A protein of the SR family of splicing factors binds extensively to exonic Balbiani ring pre-mRNA and accompanies the RNA from the gene to the nuclear pore

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We report on the molecular cloning and intracellular localization of a heterogeneous nuclear ribonucleoprotein (hnRNP), Ct-hrp45, one of the major components of pre-mRNP particles in Chironomus tentans. It is shown that hrp45 belongs to the SR family of splicing factors and exhibits high sequence similarity to Drosophila SRp55/B52 and human SF2/ASF. The distribution of hrp45 within the C. tentans salivary gland cells is studied by immunocytology. The hrp45 protein is found to be abundant in the nucleus, whereas it is undetectable in the cytoplasm. The fate of hrp45 in specific pre-mRNP particles, the Balbiani ring (BR) granules, is revealed by immunoelectron microscopy. It is observed that hrp45 is associated with the growing BR pre-mRNP particles and is being added continuously concomitant with the growth of the transcript, indicating that hrp45 is bound extensively to exon 4, which comprises 80–90% of the primary transcript. Furthermore, hrp45 remains bound to the BR RNP particles in the nucleoplasm and is not released until the particles translocate through the nuclear pore. Thus, hrp45 behaves as an hnRNP protein linked to exon RNA (and perhaps also to the introns) rather than as a spliceosome component connected to the assembly and disassembly of spliceosomes. It seems that hrp45, and possibly also other SR family proteins, is playing an important role in the structural organization of pre-mRNP particles and is perhaps participating not only in splicing but also in other intranuclear events.

[Key Words: SR proteins, hnRNP proteins; RNP particles; exon RNA; nucleo-cytoplasmic transport; nuclear pore]

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SR proteins comprise a family of evolutionarily conserved pre-mRNA splicing factors [Zahler et al. 1992; Birney et al. 1993; Fu 1995]. Each protein contains one or two amino-terminal ribonucleoprotein [RNP]-type RNA recognition motifs [RRM] and a carboxy-terminal region rich in arginine–serine dipeptide repeats [RS domain]. The SR proteins were identified initially because of a common epitope recognized by a monoclonal antibody [mAb 104], and they could be easily purified by a simple two-step salt precipitation procedure [Roth et al. 1991]. Many organisms, ranging from Drosophila to humans, were found to contain a set of SR proteins of similar molecular masses: 20, 30, 40, 55, and 75 kD [Zahler et al. 1992]. In humans, the 30-kD fraction was shown to comprise two separate proteins, SRp30a and SRp30b [Zahler et al. 1992], that had earlier been described as SF2/ASF [Ge et al. 1991; Krainer et al. 1991] and SC35/PR264 [Fu and Maniatis 1992; Vellard et al. 1992], respectively. Each SR protein seems to have been strictly conserved during evolution; the SR protein of a given molecular mass is more similar to those of similar size in other species than it is to other SR proteins within its own species [Roth et al. 1991]. New members of the SR family have been identified recently, and it is conceivable that the SR family is considerably larger than initially assumed [Blencowe et al. 1995; Neugebauer et al. 1995].

The SR proteins are essential splicing factors involved in both spliceosome assembly and the splicing process [for review, see Fu 1995]. Different SR proteins seem to have similar, but not identical, functions in constitutive and regulated splicing. The individual SR proteins can restore general splicing activity to an S 100 extract [Ge et al. 1991; Krainer et al. 1991; Mayeda et al. 1992; Zahler et al. 1992; Scretan et al. 1995], but the activity differs between the SR proteins and is dependent on the pre-mRNA species tested [Fu 1993]. The SR proteins can also
affect the choice of alternative splice sites in in vitro experiments [Ge et al. 1991; Krainer et al. 1991; Fu et al. 1992; Mayeda et al. 1992], but the individual SR proteins can favor different splice sites [Kim et al. 1992; Zahler et al. 1993a]. Furthermore, there are also differences between the various SR proteins as to how they interact with exonic splicing enhancer elements [Sun et al. 1993b; Tian and Maniatis 1993]. Thus, it seems likely from the in vitro experiments that each SR protein has a unique function in constitutive and alternative splicing.

More recently, the effects of SR proteins have also been evaluated in vivo. In transfection experiments, it was shown that SR proteins can modulate alternative splicing in the same way as observed in vitro [Caceres et al. 1994; Srebot et al. 1995; Wang and Manley 1995]. However, differences were noted, for example, Wang and Manley (1995) observed that both SRp30a [SF2/ASF] and SRp30b [SC35] inhibit splicing of SV40 early pre-mRNA in vivo, whereas they stimulate in vitro. If SRp55/B52 is overexpressed in transgenic Drosophila flies, it affects the development of the organism seriously [Kraus and Lis 1994]. In a more recent study, the corresponding gene was deleted, which interrupted development, indicating that SRp55/B52 plays a critical role in Drosophila development (Ring and Lis 1994). It was, however, not possible in this study to establish that splicing had been changed, although the splicing patterns of five specific pre-mRNAs were scrutinized. It can be concluded that the in vivo experiments confirm that SR proteins can influence the splicing process, but the in vivo situation is much more complex and the results are not easily interpreted. It is, of course, most important that the in vivo experiments clearly demonstrate that at least one of the SR proteins is critical for proper development (Ring and Lis 1994). Whether this effect is a result of aberrant splicing or is connected to failure of another cellular function remains to be demonstrated.

The SR proteins are present at the sites of transcription, which has been shown most convincingly in studies of the chromosome loops in amphibian oocytes [Roth et al. 1990] and the chromosome puffs in dipteran salivary glands [Champlin et al. 1991; Champlin and Lis 1994; Baurén et al. 1996]. In mammalian cells, SR proteins are distributed in a speckled pattern [Fu and Maniatis 1990; Roth et al. 1990; Spector et al. 1991; Carter et al. 1993; Wansink et al. 1993] that on the ultrastructural level corresponds to perichromatin fibrils and interchromatin granule clusters [Spector et al. 1991]. As perichromatin fibrils are known to contain newly synthesized heterogeneous nuclear RNA [hnRNA] [for review, see Fakan and Puvion 1980], this result supports the view that SR proteins associate with hnRNA, that is, pre-mRNA, already during transcription. The early appearance of SR proteins on nascent transcripts is in agreement with the observation that splicing can occur cotranscriptionally [Osheim et al. 1985; Beyer and Osheim 1988; Le Maire and Thummel 1990; Baurén and Wieslander 1994]. As splicing also takes place post-transcriptionally [for review, see Nevins 1983], it is to be expected that SR proteins are associated with at least some pre-mRNP in transit between the site of synthesis and the nuclear envelope. It has been difficult, however, to be more specific on the behavior of SR proteins in relation to the synthesis and flow of pre-mRNP particles, the light microscopic analysis in combination with in situ hybridization has been hampered by low resolution, and the ultrastructural analysis has not yet provided proper identification of gene-specific RNP fibrils or granules.

The precise role of SR proteins in the transcriptional and post-transcriptional events would benefit very much from studies of defined pre-mRNP particles carried out on the ultrastructural level with a concomitant assessment of the distribution of SR proteins. Now it has become feasible to perform such a study on the Balbiani ring transcription products in the salivary glands of the dipteran Chironomus tentans. The salivary gland cells contain polytene chromosomes, and the transcriptionally active regions appear as chromosomal puffs (Daneholt 1982). On chromosome IV, there are two giant puffs with exceptionally intense RNA synthesis: the Balbiani rings 1 and 2 [BR1 and BR2]. The BR1 and BR2 genes, 35–40 kb in size, belong to the same gene family and exhibit a similar exon–intron organization with five exons and four introns [for review, see Wieslander 1994]. A remarkable feature of the BR transcripts is that one of the exons (number 4) is exceptionally long (>30 kb), whereas the introns and the other exons are relatively small (<1.5 kb) and located close to the 5′ or 3′ ends [Wieslander and Paulsson 1992]. The three introns close to the 5′ end seem to be spliced before one-third of the transcript is synthesized, whereas the fourth intron close to the 3′ end is usually spliced post-transcriptionally [Baurén and Wieslander 1994]. Small nuclear RNP particles [snRNPs] appear transiently on the growing RNP fibers at or close to the introns, reflecting the rapid assembly and disassembly of spliceosomes [Kiseleva et al. 1994]. The synthesis of BR RNA and the packing of the RNA into RNP particles concomitant with transcription can be visualized in the electron microscope [Andersson et al. 1980; Olins et al. 1980; Skoglund et al. 1983]. The completed product, a spherical particle with a diameter of 50 nm, can be observed in the nucleoplasm [Stevens and Swift 1966; Skoglund et al. 1986; Mehnli et al. 1992] and also during translocation through the nuclear pore [Stevens and Swift 1966; Mehnli et al. 1992, 1995]. Thus, the formation of the BR transcription product on the gene, the intranuclear transport, and the delivery of the transcript to the cytoplasm can be scrutinized on the ultrastructural level. Furthermore, with immunoelectron microscopy [immuno-EM] it has been demonstrated recently that the flow of a specific particle protein can be followed from the gene through the nuclear pore and into polysomes [Visa et al. 1996].

In the present study, we apply the BR system in the analysis of a specific SR protein. We have shown recently that the BR RNP particles contain a protein with an apparent molecular weight of 45 kD, designated Ct-hrp45 (or hrp45 for convenience) [Wurtz et al. 1996]. Now we demonstrate that hrp45 possesses a high degree of sequence similarity to members of the SR protein family of
splicing factors, including human SF2/ASF and Drosophila SRp55/BS2. Using immuno-EM, we show that the putative splicing factor hrp45 binds to BR pre-\(\text{RNA}\) at multiple sites along the large exon, and perhaps along the entire transcript; this is in striking contrast to the behavior of snRNPs, which appear selectively at the intron regions of the transcript [Kiseleva et al. 1994]. Furthermore, hrp45 accompanies BR pre-\(\text{RNA}\) from the gene to the nuclear pore, where it is shed. We suggest that hrp45 is important for the structural organization of pre-m\(\text{RNA}\) particles and could be involved not only in splicing but also in other nuclear events.

Results

Cloning and characterization of hrp45

The mouse monoclonal antibody 2E4 recognizes the C. tentans hnRNP protein hrp45 [Wurtz et al. 1996]. The specificity of the 2E4 antibody was demonstrated by Western blot analysis of nuclear and cytoplasmic samples from C. tentans tissue culture cells (Fig. 1). The 2E4 antibody reacts specifically with a 45-kD protein, hrp45, that seems to be present in the nucleus (Fig. 1, lane a) but not in the cytoplasm (Fig. 1, lane b).

To determine the primary structure of hrp45, we used the 2E4 antibody to screen a C. tentans salivary gland, random-primed, lambda gt11 cDNA library. Two cDNA clones, p1E1 and p1E9, were isolated. The sequence of their cDNA inserts turned out to be identical [data not shown]; they were 907-bp long and contained a 109-bp 5’ untranslated sequence (UTR) followed by an open reading frame (ORF) of 788 bp with no stop codon at the 3’ end. Thus, the two clones harbored only a partial coding sequence.

To obtain a cDNA clone with the complete coding sequence, the PCR amplification product of the cDNA clone p1E1 was labeled with digoxigenin [DIG] and used as a probe to screen a C. tentans salivary gland, oligo(dT)-primed, lambda ZAP cDNA library. Seven different cDNA clones were isolated and analyzed by sequencing. The nucleotide sequence of these clones comprises the full sequence of the insert in the p1E1 and the p1E9 clones and additional sequences both at the 5’ and 3’ ends. The longest of the clones, pHRP45.1, presented in Figure 2, was shown to have a 1381-bp insert, including a 5’ UTR (133 bp), a coding region (966 bp), and a 3’ UTR (262 bp) followed by a poly[A] stretch. The 5’ UTR sequence preceding the initiation codon contains eight termination codons in frame. The 3’ UTR sequence has a single polyadenylation signal AATAAA, which is located 15 nucleotides upstream of the 3’ poly[A] stretch, and is likely to represent an authentic 3’ end of the corresponding mRNA according to the requirements for a polyadenylation region [for review, see von Heijne 1988].

The putative translation initiation site, AAGATG-GTT, is in agreement with Kozak’s rule for initiation site consensus in eukaryotic mRNAs [Kozak 1986]. The ORF in the pHRP45.1 sequence translates into 322 amino acids, starting at position 133 and ending with a stop codon at position 1101; the predicted molecular weight of the protein is 37,420. Two amino acids are especially abundant, arginine (65 residues, or 20% of the total amino acid content) and serine (52 residues, or 16% of the total content), and both are mainly located in the carboxy-terminal part. At positions 193–212 in the carboxy-terminal part, there is a putative bipartite nuclear localization signal [Dingwall and Laskey 1991]. The ORF sequence of the clone HRP45.1 was confirmed by analysis of the six additional cDNA clones.

To demonstrate that the protein encoded in the isolated cDNA clones is indeed recognized by the 2E4 antibody, the protein was expressed in E. coli. A protein of the expected molecular size, 37–38 kD, appeared among the bacterial proteins (Fig. 1B, cf. lanes a and b), and in a Western blot experiment the antibody reacted exclusively with this protein (Fig. 1B, cf. lanes c and d). It
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hrp45 is an analog to the mammalian RNA splicing factor SF2/ASF and the Drosophila SRp55/B52 protein. Searches of the EMBL DNA data base (Rice et al. 1993) and the SWISS-PROT data base (Bairoch and Boeckmann 1993) revealed that hrp45 has a strong sequence similarity to the proteins of the SR family of RNA splicing factors. The most characteristic feature of these proteins is the RS domain (Zahler et al. 1992; Birney et al. 1993). The highest score, ~78% identity and ~84% similarity, was obtained with the Drosophila protein SRp55/B52 (Champlin et al. 1991; Roth et al. 1991). Furthermore, hrp45 shows ~48% identity and ~64% similarity to the human splicing factor SF2/ASF (Ge et al. 1991; Krainer et al. 1991). A direct alignment of hrp45 with SRp55 and SF2 highlights the sequence conservation between the three proteins (Fig. 3A). Sequence conservation is higher in the amino-terminal than in the carboxy-terminal part.

SF2/ASF has a modular primary structure comprising two RRMs and an auxiliary RS domain (Ge et al. 1991; Krainer et al. 1991); a small fourth domain, exhibiting a weak similarity to the RS domain, has also been recognized more recently between the two RRMs (Kim et al. 1991). As shown in the DOTPLOT comparison in Figure 3B, hrp45 has the same modular structure; regions A and C correspond to the two RRMs, region B to the small intervening domain, and region D to the carboxy-terminal RS domain. The sequences of the two RRMs in hrp45 are very similar to the RRMs in SF2 and Drosophila SRp55 (Fig. 3A). Region B in hrp45 is not only rich in arginines, but it is also relatively rich in glycines, although it does not contain a stretch of glycines, which is characteristic for SF2/ASF (Ge et al. 1991; Krainer et al. 1991). The RS domain in hrp45 is about three times longer than in SF2/ASF, but still shorter than the RS domain in SRp55/B52.

The conclusion that hrp45 is an SR protein is supported from several experimental observations on the properties of hrp45. First, as mentioned above, hrp45 is heavily phosphorylated, which is a characteristic of SR proteins (Roth et al. 1991; Zahler et al. 1993b). Furthermore, the mAb 104 antibody, which is known to be specific for SR proteins (Roth et al. 1991), reacts with a nuclear 45-kD protein in C. tentans tissue culture cells as shown by Western blotting, the hrp45 protein constitutes one of three predominant SR proteins (Fig. 1A, lane c). In a follow-up experiment, the 45-kD protein was immunoprecipitated from the nuclear extract by 2E4 and in a subsequent Western blot analysis shown to be recognized by the 104 antibody (data not shown). Finally, it has been observed that the 2E4 antibody also recognizes the SR protein SF2/ASF [A. Alzhanova-Ericsson, A. Kanopka, Y. Aissouni, G. Akusjärvi and B. Daneholt, in prep.], indicating that hrp45 and SF2/ASF have the 45-kD epitope in common, it should be noted that this epitope has to be different from the phospho-epitope recognized by mAb104 (Roth et al. 1991), as it is insensitive to dephosphorylation (Fig. 1C). Thus, the observed properties of cellular hrp45 are in agreement with the prediction from the cDNA sequence that hrp45 is an SR protein.

hrp45 is confined to the cell nucleus To determine the intracellular distribution of hrp45, the 2E4 antibody was used for immunocytology on semithin cryosections of fixed salivary glands. This assay at the light microscopy level provided information on the gen-
eral distribution of hrp45 in the cell compartments. Both the nucleus and the cytoplasm of the salivary gland cells could be readily identified by phase contrast microscopy (Fig. 4A). Observation of the same section under bright field conditions showed that immunostaining was restricted to the cell nucleus; no significant labeling was observed in the cytoplasm (Fig. 4A). The BRs were the most intensely labeled structures in the nucleus, but several other puffs as well as the nucleoplasm were stained. In the negative control, no labeling was detectable in either the nucleus or the cytoplasm of the salivary gland cells (Fig. 4B and B'). It was concluded that hrp45 is essentially confined to the cell nucleus (Fig. 1); the small amount of nascent hrp45 likely to be present in the cytoplasm is evidently below the detection limit of the method used.

hrp45 accumulates in the BR RNP complexes concomitant with transcription

To analyze the distribution of hrp45 along the BR genes, we carried out immuno-EM on isolated polytene chromosomes using the 2E4 antibody. The isolated chromosomes were fixed in 3.7% formaldehyde, challenged with antibodies, postfixed in 2% glutaraldehyde, embedded in plastic, sectioned, and stained (preemitting technique). The transcriptionally active BR genes could be visualized in the electron microscope as transcription loops (for review, see Daneholt 1992). At low magnification BR1 and BR2 were easily recognized and segments of transcription loops discerned within the BRs (Fig. 5A). The assembly of the BR RNP complexes could be followed along the gene; in the promoter-proximal region, the nascent RNP5s appear as thick RNP fibers growing longer and longer, whereas in the middle and distal regions the 5' end of the RNP fiber is being packed into a dense globular structure that increases in diameter along the gene. Loop segments representing the various portions of the BR genes are denoted in Figure 5B: Thick RNP fibers are seen in proximal portions, small stalked granules in middle portions, and large stalked granules in distal portions. In our immuno-EM experiments, we studied the binding of the 2E4 antibody to the three loop segments to establish when hrp45 is being incorporated into the growing RNP particles.

The results of the immuno-EM experiments are presented in Figure 5C–E. There are gold particles over all
Figure 4. Distribution of Ct-hr45 in the salivary gland cells. A
representative salivary gland section immunolabeled with mAb
2E4 is visualized and photographed by light microscopy. (A) and
(A') show bright field and phase contrast images, respectively, of
the same section. Note that the nucleus (NUC) is stained
whereas the cytoplasm (CYT) is not stained. The most strongly
labeled structures in the nucleus are the Balbiani rings (BR).
(B) and (B') show bright field and phase contrast images, respec-
tively, of a section treated with the negative control antibody.
Dashed lines in the bright field images outline the nuclear and
cytoplasmic compartments. Bar, 40 μm.

three regions: the proximal, middle, and distal regions. It
is also evident that the signal grows stronger in parallel
with the increase in size of the transcription product. No
significant labeling was observed in the negative control
(Fig. 5F). Thus, hr45 seems to associate with nascent
RNA early during transcription and is being added sub-
sequently concomitant with the growth of the tran-
script.

To determine more accurately the appearance of hr45
along the BR genes, the distribution of the gold particles
among the three gene segments was established for BR1
as well as for BR2 following the procedure described by
Kiseleva et al. (1994). The quantitative analysis of the
gold particles in the proximal, middle, and distal regions
of the BR1 and BR2 loops is presented in Table 1. The
increase in labeling along the gene is confirmed. Furth-
more, the relative distribution of hr45 in the proximal,
middle, and distal segments corresponds well to the pre-
dicted relative amounts of growing BR RNA in the three
segments (i.e., 1:3:5). Thus, it seems as if hr45 is being
added continuously to the growing RNP complex and
that hr45 is bound essentially along the entire length of
both the BR1 and BR2 transcripts.

The 35–40-kb BR transcripts consist of 80–90% exon
RNA; exon 4 is by itself 30–35 kb and one of the longest
exons known (Wieslander 1994). Therefore, the observed
distribution of hr45 along the entire transcripts strongly suggests that hr45 is bound extensively to
exon RNA. Whether hr45 is also bound to intron RNA
cannot be determined because the resolution of the
method does not allow a sufficiently detailed analysis of
the hr45 distribution at the 5' and 3' ends of the tran-
scripts, where the relatively short introns are located.

hr45 is present in nucleoplasmic BR particles and in
BR particles translocating through the nuclear pores

To further follow the fate of hr45, we studied the local-
ization of hr45 in BR RNP particles in sections of sali-
vary glands using immuno-EM and the 2E4 antibody. In
these experiments, the glands were fixed in 4% formal-
dehyde and 0.1% glutaraldehyde, cryoprotected, frozen,
and sectioned. The immunoreactions were carried out
on the sections, and subsequently the specimens were
stained and embedded in polyvinyl alcohol. In spite of
the relatively mild fixation, the ultrastructural morphol-
ogy was adequate to allow the BR RNP particles to be

Figure 5. Electron micrographs of transcriptionally active BR
genes and immuno-EM analysis of the distribution of hr45
along BR transcription units. An isolated chromosome IV is
shown in A; BR1 and BR2 are indicated by arrows. Proximal (p),
middle (m), and distal (d) regions of BR transcription units are
shown in B. Three examples of immunolabeling with mAb 2E4
against hr45 are given in C–E; note the gradient of accumula-
tion of gold labeling from proximal to distal regions of the tran-
scription unit. No labeling can be seen with the negative control
antibody (F). Bar in A, 5 μm; in B, 0.5 μm; and in F, 250 nm
(magnification the same in C–F).
Quantitative immuno-electron microscopy analysis of the distribution of hrp45 along transcription loops in BR1 and BR2

| Regions | BR1 | BR2 |
|---------|-----|-----|
| proximal | 699 (11.5%) | 720 (12.5%) |
| middle  | 1949 (32.2%) | 1818 (31.5%) |
| distal  | 3424 (56.3%) | 3241 (56.0%) |
| Total   | 6072 (100%) | 5779 (100%) |

The number of gold particles recorded in the proximal, middle, and distal regions of BR1 and BR2 transcription loops. BRs from six chromosomes (collected from six larvae) were studied; for each BR, three randomly chosen areas (4 x 6 μm) were selected for quantitative analysis. Each gold particle was referred to the proximal, middle, or distal region of a transcription loop.

As shown in Figure 6A, gold particles were found on growing BR RNP particles in the BRs, confirming the conclusion drawn above that hrp45 is being associated already with nascent BR RNA. The BR RNP particles are known to be released from the chromosome and appear in the nucleoplasm as dense RNP granules with a diameter of ~50 nm (e.g., Mehlin et al. 1992). In our immuno-EM experiments, the BR granules in the nucleoplasm were often decorated by gold particles; two examples are shown in Figure 6, B and C. Thus, hrp45 can still remain in the BR RNP particle during intranuclear transport.

BR RNP particles translocating through the nuclear pore are presented in Figure 6D–F. Labeling was observed on BR particles at this stage and the gold markers were visualized in the BRs, in the nucleoplasm, and during pore passage.

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BR RNP particles translocating through the nuclear pore are presented in Figure 6D–F. Labeling was observed on BR particles at this stage and the gold markers were
exclusively associated with the nuclear portion of the RNP particle in transit. As illustrated in Figure 6, E and F, the portion of the RNP fiber extending into the cytoplasm was unlabeled. This indicates that hrp45 can reach the pore associated with BR RNA but that it is released from the particle in conjunction with the translocation of the particle through the pore. No significant labeling was found in the cytoplasm of the salivary gland cells (Fig. 6G), which is in agreement with the light microscopy data presented above (Fig. 4). Thus, the hrp45 protein does not enter the cytoplasm with the BR RNA molecule.

We conclude that hrp45 is added onto the nascent transcript concomitant with transcription, accompanies the pre-mRNA to the nuclear pore, and is released from the transcript during the translocation of the particle through the pore. To our knowledge, this is the first time it has been shown that a specific hnRNP protein reaches the nuclear pore as a constituent of an RNP particle, and is shed at or within the nuclear pore complex, when the RNA enters cytoplasm.

Discussion

hrp45 is an SR protein

The sequence analysis of hrp45 revealed that hrp45 belongs to the family of SR proteins, which are known to act as pre-mRNA splicing factors. A high degree of sequence similarity was observed to human SF2/ASF (Ge et al. 1991; Krainer et al. 1991) and Drosophila SRp55/B52 proteins (Champlin et al. 1991; Roth et al. 1991). The hrp45 protein has the same domain structure as SF2/ASF (and SRp55/B52): it contains two RRM sequences (regions A and C), a relatively short arginine-rich region between the RRM regions (region B), and a carboxy-terminal RS domain (region D).

Similar to SRp55 (Mayeda et al. 1992), hrp45 can substitute for SF2/ASF in an in vitro constitutive splicing system (A. Alzhanova-Ericsson, A. Kanopka, Y. Aissouni, G. Akusjärv, and B. Daneholt, in prep.). Therefore, hrp45 is not only structurally similar, but also functionally related to human SF2/ASF and SRp55/B52, and probably to other SR proteins. The members of the SR family seem to have similar or overlapping functions in the splicing of different pre-mRNA substrates because there are observations of both redundancy and substrate specificity of these proteins (see the introductory section). We have shown that hrp45 can be detected only at some, but not all chromosome puffs on C. tentans chromosomes (Wurtz et al. 1996). This result is in agreement with the view that hrp45 is a splicing factor with a differential substrate specificity: It seems to act on some but not all pre-mRNA substrates.

hrp45 is distributed along the BR transcript

The BR system allowed us to investigate the localization of hrp45 along growing pre-mRNP particles in situ. It was found that hrp45 is added cotranscriptionally to the growing BR pre-mRNA, and the amount of hrp45 is increasing in parallel with the growth of the primary transcript (Fig. 7). This result is in agreement with the earlier observation that hrp45 is an abundant protein in BR particles and that it is distributed along the RNP fiber of unfolded BR particles (Wurtz et al. 1996). Thus, hrp45 appears to be frequently bound along the BR transcript. As the transcript is dominated by exon 4, representing 80–90% of the entire transcript, it can be concluded that hrp45 binds extensively along this exceptionally large exon. The immuno-EM technique used here is not sensitive enough to establish whether hrp45 is also bound to the other, smaller exons and to the four introns. Antibody labeling of growing RNPs on both the most promoter-proximal and the most promoter-distal parts of the BR genes suggests, however, that hrp45 is likely to be present along the entire transcript. Such a result is in agreement with the observation that both SR proteins (Roth et al. 1991; Wu et al. 1991) and hnRNP proteins (Wu et al. 1991) are distributed along the entire loops in lampbrush chromosomes.

As an SR protein, hrp45 is likely to participate in the splicing process. The observed distribution of hrp45 along the BR transcript indicates, however, that most, if not all, hrp45 is not a constitutive component of spliceosomes. The appearance of spliceosomes along the active BR genes was studied recently by immuno-EM using antibodies to snRNP particles (Kiseleva et al. 1994). The antibody labeling was found to be restricted to the proximal third of the gene and to the very distal end of the gene; this distribution corresponds to the regions where the introns are located. It was concluded from this study that both the assembly and the disassembly of the spliceosomes are closely linked to the splicing process, the three 5’ end introns known to be spliced in the proximal third of the gene and the 3’ end intron at least to some extent spliced before the completed transcript is released.
from the template (Baurén and Wieslander 1994). Thus, the more or less even distribution of hrp45 along the transcript does not correspond to that expected for a spliceosome component.

It is conceivable that hrp45 is being added to the transcript immediately upon transcription and is present along the primary transcript when spliceosomes are assembled on pre-mRNA. Such a view is supported by recent studies that show that SR proteins promote the assembly of the earliest detectable splicing complexes (Staknis and Reed 1994) and, in fact, commit pre-mRNA to the splicing pathway (Fu 1993). Furthermore, it has been shown that SF2/ASF presumably bound to pre-mRNA, helps recruiting U1 snRNP to the 5' splice site (Kohtz et al. 1994; Jamison et al. 1995). The SR proteins are also involved in several of the subsequent steps in spliceosome assembly and in the splicing process (for review, see Fu 1995).

In this context, it should be emphasized that SR proteins are known to be associated with specific exon sequences, the splicing enhancers [Lavigueur et al. 1993; Sun et al. 1993b; Tian and Maniatis 1993; Staknis and Reed 1994], which stimulate splicing of the upstream intron (Reed and Maniatis 1986; Watakabe et al. 1991; Xu et al. 1993). The interaction between the SR proteins and the enhancer sequence can be direct (Lavigueur et al. 1993; Sun et al. 1993b) or through complex formation with specific binding factors (Tian and Maniatis 1994; Lynch and Maniatis 1995). The bound SR proteins promote splicing by interacting with the splicing factor U2AF, stimulating and/or stabilizing complexes assembled at the nearby 3' splice site [Wang and Manley 1995]. As SR proteins can interact with each other and with other splicing factors (for review, see Fu 1995), it has been proposed that they exert their effect by forming a network of proteins between the enhancer sequence and the adjacent upstream 3' splice site (Staknis and Reed 1994). Whether or not the BR exon 4 has enhancer activity has not been established, but it is interesting to note that the exon has a high purine content (45% adenine and 20% guanine in the repeat unit of exon 4 of the BR2.1 transcript) [Sfimegi et al. 1992], which is known to be a characteristic of most exonic enhancers [Lavigueur et al. 1993; Sun et al. 1993a; Xu et al. 1993; Tanaka et al. 1994; Lynch and Maniatis 1995]. The remarkable observation in the present study is that hrp45 resides along the entire exon 4 rather than at a specific site close to the upstream intron. The simplest explanation would be that because the exon sequence is internally repetitive (Wieslander 1994), a binding site (and a putative enhancer) would be present in a large number of copies evenly distributed along the exon. It is less likely that the exonic hrp45 molecules specifically mediate a stimulatory effect of a downstream 5' splice site enhancer (Robberson et al. 1990; Talerico and Berget 1990; Hoffman and Grabowski 1992; Staknis and Reed 1994), because in the case of BR exon 4 splicing of the upstream intron 3 is completed far ahead of the synthesis of the downstream intron 4 (Baurén and Wieslander 1994; Kiseleva et al. 1994). However, it is not excluded that the SR proteins bound along exon 4 could play a role in exon definition, a process crucial for proper splicing (Robberson et al. 1990). Finally, it should be noted that an hnRNP A1-like protein, Ct-hrp36, also appears along exon 4 (Visa et al. 1996), which could be functionally relevant as SR proteins and hnRNP A1 are known to affect at least alternative splicing in an antagonistic manner (Mayeda and Krainer 1992; Mayeda et al. 1994; Cáceres et al. 1994).

hrp45 remains bound to the BR transcript from the gene to the nuclear pore

Similar to other SR proteins [Fu and Maniatis 1990; Roth et al. 1990; Specter et al. 1991], hrp45 is essentially confined to the cell nucleus; the antibody labeling of cytoplasm is not above background. Our immunocytological analysis showed that hrp45 is present not only on the chromosomes but also in the nucleoplasm. Furthermore, in the immuno-EM analysis of the BR particles it could be demonstrated that hrp45 is associated with the particles in the nucleoplasm and in transit through the nuclear pores (Fig. 7). It has been shown earlier by immuno-EM of isolated, unfolded BR particles that hrp45 is distributed along the BR transcript also in nucleoplasmic particles (Kiseleva et al. 1994), almost all of them containing fully spliced RNA (Baurén and Wieslander 1994).

Although it is clear that the hrp45 protein is present in BR particles at or within the nuclear pore complex, it is more difficult to decide whether all the hrp45 protein molecules remain associated with the RNA during the transport from the gene to the pore. It is difficult to compare accurately the antibody labeling of the clustered growing RNP particles with that of the solitary ones in the nucleoplasm and in the pore, although it is evident that the labeling is of the same order of magnitude. It is, therefore, unlikely that hrp45 has been removed substantially from the BR RNP complex during the intranuclear transport. Furthermore, as hrp45 is an abundant protein component in the BR particle [Wurtz et al. 1996], a major loss of hrp45 should affect the structure and composition of the BR particle. Comparisons of three-dimensional reconstructions of growing BR particles on the genes, released particles in the nucleoplasm, and particles translocating through the pores did not indicate any such substantial losses of material from the particles inside the nucleus [Skoglund et al. 1986; Mehlin et al. 1992, 1995]. Finally, and most important, it was shown by isopycnic centrifugation in CsCl gradients that essentially all nucleoplasmic BR particles have the same, well-defined protein to RNA ratio, indicating that there is no gradual loss of proteins from the particles [Wurtz et al. 1990]. Therefore, we regard it as less likely that a considerable part of the hrp45 is being released in the nucleoplasm prior to the unfolding and disassembly of the particle at the pore complex (Fig. 7).

In conjunction with the translocation, hrp45 is being shed from the particle. It is not possible to decide more precisely when the release takes place, but most likely, hrp45 does not pass through the central channel of the
nuclear pore complex. hrp45 would belong then to the nonshuttling group of hnRNP proteins [Piñol-Roma and Dreyfuss 1992; Dreyfuss et al. 1993].

Being distributed along the BR transcript and remaining associated with the transcript all the way to the nuclear pore, hrp45 shows the anticipated properties of a structural hnRNP protein and not those of a spliceosome component. This suggests that apart from its effects on splicing, other functions should be considered for hrp45. It is likely to participate in the packing of the BR RNA into a well defined higher-order RNP structure: a 7-nm RNP fiber [Lönnroth et al. 1992] tightly folded into an RNP ribbon that is being bent into a ring [Skoglund et al. 1986]. It is also conceivable that hrp45 could affect the properties of the BR particle in processes such as interactions with the nuclear matrix, intranuclear transport, binding to the nuclear pore complex, and translocation through the central channel of the pore complex. Two recent studies in mammalian cells also support a postsplicing role for SR proteins. First, it was demonstrated that some SR proteins stay bound to the splicing complexes through both steps of the splicing reaction and also remain in the exon–product complex [Blencowe et al. 1995]. Second, it was shown that overexpression of the SR protein SC35 can inhibit splicing and can result in accumulation of unspliced pre-mRNA that is transported efficiently into cytoplasm, indicating that SR proteins can affect the stability and/or facilitate transport of pre-mRNA [Wang and Manley 1995]. Thus, the hrp45 data, as well as other lines of evidence, suggest that SR proteins could be active not only in splicing but also in other processes within the cell nucleus.

Materials and methods

Experimental material

C. tentans larvae were raised under laboratory conditions [Lezzi et al. 1981]. C. tentans tissue culture cells were grown in suspension at 24°C as described by Wyss [1982].

Antibodies

The mouse monoclonal antibody 2E4 against C. tentans hrp45 was prepared in our laboratory as described [Wurtz et al. 1996] and used either as purified antibody or as 2E4-hybridoma supernatant as indicated. A monoclonal mouse anti-human von Willebrand factor antibody [DAKO-vWF, F8/86] was used as a primary antibody negative control for immunocytoology and immunostaining. The monoclonal antibody mAb104 [American Type Culture Collection No. CRL-2067] was used to detect SR proteins [Roth et al. 1990].

SDS–PAGE and Western blot analysis

Protein extracts for Western blot analysis were prepared and analyzed as described by Wurtz et al. [1996]. β-Glycerophosphate was added to all buffers at 5 mM to inhibit endogenous phosphatase activities. C. tentans tissue culture cells were homogenized in TNM buffer [10 mM triethanolamine·HCl (pH 7.0), 100 mM NaCl, and 1 mM MgCl₂ containing 0.2% NP-40 and 0.1 mM PMSF using a glass tissue grinder. The homogenate was centrifuged at 200g for 5 min at 0°C and the resulting supernatant constituted the cytoplasmic extract used for Western blot analysis. The pellet was washed with TNM, and RNP extracts I and II were prepared according to Wurtz et al. [1996]. The RNP extracts I and II were combined and used as nuclear fraction for Western blot analysis. In all cases, the proteins were precipitated with acetone, separated by electrophoresis in SDS–polyacrylamide gels, and transferred to PVDF membranes (Millipore) using a semidyed electrophoretic system [BioRad Labs]. The membranes were cut into strips, blocked with 10% nonfat dry milk in PBS for 1 hr, and incubated with the first antibody [hybridoma supernatant from clone 104 diluted 1/100 or 2E4 diluted 1/1000] in PBS containing 1% dry milk and 0.05% Tween-20. Labeling was visualized with alkaline phosphatase-conjugated anti-mouse immunoglobulins using the NBT/BCIP system [Promega].

Immunoprecipitation and alkaline phosphatase treatment

Immunoprecipitation of hrp45 from RNP extract I was performed essentially as described by Wurtz et al. [1996] using mAb 2E4 directly cross-linked to protein G–agarose [Zymed Laboratories, Inc]. After 1.5 hr of incubation at 4°C, the resin was sedimented and washed three times with PBS containing 0.1% NP-40. The washed resin was resuspended in 50 mM Tris-HCl [pH 9.0], 1 mM MgCl₂, 0.1 mM ZnCl₂, and 1 mM spermidine and divided into two equal parts. One of them was treated with 100 U/ml calf intestine alkaline phosphatase [Boehringer Mannheim] for 10 min at 37°C to dephosphorylate hrp45. The other half was incubated in the same conditions without enzyme [mock reaction]. In both cases, the hrp45 protein was finally eluted from the agarose resin with 0.5% SDS, precipitated with acetone, and analyzed by Western blot as described above.

Isolation of cDNA clones

Mouse mAb 2E4 was used at 1:1000 dilution to screen a random primed lambda gt11 cDNA library from salivary glands of C. tentans with Protoblott Immunoscreening System [Promega Biotech] according to the manufacturer's instructions. Purified positive plaques were used directly as a source of template DNA for PCR amplification with GeneAmp PCR Core Reagents [Perkin Elmer Cetus] and lambda gt11 forward and reverse primers [Promega Biotech]. The Magic PCR Prep DNA Purification System [Promega Biotech] was used for purification of DNA fragments amplified by PCR. PCR-amplified DNA fragments were nonradioactively labeled with the DIG-System of Nonradioactive Labeling and Detection [Boehringer Mannheim] and used for screening an oligo{dT}-primed lambda ZAP cDNA library from the salivary glands of C. tentans.

DNA sequencing and sequence analysis

To avoid subcloning, we PCR-amplified cDNA inserted into either lambda gt11 or lambda ZAP cDNA and used the amplified DNA as a source of DNA template for sequencing with walking primers. Nucleotide sequencing was carried out with Taq DyeDeoxy Terminator Cycle Sequencing Kit [Applied Biosystems] using double-stranded DNA templates. Multiple synthetic oligonucleotides (20-mers or longer) were used to sequence both DNA strands. The sequencing gel was run on a 373A Automated DNA Sequencer [Applied Biosystems]. Sequence analysis was performed with the University of Wisconsin Genetics Computer Group Sequence Analysis programs [Devereux et al. 1984] and EGGX extensions to the Wisconsin Package Sequence Analysis programs.

Expression of cDNA-encoded protein in bacteria

eDNA clone plhrp45.1 was used as a template in a PCR reaction with Taq polymerase [Boehringer Mannheim] to am-
pify the ORF. The oligonucleotides were designed to introduce a NeoI restriction site at the 5’ end of the amplified PCR product and a BamHI site at the 3’ end. The PCR product was digested with NeoI and BamHI and inserted into the corresponding sites of the expression vector pET21d [Novagen, Inc.]. The resulting plasmid pET21d-hrp45 was transformed into E. coli BL21(DE3)Es3 cells according to the protocol supplied by Novagen. Transformants resistant to both ampicillin and to chloramphenicol were selected and analyzed by sequencing. It was confirmed that the clone pET21d-hrp45-2 contains the cDNA insert with an ORF identical to the one in the original clone pHrP45.1. Therefore, clone pET21d-hrp45-2 was used further for protein expression. Bacterial cultures grown in Luria-Bertani medium with ampicillin and chloramphenicol were induced for 3 hr with 0.4 mM isopropylthiogalactoside (IPTG) when their OD600 reached 1. IPTG-induced and mock-induced supernatant or as a negative control 20 µg/ml anti-von Willebrand factor (DAKO). In the second step of immunostaining we used a 1:1000 dilution of mouse anti-human antibody (DAKO). The slides were washed three times for 5 min each in 0.1% Tween 20 in TKM and then incubated for 90 min with 40 µl of the primary antibody, diluted in TKM to 1:100 for 2E4, or 1:5 for the negative control anti-human von Willebrand factor antibody (DAKO). The slides were washed three times for 5 min each in 0.1% Tween 20 in TKM and then incubated for 90 min with the gold-conjugated secondary goat anti-mouse antibody [AuroProbe EM GAM IgG G5, Amersham, 5 nm in diameter]. The immunogold labeling was silver-enhanced with IntenSEM (Amersham).

Immuno-EM on isolated chromosomes

Transcription of BR genes was stimulated in vivo by adding pilocarpine nitrate to the cultivation water [0.1 mg/ml] and incubating larvae for 3–5 hr at 22°C as described previously [Visa et al. 1996]. Briefly, C. tentans salivary glands were fixed for 1 hr with 4% formaldehyde in 0.1 M cacodylate buffer [pH 7.2] at room temperature, cryoprotected with 2.3 M sucrose, and frozen by immersion in liquid nitrogen. Sections were obtained in a cryo-ultramicrotome Ultracut S/FC S (Reichert) and mounted on glass slides. Before immunolabeling, sections were blocked with 2% BSA in phosphate-buffered saline (PBS) [157 mM NaCl, 3 mM KCl, 8 mM Na2HPO4, and 2 mM NaH2PO4, pH 7.2] with 0.1 M glycine. As a first antibody, we used either undiluted 2E4-hybridoma supernatant or as a negative control 20 µg/ml anti-von Willebrand factor (DAKO). In the second step of immunostaining we used a gold-conjugated goat antibody against mouse IgG [AuroProbe EM GAM IgG G5, Amersham, 5 nm in diameter]. The immunogold labeling was silver-enhanced with IntenSEM (Amersham).

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