the intron as a lariat, which later is debranched (for reviews, see Green 1986; Padgett et al. 1986).

One function of the snRNPs in mRNA splicing is to recognize essential sequences on the mRNA precursor. The U1 snRNP recognizes the 5' splice site (Mount et al. 1983; Black et al. 1985) in an interaction that depends, at least in part, on base-pairing between the 5' end of U1 snRNA and the 5' splice site (Zhuang and Weiner 1986; Seraphin et al. 1988; Siliciano and Guthrie 1988). The U5 snRNP is believed to bind to the 3' splice site (Chabot et al. 1985) by interacting with a 3' splice-site-binding protein (Gerke and Steitz 1986; Tazi et al. 1986). In yeast, the U2 snRNP recognizes the invariant branchpoint sequence UACUAAC by base-pairing (Parker et al. 1987), and current evidence suggests that the same is true in mammalian systems, although the branchpoint...
consensus PyUPuAC is much weaker (Reed and Maniatis 1988; Zhuang et al. 1989). In mammals, binding of the U2 snRNP to the branchpoint (Black et al. 1985) also requires an auxiliary factor called U2AF, which independently recognizes the 3' splice site (Ruskin et al. 1988). In addition, interactions between U-snRNPs may help to assemble the spliceosome and to bring distant 5' and 3' splice sites together. Various hnRNP [heterogeneous nuclear ribonucleoprotein particle] proteins also may contribute to assembly of the spliceosome and recognition of specific sequences or RNA structures within the mRNA precursor (Swanson and Dreyfuss 1988).

Using the adenovirus E1A gene, we have shown previously that a 5' splice site mutation known as hr440 can be suppressed efficiently in vivo by a compensatory base change in U1 snRNA (Zhuang and Weiner 1986). The suppressor U1 snRNA was designated U1-4u to indicate that position 4 in the U1 sequence was changed from C to U to compensate for the G → A change at position +5 in the 12S 5' splice site introduced by the hr440 mutation. The existence of a suppressor U1 snRNA provides a tool for a genetic analysis of the role of U1 snRNP in splicing. Second-site mutations can now be introduced into the suppressor U1 RNA to identify U1 nucleotides that are essential for function. This kind of second-site reversion analysis should enable us to identify U1 nucleotides that are involved in steps such as snRNP assembly, snRNP binding to the mRNA precursor, interaction between snRNPs within the spliceosome, and possibly the actual phosphoester bond transfers that accomplish splicing. In this initial paper we describe the characterization of point mutants in stem-loop I of the suppressor U1-4u snRNA.

Results

Construction and analysis of suppressor U1-4u second-site mutations

To begin our genetic analysis of U1 structure and function, we chose to make point mutations in stem-loop I because both the structure of this stem and the sequence of the loop have been highly conserved through evolution (Branlant et al. 1980; Mount and Steitz 1981; Forbes et al. 1984; Kato and Harada 1985; Yu et al. 1986; Kretzner et al. 1987; Siliciano et al. 1987; van Santen and Spritz 1987; Kiss et al. 1988; van Santen et al. 1988). Figure 1 shows the location of 22 point mutations that were introduced by site-directed mutagenesis into stem-loop I of the suppressor U1-4u.

To determine the suppressor activity of the U1-4u mutants, we performed a transient expression assay using the hr440 mutant of the adenovirus E1A mRNA precursor as substrate. HeLa cells were cotransfected with a mutant U1-4u gene and an E1A gene containing the hr440 mutation that abolishes 12S splicing (Zhuang and Weiner 1986). Suppression of the defect in 12S splicing then was measured by subjecting total cytoplasmic RNA from the transfected cells to RNase protection analysis. As shown in Figure 2, RNA from cells transfected with the hr440 gene alone and no suppressor U1-4u gene protected only two fragments derived from the 5' and 3' exons of E1A 13S RNA (Fig. 2A, lane 3; Fig. 2B, lanes 3 and 11; a residual level of 12S 5' exon is detected also). RNA from cells transfected with the hr440 gene, together with the suppressor U1-4u gene, gave a much stronger 12S 5' exon signal (Fig 2A, lane 4; Fig. 2B, lanes 4 and 12). RNAs from cells transfected with the hr440 gene and different U1-4u mutants protected variable amounts of the 12S 5' exon fragment. Levels of suppression obtained with different U1-4u mutants were quantified by densitometry. To do this, we measured the level of protected 12S 5' exon fragment relative to 3' exon fragment to normalize suppression to the total amount of stable 9S + 12S + 13S mRNA derived from the E1A transcription unit (see Zhuang and Weiner 1986). The suppressor activities of the U1-4u mutants are shown in Table 1 as a percentage of the suppression obtained with the parental U1-4u.

Disruption of stem I reduces suppression

Mutations that disrupt base-pairing in stem I generally result in poor suppression (Table 1), this is especially true for the four mutations that disrupt base-pairing in...
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Figure 2. Effect of second-site mutations in stem-loop I on suppression by U1-4u. HeLa cells were transfected with the adenovirus E1A hr440 mutant and different second-site mutants of U1-4u. Cytoplasmic RNA from transfected cells was hybridized to a labeled SP6 antisense transcript of the E1A hr440 gene, as shown in C. The resulting hybrids were then digested with T1 and pancreatic RNase and resolved on a 5% denaturing polyacrylamide gel. |A| Cells were transfected with hr440 and pUC13 vector (lane 3), with hr440 and the U1-4u gene (lane 4) or with hr440 and mutants in the stem of stem-loop I of U1-4u (lanes 5–15). |B| Mock-transfected cells without added DNA; |lanes M| denatured DNA size markers from an end-labeled HpaII digest of pBR322. |B| Cells were transfected with hr440 and mutants in the loop of stem-loop I of U1-4u (lanes 5–10,13–16). Other lanes are as in A.

Loop I mutations have a wide range of phenotypes

Mutations in loop I yield a range of phenotypes from normal to null [Fig. 2B]. As expected, mutation of the phylogenetically conserved C{sub 31} [see Table 2] almost completely abolishes suppression [mutant 31U], but mutation of the unconserved C{sub 33} has little effect; mutant 33A has wild-type suppressor activity, and 33G has ~86% activity. Mutations in other highly conserved loop nucleotides, however, have only intermediate effects [29C, 32C, 34U, 35U, 36U, and 38U]. Perhaps even more surprising is the observation that duplication of the highly conserved nucleotides 28–31 does not abolish suppression [mutant r28-31; 19% residual suppressor activity]. Mutation 38U, which might have been expected to disrupt a potential U2{sub 27} : G{sub 39} base pair at the

the middle of the upper stem [mutants 24A, 25A, 40U, and 41C]. The compensatory double-point mutation 25A/40U was made by combining mutations 25A and 40U, thereby replacing a C : G with an A : U base pair. Although the component point mutations 25A and 40U are both severely defective (~7–13% suppression), the compensatory double-point mutation 25A/40U restores suppressor activity to nearly normal levels (~78%). The importance of a stable stem I is supported further by the effect of mutation 20A, which presumably strengthens stem I by replacing the normal G : U base pair with a stronger A : U base pair and reproducibly gives higher levels of suppression than the parental U1-4u. Mutation 43A appears to disrupt base-pairing within the stem and yet has little effect on suppression. Superficially, this result would appear to contradict the idea that U1 function requires a stable stem I; however, it is also possible that A{sub 43} base-pairs with U{sub 22}, restoring the stable stem and bulging A{sub 21} instead of U{sub 22}. Consistent with the interpretation that the identity of the bulged nucleotide in stem I is not critical, mutation 22G replaces the bulged U residue with a G but does not seriously impair suppression [85% suppression]. From these results, we conclude that a stable stem I structure is essential for U1 snRNA to function in splicing.
Table 1. Suppressor activities and cellular abundances of the U1-4u mutants

| Mutant            | Suppressor activity (%) | Abundance (%) |
|-------------------|-------------------------|---------------|
|                   | Exp. 1 | Exp. 2 | Exp. 3 | Average | Stem mutants |
| 25A               | 10     | 4      |        |         | 15           |
| 40U               | 11     | 14     |        | 13      | 20           |
| 41C               | 22     | 15     |        | 19      | 32           |
| 24A               | 27     | 13     |        | 20      | 34           |
| 23U               | 43     | 39     |        | 41      | ND           |
| 19U               | 41     | 47     |        | 44      | ND           |
| 19G               | 65     | 57     |        | 61      | ND           |
| 25A/40U           | ND     | 78     |        | 78      | 151          |
| 22G               | 92     | 77     |        | 85      | ND           |
| 43A               | 64     | 109    |        | 87      | 88           |
| 20A               | 114    | 143    |        | 129     | ND           |
| Loop mutants      |         |        |        |         |              |
| 27A/31U           | 1      | 0      | 0      | 0       | ND           |
| 31U               | 2      | 0      | 0      | 1       | 8            |
| r28-31            | ND     | 22     | 15     | 19      | 25           |
| 36U               | 25     | 43     | 27     | 32      | 33           |
| 32C               | 57     | 51     | 31     | 46      | 60           |
| 29C               | 56     | 56     | 35     | 49      | 45           |
| 34U               | 62     | 69     | 38     | 56      | ND           |
| 38U               | 66     | 70     | 43     | 60      | ND           |
| 35U               | 52     | 85     | 67     | 68      | ND           |
| 33G               | 95     | 99     | 65     | 86      | ND           |
| 33A               | 100    | 133    | 92     | 108     | 122          |
| Insertion mutants |         |        |        |         |              |
| +G                | 34     | 34     |        | 34      | ND           |
| +GG               | 29     | 19     |        | 24      | 105          |

a Percent relative to U1-4u.
b The abundance of stem and loop mutants was determined by densitometry of the autoradiogram shown in Fig. 3. Percentages are relative to U1-4u. The abundance of the + GG insertion mutant was determined by densitometry of the autoradiogram shown in Fig. 6. Due to proximity of the + G and wild-type U1 bands, the abundance of the + G mutant could not be quantified by densitometry; however, visual comparison of two different exposures of the gel shown in Fig. 6 suggest that the + G and + GG mutants are equally abundant. Percentage is relative to internal control U1.

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Mutations that inhibit suppression severely destabilize U1 snRNA

As a first step toward understanding how mutations in stem-loop I of U1-4u affect suppressor activity, we determined the level of suppressor U1-4u RNAs in transfected cells using a primer extension assay. We used a synthetic oligodeoxynucleotide primer complementary to nucleotides 8-23 of U1 and performed a primer extension assay with the Klenow fragment of DNA polymerase I in the presence of dTTP, dCTP, and dATP. Klenow polymerase gives more faithful transcription than reverse transcriptase when one dNTP is omitted from the reaction (Zhuang and Weiner 1986). This assay allowed us to distinguish the suppressor U1-4u RNA from the endogenous U1 because the extension product for wild-type U1, which has a C residue at position 4, will be 4 nucleotides shorter than that for U1-4u in the absence of dGTP. To compare different U1-4u mutants, we cotransfected each of them with another U1 mutant as an internal control. The internal control U1 was made by replacing nucleotides 3 and 4 of U1 with a pentanucleotide sequence (GAAUU) that does not include a C residue. Thus, in the same primer extension assay, the extension product for the internal control U1 is 3 nucleotides longer than for U1-4u (Fig. 3; the major products for U1-4u and the internal control U1 are 1 nucleotide shorter than expected, presumably because of the 5'-terminal cap structure).

Primer extension assays performed on some U1-4u mutants are shown in Figure 3 and summarized in Table 1. The parental U1-4u RNA is synthesized abundantly (Fig. 3, lanes 2 and 9) and, as observed previously, may represent as much as 25% of the total U1 in transfected cells (Zhuang and Weiner 1986). The internal control U1 is as abundant as U1-4u RNA. Mutations that have a severe effect on suppression also yield low levels of suppressor RNA (40U, 25A, 41C, 24A, 31U, and r28-31), top of stem I, likewise has only a modest effect, because other mutations that disrupt the stem have relatively severe effects (see Fig. 2A), the U27 : G38 base pair, if it exists, may contribute little to the stability of stem I. Finally, as expected from the behavior of the parental mutation 31U, the double-point mutation 27A/31U also abolishes suppression.
Table 2. Phylogenetic comparison of the 5' end and stem-loop I of U1 snRNA

|                  | 5' Splice site | Stem I [5' half] | Loop I | Stem I [3' half] |
|------------------|----------------|------------------|--------|------------------|
| Human            | AUACUUACCUG    | GCAGG            | GAAGAUACCAU | GAUCAGGAAG |
| Rat              |                |                  | G- C-     |                 |
| Mouse            |                |                  | U-         |                 |
| U1a              |                |                  | C-         |                 |
| U1b-2            |                |                  | A-         |                 |
| U1b-5            |                |                  | A-         |                 |
| Chicken          |                |                  | G-         |                 |
| Xenopus          |                |                  | C-         |                 |
| Drosophila       |                |                  | C-         |                 |
| Sea urchin       |                |                  | C-         |                 |
| Bean             |                |                  | C-         |                 |
| Soybean          |                |                  | C-         |                 |
| Green alga       |                |                  | U- C-      |                 |
| Yeast            | AAGAU          | AUC- G- GG-       | G- A-      |                 |

The 5’ cap nucleotide is not shown. U residues at positions 5–6 are actually pseudouridines in some species. Only bases that differ from human U1 snRNA are indicated, and spaces are introduced into the sequence for comparison. The bulged nucleotides in the stem are underlined. Sequences for bean, soybean, yeast, and part of Xenopus U1 are derived from gene sequences. References for the U1 sequences are as follows: (Human) Branlant et al. (1980); (rat) Branlant et al. (1980); (mouse) Kato and Harada [1985]; (chicken) Branlant et al. (1980); (Xenopus) Forbes et al. (1984); Zeller et al. (1984); Krol et al. (1985); (Drosophila) Mount and Steitz (1981); (sea urchin) Yu et al. (1986); (bean) van Santen and Spritz (1987); (soybean) van Santen et al. (1988); [green alga (Chlorella saccharophila)], Kiss et al. (1988); (yeast) Kretzner et al. (1987); Siliciano et al. (1987).

Whereas mutations that have little effect on suppression produce high levels of suppressor RNA [25A/40U, 33A, and 43A; although the unusual abundance of 25A/40U and 33A could be real, it is likely to reflect the range of experimental error in a single data set]. Because mutations within the U1-coding region do not appear to affect the efficiency of transcript-initiation or 3'-end formation (Skuzeski et al. 1984; Hernandez and Weiner 1986; Neuman de Vegvar et al. 1986; Hamm et al. 1987), we conclude that most mutations in stem-loop I that interfere with U1 suppressor activity do so by destabilizing U1 snRNA or the U1 snRNP.

Severe mutations 25A and 31U affect binding of U1 snRNP-specific proteins

Stem-loop I recently has been shown to be the binding site for the U1 snRNP-specific proteins [Hamm et al. 1987, Patton et al. 1987]. As an enriched source of snRNP proteins, we treated HeLa cell nuclear extract with micrococcal nuclease to digest the U-snRNAs and liberate free snRNP proteins, and inhibited micrococcal nuclease with EGTA to chelate calcium ions. Pretreatment with micrococcal nuclease increases the amount of U1-4u SP6 RNA that can be precipitated with anti-(U1)RNP serum from <5% to ---25% of the input RNA (Fig. 4A). When we tested U1-4u SP6 RNAs containing the second-site mutation 25A or 31U (Fig. 4B), the mutant RNA was precipitated about fivefold less efficiently by anti-(U1) RNP serum than was the parental U1-4u SP6 RNA. We conclude that mutations 25A and 31U reduce the affinity of U1 snRNA for U1 snRNP-specific proteins.

Suppressor activity is reduced by insertions at position 11 of U1-4u

Nucleotides 3–11 at the 5’ end of U1 snRNA are exposed on the surface of the U1 snRNP [Lerner et al. 1980, Rinke et al. 1984] and are thought to form base pairs with the 5’ splice site [Lerner et al. 1980, Rogers and Wall 1980; Zhuang and Weiner 1986; Seraphin et al. 1988; Siliciano and Guthrie 1988]. To test whether the position of U1 nucleotides 3–11 is rigidly specified relative to the body of the U1 snRNP, we inserted either one or two G residues at position 11 of U1-4u. These mutations, designated + G and + GG, reduced suppressor activity to ~34% and 24%, respectively, as determined by the RNase protection assay (Fig. 5 and Table 1). We also determined the level of these two RNAs in transfected...
cells by using a synthetic oligodeoxynucleotide complementary to nucleotides 17–46 of U1 to prime reverse transcriptase in the presence of all four deoxynucleoside triphosphates. The extension products for +G, +GG and the internal control U1 RNA will be longer than that for endogenous U1 by 1, 2, and 3, nucleotides, respectively. As shown in Figure 6A, the +G and +GG RNAs are as abundant as the internal control U1, which was shown above to be as abundant as U1-4u RNA [Fig. 3]. The +G RNA can be immunoprecipitated by anti-Sm and anti-[U1]RNAP antibodies [Fig. 6B], indicating that it is assembled efficiently into a snRNP. Thus, the +G and +GG mutations appear to allow efficient assembly of a relatively inactive snRNP. This suggests, as might have been expected from phylogenetic evidence (see Table 2 below), that changes in the position of the 5′ end of U1 RNA relative to stem-loop I interfere with U1 snRNP function.

Discussion

Functional aspects of the conserved U1 sequence

We chose initially to make point mutations in stem-loop I of U1 because the structure of this stem, the sequence of the loop, and the position of the stem-loop relative to the 5′ end of U1 snRNA have been highly conserved through evolution. We found that all three phylogenetically conserved features are important for the stability and/or function of U1 snRNA. As shown in Table 2, the base of stem-loop I is always found 16 nucleotides from the 5′ end of U1 snRNA. The loop contains 10 nucleotides (or 12 nucleotides if the base-pairing at the top of the stem does not exist), and the stem consists of 9 or 10 bp interrupted by one or two bulged nucleotides. The sequence of the stem can vary, but most of the loop nucleotides are conserved.

Phylogenetic comparison suggests that the structure of the stem, but not the sequence, is important for function [Fig. 2A]. As expected, point mutations that weaken the stem weaken suppression [e.g., 25A or 40U]; compensatory base changes that restore the stem restore suppression [25A/40U]; and substitution of a presumably stronger A : U base pair for the normal G : U base pair may even increase suppression [20A]. Phylogenetically, the bulge in stem I can be U, C, A, or even two U residues, and we confirm this phylogenetic flexibility by showing that mutation of the bulged nucleotide from U to G (22G) has little effect on suppression [we assume the mutation does not cause rearrangement of the base-paired stem]. Mutation 38U might have been expected to disrupt a potential U27 : G38 base pair at the top of stem I but has only modest effects compared to other mutants that disrupt the stem. Thus, the U27 : G38 base pair may contribute little to the stability of stem I or may not exist at all (see also Kretzner et al. 1987).

Except for position 33, the loop sequence is highly but not absolutely conserved; all 4 bases can be found at position 33, which is the only nucleotide in this loop that differs between human U1 and a variant mouse U1 (Kato and Harada 1985) or the U1 of Xenopus (Zeller et al. 1984), sea urchin (Yu et al. 1986), soybean (van Santen et al. 1988), and budding yeast (Kretzner et al. 1987; Siliciano et al. 1987). As expected on phylogenetic grounds, nucleotide 33 can be changed without significantly affecting suppression [Fig. 2B, mutants 33G and 33A]. Also consistent with the phylogenetic data, increasing the size of the loop decreases suppression (mutant r28-31). Interestingly, although some mutations in highly conserved loop nucleotides have a severe effect on suppression (31U), others have less than a twofold effect [29C, 32C, 34U, 35U, 36U, and 38U]. These data suggest that a twofold effect on function may be sufficient to fix the sequence of U1 snRNA on an evolutionary timescale. This observation is particularly interesting because human U1 is encoded by a large homogeneous multigene family whose copy number could, in principle, expand to compensate for deleterious muta-

Figure 3. Expression of second-site mutants of U1-4u in transfected cells, as determined by primer extension assay. HeLa cells were transfected with the adenovirus E1A hr440 mutant, different second-site mutants in stem-loop I of U1-4u, and an internal control U1 gene (see text). Using total RNA from transfected cells as template, an oligodeoxynucleotide complementary to nucleotides 8–23 of U1 was used to prime DNA synthesis by the Klenow fragment of DNA polymerase I in the presence of dTTP, dCTP, and dATP. Extension products were resolved on a 15% denaturing polyacrylamide gel. The hr440 mutant and the internal control U1 gene were introduced into cells with pUC13 vector alone (lanes 1, 8), the parental U1-4u gene (lanes 2, 9), stem mutants (lanes 3–7, 16), or loop mutants (lanes 10–15).
Figure 4. Effect of the 31U and 25A mutations on binding of U1-specific proteins in vitro. (A) A labeled SP6 transcript spanning nucleotides 1–75 of U1-4u was incubated in a HeLa nuclear extract with (+) or without (−) prior treatment with micrococcal nuclease (MN) and then immunoprecipitated with anti-[U1]RNP serum (lanes U1) or nonimmune serum (lanes NI). RNA from precipitates (ppt) and supernatants (sup) was resolved on a 6% denaturing polyacrylamide gel. (B) Labeled SP6 transcripts spanning nucleotides 1–75 of U1-4u and the severe second-site mutants 31U and 25A were incubated in HeLa nuclear extracts pretreated with micrococcal nuclease. Following immunoprecipitation with anti-[U1]RNP serum, total RNA (total), RNA from precipitates (ppt), and RNA from supernatants (sup) was resolved on a 6% denaturing polyacrylamide gel. Labeled SP6 transcripts span nucleotides 1–75 or U1-4u [lanes 1, 5, 9], the second-site mutant 31U [lanes 2, 6, 10], the second-site mutant 25A [lanes 3, 7, 11], or an unrelated RNA derived from the third exon of the rabbit β-globin gene, as described in Methods (lanes 4, 8, 12).

Do mutations in stem-loop I affect U1 snRNP assembly or function?

For all mutations tested in stem-loop I (Fig. 3), mutations that affect suppression severely lead to low levels of suppressor RNA, whereas mutations that have little effect on suppression produce high levels of suppressor RNA. The correlation between suppression and abundance suggests that mutations in stem-loop I usually interfere with suppressor activity by destabilizing U1 snRNA or the U1 snRNP. As a result, it is possible that many or all of the mutations we have characterized in stem-loop I affect the stability of U1 snRNA or the U1 snRNP rather than the ability of the mutant U1 snRNPs to participate in splicing. In principle, the low levels of certain mutant U1 RNAs could also be explained by poor transcription; however, this would not explain the effect of the compensatory base change in stem I (25A/40U), and it also seems unlikely in light of evidence (Skuzeski et al. 1984; Hernandez and Weiner 1986; Neuman de Vegvar et al. 1986; Hamm et al. 1987) suggesting that mutations within the U1-coding region do not affect the efficiency of transcriptional initiation or 3' end formation (for review, see Dahlberg and Lund 1988).
sequence. Unlike mutations in stem-loop I, insertion of one or two G residues at position 11 decreases suppression (Fig. 5, mutants +G and +GG) without affecting the abundance of the snRNAs or their assembly into snRNPs, as judged by precipitation with anti-Sm and anti-(U1)RNP antibodies (Fig. 6). Although we cannot say with certainty how +G and +GG affect splicing, phylogenetic conservation of the distance between the 5′ end of U1 and stem-loop I suggests that this spacing is important. Proper spacing might be required for binding of the U1 snRNP to the mRNA precursor and/or for correctly positioning the RNA duplex formed between the 5′-terminal sequence of U1 and the 5′ splice site (Zhuang and Weiner 1986; Seraphin et al. 1988; Siliciano et al. 1987), relative to the rest of the spliceosome so that the cleavage and ligation reactions can occur.

In conclusion, we have begun a second-site reversion analysis of mammalian U1 snRNA. We showed that a stable stem I structure and the conserved nucleotides within loop I are essential for U1 snRNA function. However, many mutations in stem-loop I were found to affect splicing indirectly by destabilizing U1 snRNA or affecting U1 snRNP assembly. Further analysis will be needed to identify U1 nucleotides that are directly involved in mRNA splicing. Ultimately, we hope to identify any U1 nucleotides that might base-pair with other snRNAs in the course of splicing and to test the nature of these interactions by making compensatory base changes in both snRNAs.
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Methods

Mutant constructions

The template for site-directed mutagenesis of U1-4u was an M13 mp8 clone containing the U1-4u gene as a 681-bp HaeIII fragment fitted with BamHI linkers (Zhuang and Weiner 1986). To make point mutations in stem-loop I of U1-4u, a mixed 30-mer oligodeoxynucleotide complementary to positions 17–46 of U1 was synthesized so that each position was, on average, 97%, wild type and 1% each of the other three nucleotides. Site-directed mutagenesis was performed as described (Kunkel 1985). Mutants were identified by dideoxy sequencing of single-stranded DNAs from the recombinant phage, using a primer complementary to positions 121–142 of U1. Mutant U1-4u genes were recloned into the pUC13 vector as a HindIII–EcoRI fragment for use in transfection. Mutant 25A/40U was constructed by replacing the EcoRI–BclI fragment (extending from position –425 in 5’ flank to position 27 in U1-coding region) of mutant 40U in pUC13 with the corresponding fragment from mutant 25A. Mutant r28-31 was constructed by cutting the U1-4u gene in pUC13 at position 27 with BclI, filling the ends in with Klenow fragment, and religating. To make mutants + G, + GG, and internal control U1, site-directed mutagenesis was performed (as above) using the oligodeoxynucleotides 5’-TCTCCCTGGCCAGGTAAATA-3’, 5’-TCTCCCTGCAGGTAAATA-3’, and 5’-TCTCCTGAGGTAAATA-3’, respectively.

To construct SP6 clones used in snRNP protein-binding experiments (Fig. 4), the BgIII–BamHI fragments (extending from position –6 in 5’ flank to position +91 in 3’ flank) of the U1-4u gene as well as mutants 25A and 31U, were isolated and cut at position 73 of the coding region with HpaII. The BgIII–HpaII fragments were filled in with Klenow fragment and cloned between the filled-in HindIII and SalI sites of the vector pSP64 (Melton et al. 1984). The non-U1 SP6 clone was constructed by cloning a PvuII–BglII fragment of the rabbit β-globin gene (extending from position –9 in 5’ flank to position 1195 in exon 3) in opposite orientation between the BamHI and Smal sites of the vector pSP64. DNA from this clone was linearized with EcoRI at position 1115 to give a 118-nucleotide transcript.

Transfection and RNA preparation

Transfections were performed as described previously (Hernandez 1985), using 6 μg of the U1-4u gene in pUC13 and 6 μg of plasmid pSVhr440 (Zhuang and Weiner 1986) per 6-cm plate. About 48 hr after transfection, cytoplasmic RNA was extracted from the postnuclear supernatant of cells lysed with NP-40 (Hernandez 1985). In experiments comparing the abundance of various suppressor RNAs, HeLa cells were transfected with 4 μg each of the U1-4u genes, the internal control U1 gene, and pSVhr440. Total RNA was prepared from these cells lysed with SDS and proteinase K. Following phenol–chloroform extractions, one volume of 95% ethanol was added and the resulting DNA clump was removed. Another volume of 95% ethanol was then added to precipitate the RNA.

RNase protection analysis

RNase mapping was performed essentially as described (Melton et al. 1984). RNase digestion was allowed to proceed for 30 min at 30°C and stopped with 2% SDS. The samples were then treated with 400 μg/ml of proteinase K at 37°C for 15 min, extracted with phenol–chloroform, ethanol-precipitated, and fractionated on denaturing polyacrylamide gel.

Primer extension

Synthetic oligodeoxynucleotide primers were labeled at the 5’ end by kinasing. For mutants in stem-loop I of U1-4u, primer extension was performed with the Klenow fragment of DNA polymerase I at 37°C for 20 min in the presence of 38 μM each of dTTP, dCTP, and dATP (Zhuang and Weiner 1986). For the + G and + GG mutants, primer extension was performed with reverse transcriptase as follows. RNA was mixed with the labeled primer in 12 μl of 125 mM KCl and 83 mM Tris-HCl (pH 7.6), heated to 85°C for 2 min, incubated at 65°C for 10 min, and allowed to cool to room temperature for 30 min. The mixture was made 3 mM in MgCl₂, 10 mM in DTT, 50 μg/ml in actinomycin D, and 0.5 mM in each dNTP; 200 units of M-MLV reverse transcriptase [BRL] was added per 20-μl reaction, and the reaction was incubated at 37°C for 1 hr. Reactions were stopped with 25 mM EDTA, ethanol-precipitated, and fractionated on a denaturing polyacrylamide gel.

In vitro snRNP protein binding and immunoprecipitation

HeLa cell nuclear extracts (Dignam et al. 1983) were made 1 mM in CaCl₂, treated with 50 U/μl of micrococcal nuclease [Cooper Biomedical] at 37°C for 30 min, and the reaction stopped with 5 mM EGTA. Labeled SP6 RNA of sp. act. >10⁶ cpm/μg was synthesized as described (Melton et al. 1984) and gel-purified. About 10⁵ cpm of the SP6 RNA was incubated in a 15-μl reaction containing 60% (vol/vol) micrococcal nuclease-treated extract, 20 mM creatine phosphate, 0.5 mM ATP, 1 U/μl RNasin [Promega], and 1 μg/μl poly(A) [Sigma] at 37°C for 30 min. The reactions were then immunoprecipitated with anti-(U1)RNP serum [Pettersson et al. 1984], as described [Mimori et al. 1986]. Assuming complete recovery of cellular U1 in the nuclear extract, there should be at least a fivefold excess of endogenous U1 snRNP proteins over added SP6 U1 transcripts under these assay conditions.

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