Molecular Cloning and Characterization of a Putative Nuclear DEAD Box RNA Helicase in the Spruce Budworm, Choristoneura fumiferana

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RNA helicases play important roles in cellular processes such as pre-mRNA splicing, RNA processing, ribosomal biogenesis, and translation. A full-length DEAD box RNA helicase cDNA (CfHL-113) was isolated from the spruce budworm, Choristoneura fumiferana. CfHL-113 contained the eight functional motifs, which are highly conserved in the DEAD box RNA helicase family, and an arginine-aspartate (RAD) domain at its N-terminal end. CfHL-113 was highly homologous to Rattus norvegicus HEL117 and human p68 genes, both of which are suggested to be involved in RNA splicing. The results of Northern and Western blotting showed that expression of the CfHL-113 gene was low or undetectable in eggs, larvae, pupae, and adults. High levels of expression were, however, detected in the three in vitro cultured cell lines, CF-203, CF-124T, and CF-70, which were developed from the midgut, ovaries, and mature larvae, respectively. Immunocytochemistry revealed that CfHL-113 protein was present exclusively in the nuclei of these cell lines. Arch. Insect Biochem. Physiol. 61:209–219, 2006. © 2006 Wiley-Liss, Inc.

KEYWORDS: RNA unwinding; RNA splicing; ribosome biogenesis; translation

INTRODUCTION

ATP-dependent RNA helicases are a family of proteins that are capable of unwinding double-stranded RNA and DNA/RNA hybrids (Lücking et al., 1998). Some members of this family have been experimentally demonstrated to play important roles in transcription, translation, mRNA splicing, and DNA replication (Lücking et al., 1998; de la Cruz et al., 1999). The most extensively studied samples include: eukaryotic initiation factor-4A (eIF-4A, Abramson et al., 1987; Dorn et al., 1993); pre-RNA processing proteins (pprp proteins, Dalbadie-McFarland and Abelson, 1990; Dayyeh et al., 2002); Drosophila vasa (Lasko and Ashburner, 1988); rat HEL117 (Sukeyegawa and Blobel, 1995); and human p68 (Ford et al., 1988; Hloch et al., 1990). This helicase family is characterized by a common core region that consists of eight highly conserved signature motifs (Lücking et al., 1998; Aubourg et al., 1999). Many members of this family have a DEAD (Asp-Glu-Ala-Asp) box in Motif IV. Therefore, they are called DEAD box RNA helicases (Schmid and Linder, 1992; Lücking et al., 1998).

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RNA helicases have been identified from almost all of organisms. In insects, several Drosophila RNA helicases have been well studied. Drosophila maleless protein (Mle) is an RNA helicase required for X chromosome dosage compensation and essential for male viability (Kuroda et al., 1991). A Drosophila homologue of the eukaryotic initiation factor 4A (eIF-4A) gene is maternally expressed in the oocyte; mutation of the gene is recessive lethal (Dorn et al., 1993). Drosophila ME31B is also a maternally expressed DEAD box RNA helicase and is strongly expressed during oogenesis (de Valoir et al., 1991). Drosophila hel protein is an enhancer of white variegation and is associated with chromosomes in cell nuclei of embryos and ovaries (Eberl et al., 1997). Two vasa genes were isolated from insects. Drosophila vasa gene is homologous to eIF-4A and its product is required in only the female germ line (Lasko and Ashburner, 1988) and is involved in the formation of polar granules and germ cells (Hay et al., 1988). The Bombyx mori vasa gene is the first reported lepidopteran RNA helicase and is expressed only in the germline (Nakao, 1999). The biological functions of most insect RNA helicases are not clear and need to be further investigated.

We have previously identified a 64-kDa DEAD box RNA helicase from the spruce budworm, Choristoneura fumiferana (Zhang et al., 2004), which is a homologue of human nucleolar RNA helicase NOH61 (Zirves et al., 2000) and yeast RNA helicase Dpb9p (Daugeron et al., 2001). We report here molecular cloning and characterization of another DEAD box RNA helicase cDNA (CfrH1c113) from C. fumiferana. CfrH1c113 was a homologue of rat HEL117 (Sukegawa and Blobel, 1995) and human prp5 (Will et al., 2002) genes, which are involved in RNA splicing.

MATERIALS AND METHODS

Experimental Insects

East spruce budworm (Choristoneura fumiferana Clem., Lepidoptera: Tortricidae) eggs were maintained at 22°C and 70% relative humidity (RH) and allowed to hatch into first instar larvae on the balsam fir needles. The first instar larvae molted into second instar in 6 days. The second instar larvae were maintained at 16°C for one week and then the diapausing second instar were stored at 2°C for 27 weeks to satisfy the obligatory requirement for cold treatment. At the end of this period, the larvae were moved from 2°C to 16°C for 1 week and then placed on artificial diet (McMorrin, 1965) at 22°C, 70% RH and a photoperiod of 12-h light and 12-h darkness and reared until they reached the adult stage.

Cell Lines

Three spruce budworm cell lines, FPMI-CF-203, IPRI-CF-124T, and IPRI-CF-70, were used in this study. FPMI-CF-203 was developed from the midgut tissues (CF-203, Sohi et al., 1993); IPRI-CF-124T was derived from neonate larvae (CF-124T, Billmoria and Sohi, 1977), and IPRI-CF-70 was from ovary (CF-70, Caputo Guido, personal communication). CF203 cells were grown in 25-ml flasks with SF900 medium supplemented with 5% FBS (Life Technologies, Inc., Gaithersburg, MD). Cells of CF-124T and CF-70 were grown in modified Grace’s media (Grace, 1962).

Construction and Screening of cDNA Library

Messenger RNA was isolated from the midgut cell line CF-203, which is responsive to juvenile hormone (Feng et al., 1999). A cDNA library was constructed in UniZAP XR vector (Stratagene, La Jolla, CA) according to the manufacturer’s instructions. Polyclonal antibodies raised against Rst(1)JH protein (formally called MEI; Ashok et al., 1998; Pursley et al., 2000) of D. melanogaster were used to screen the cDNA library for Rst(1)JH homolog in the spruce budworm. Positive plaques were selected after three rounds of screenings and plasmid cDNA was isolated.

Sequence Analysis

Sequencing was performed using AlFexpress™ AutoRead™ sequencing Kit and AlFexpress™ DNA
sequencer (Amersham Pharmacia Biotech, Piscataway, NJ). Annotation, comparison, and alignment of sequences were performed using the National Center for Biotechnology Information BLAST search services (Altschul et al., 1990) and Clustal Alignment Program (Higgins and Sharp, 1988) of DNASTAR (DNASTAR, Inc., Madison, WI). Prediction of the subcellular localization of protein was conducted using PSORT program (Nakai and Kanehisa, 1992; http://psort.nibb.ac.jp/).

Production of Recombinant Protein in Baculovirus Expression System

The open reading frame of CfrHlc113 was cloned into the pFastBac1 donor plasmid (Invitrogen, Burlington, Ontario, Canada). The generated recombinant plasmid was used to transform DH10Bac cells, in which the CfrHlc113 cDNA was transposited to Bacmid DNA (modified Autographa california multicapsid nucleopolyhedrovirus DNA, AcMNPV). Insertion of CfrHlc113 cDNA into the AcMNPV genome was confirmed by using CfrHlc113-specific primers and PCR according to the manufacturer's instruction. Sf21 cells were transfected with the recombinant AcMNPV. Expression of CfrHlc113 was examined using SDS-PAGE and Western blotting.

SDS-PAGE

Proteins were denatured at 100°C for 5 min in an equal volume of 2× protein loading buffer (0.1 M Tris buffer, pH 6.8, 4% SDS, 0.2% β-mercaptoethanol, 40% glycerol and 0.002% bromphenol blue) and were then separated in 8% acrylamide gels in Tris-glycine-SDS buffer (10 mM Tris, 50 mM glycine, 0.1% SDS, pH 8.0) in a mini vertical electrophoresis system (Bio-Rad Laboratories, Hercules, CA). Thirty micrograms of protein was used per lane. The gels were stained with Coomassie Blue R-250.

Antibody Production

The recombinant protein expressed in an AcMNPV recombinant was excised from SDS-PAGE gels. Polyclonal antiserum was made in a rabbit by Cedarlane Inc. (Hornby, Ontario, Canada). Antiserum was collected after three boost injections, each with 200 µg protein in the Freund’s adjuvant. Pre-immune serum collected from the same rabbit prior to immunization was used as a control.

Western Blotting

Proteins were transferred from SDS-PAGE gels to nitrocellulose membranes after electrophoresis. Thirty micrograms of protein was used per lane. The anti-CfrHlc113 antibodies were used as primary antibodies at 1:1,000 dilution. The goat anti-rabbit IgG alkaline phosphatase conjugate (Sigma Chemical Company, St. Louis, MO) was used as secondary antibody at 1:2,000 dilutions. Color development of the alkaline phosphatase reaction was performed using 5-bromo-4-chloro-3-indolyl phosphate and nitroblue tetrazolium as substrates.

Immunofluorescence Localization

Immunofluorescence localization of CfrHlc113 was performed according to the general procedures described by Watkins (1996) for insect cells. Cells were grown on glass slides and fixed with 2% paraformaldehyde fixative plus 0.1% Triton X-100 on ice for 30 min. Cells were then permeabilized in cold methanol for 5 min. Antibodies at 1:1,000 dilution were applied to the cells on the slide and incubated for 1 h. The cells were blocked for 1 h in 1.5% bovine serum albumin and were then washed four times in phosphate buffered saline (PBS), followed by treatment with fluorescein conjugated anti-rabbit IgG (Sigma-Aldrich Canada Ltd. Oakville, ON, Canada) (1:500 dilution) for 1 h. The cells were washed three times in PBS and counter-stained with 4′,6-diamidino-2′-phenylindole dihydrochloride (DAPI) for 30 min and examined by fluorescence microscopy. The photographs were taken as double exposures using fluorescence and DAPI filters.
Northern Blotting

Ten micrograms of total RNA per lane were separated on 1% formaldehyde-agarose gels. The RNA was visualized by staining with ethidium bromide and photographed under UV light. The RNA was then transferred to nylon membranes. The blots were pre-hybridized in a pre-hybridization solution (Rapid-hyb buffer, Amersham Pharmacia Biotech) for 4 h and then hybridized in a hybridization solution (pre-hybridization solution plus \(^{32}\)P-dCTP labeled CfrHlc113 DNA probe) at 65°C for at least 12 h. After hybridization, the membranes were washed twice in 2× SSC plus 0.1% SDS at 42°C for 15 min and twice with 0.5× SSC plus 0.1% SDS at 55°C for 15 min and once with 0.1× SSC plus 0.1% SDS at 65°C for 15 min.

RESULTS

Cloning and Sequence of CfrHlc113

In an attempt to clone Drosophila Rst(1)JH protein homologue, which was formally called MET and has been suggested to be a JH receptor (Ashok et al., 1998; Pursley et al., 2000), from the spruce budworm, we used the anti-Rst(1)JH protein antibodies to screen an expression cDNA library of the spruce budworm Cf-203 cell line, which was responsive to JH I (Feng et al., 1999). Ten immunologically positive clones were isolated from a total of approximately \(10^6\) plaques. Sequencing and comparison with the sequences in the GenBank database revealed that three of the ten positive clones were identical and encoded an already identified C. fumiferana member (64 kDa) of the DEAD box RNA helicases family (CfrHlc64, Zhang et al., 2004). Four other clones were identical and encoded a different 113-kDa RNA helicase that also belongs to the DEAD box family of RNA helicases (this report). Another clone encoded a nuclear exportin (unpublished data) and the remaining two clones encoded two unknown proteins (unpublished data). The present report describes the C. fumiferana 113-kDa RNA helicase, CfrHlc113. The CfrHlc113 cDNA was 3,196 base pairs in length. The longest open reading frame was 3,039 nucleotides in length, potentially encoding a protein of 1,012 amino acids with a calculated molecular mass of 113 kDa and a pI of 9.58 (GenBank accession number for the sequence is AY559246). All of the eight conserved functional motifs that define the RNA helicase family were found in CfrHlc113. A tetrapeptide Asp-Glu-Ala-Asp (DEAD) was present in the sequence, indicating that it was a member of the DEAD box RNA helicase family.

Comparison of the deduced amino acid sequence of CfrHlc113 with 43 RNA helicases from different species revealed that CfrHlc113 showed high identities to homologues of Mus musculus (70%, NP_666087.1), Anopheles gambiae (70%, XP_311375.1), D. melanogaster (64%, NP_573020.2), Rattus norvegicus HEL117 (57%, Sukegawa andBlobel, 1995. NP_620798.1), and Homo sapiens prp5 (57%, Will et al., 2002. AAH12304.1). CfrHlc113 also showed 18–46% identity to homologues from other species ranging from fungus to plants. In general, these RNA helicases can be clustered into three broad phylogenetic groups (Fig. 1A). Group I includes CfrHlc113, the well-characterized R. norvegicus HEL117 (Sukegawa and Blobel, 1995; NP_620798.1), and human prp5 (Will et al., 2002; AAH12304.1). It has been suggested that these two proteins are involved in mRNA splicing. CfrHlc113 also showed 30% identity to human p68 (Ford et al., 1988; NP_004387), and 27% identity to D. melanogaster vasa (Lasko and Ashburner, 1988; CAA31405) and Bombyx mori vasa-like gene (Nakao, 1999; BAA19572), which are believed to be involved in early embryogenesis and oogenesis. Functions of most other members in this group have not been reported. Group II includes bacterial RNA helicases (NP_755783, Q8XA87 and NP_457662) from E. coli and Buchnera aphidicola, which have been suggested to be cold-shock helicases (Jones et al., 1996). The well-characterized transcription factor DmElf4A (Dorn et al., 1993; CAA48790), DmME31B (De Valoir et al., 1991; AAA28603), and Dmhel (Eberl et al., 1997; AAB65835) are also clustered into this group. Group III is represented by CfrHlc64 (Zhang et al.,
Fig. 1. A: Phylogenetic analysis of 44 RNA helicases from different species: CfrHlc113 (C. fumiferana, this study, AY559246), CfrHlc64 (C. fumiferana, Zhang et al., 2004; AY460342), AgHlc-2 (A. gambiæ, XP_311375.1), DmHlc-2 (D. melanogaster, Lasko and Ashburner, 1998; NP_573020.2), RnHeli117 (R. norvegicus, Sukegawa and Blobel, 1995; NP_620798.1), Hsp90p (H. sapiens, Will et al., 2002; NP_12304.1), MmHlc-2 (M. musculus; NP_660887.1), CefHlc-2 (Caenorhabditis elegans, NP_500063.1), ChHlc (C. briggsae, CAE70203.1), OsHlc (Oryza sativa, NP_013959.1), AtHlc-2 (Arabidopsis thaliana, NP_173516.1), MgHlc (Magnaporthe grisea, EAA50614.1), NcHlc (Neospora canina, XP_331895.1), SpHlc (Schizosaccharomyces pombe, NP_587586.1), Hsp68 (H. sapiens, Ford et al., 1988; NP_004387), Dnvasa (D. melanogaster, Lasko and Ashburner, 1988; CAA31405), Bnvasa (Bombyx mori, Nakao, 1999; BAA19572), PyyHlc (Plasmodium yoelii yoelii, EAA17238.1), PfHlc (P. falciparum, NP_703432.1), Dnabstrakt (D. melanogaster, AA04040), BtHlc (Bacteroides thetaiotaomicron, NP_810798), EcHlc (Escherichia coli cold-shock helicase, NP_755783), SeHlc (Salmonella enterica, NP_457662).

EcsdA (E. coli cold-shock helicase A, Q8X87), PfHlc (Photorhabdus luminescens, NP_013687), Ba(Ap)Hlc [Buchnera aphidicolæ str. Aps (Acyrthosiphon pisum), NP_240190], Ba(Sg)Hlc: [B. aphidicolæ str. Sg (Schizzius graminum), NP_660702], HiHlc (Haemophilus influenzae, NP_438403), CacHlc (Clostridium acetobutylicum, NP_349534), DmE14A (D. melanogaster, CAA48790), DmME31B (De Valoir et al., 1991; D. melanogaster, AAA28603) Dmhel (D. melanogaster, Eber et al., 1997; AAD65835), AgHlc (A. gambiæ, XP_309500), DmHlc (D. melanogaster, NP_523434), MmHlc (M. musculus; NP_080814), RnHeli117 (R. norvegicus, XP_214091), HsNOH61 (H. sapiens, NP_061955), CefHlc (C. elegans, NP_740966), CaHlc (Candida albicans, CA21924), ScDbp9p (Saccharomyces cerevisiae, NP_013378), SpHlc (S. pombe, NP_588531); AtHlc (A. thaliana, NP_195217). GfHlc (Giara lambia, EAA46394). B: Octapeptide repeats of the RSD domain in CfrHlc113. The residues that are identical to the HEL117 octapeptide are shaded. The consensus residues are given when at least three residues match in the five repeats.
The eight highly conserved helicase core motifs were found in all these sequences, but their C- and N-termini were variable. The members in Group I have an RSD domain, EK domain, and/or Q domain at either the C- or N-terminal ends (Aubourg et al., 1999). The members of Group II and III do not usually have extending domains at either end. There was an arginine-serine-aspartate-rich (RSD) domain at the N-terminus of CfHlc113 (Fig. 2A). In the RSD domain, 48 out of 79 amino acids residues (61%) were arginine-serine/aspartate and present as Arg-Ser (RS) or Ser-Arg (SR) or Arg-Asp (RD) dipeptides. Five repeats of an octapeptide motif, SRDRxRxR, were found in this domain (Fig. 1B). The repeats were conserved with that in R.

(A)

![Alignment of amino acid sequences of CfHlc113 and CfHlc64.](image)

(B)

<table>
<thead>
<tr>
<th>DEAD box RNA helicases</th>
<th>I</th>
<th>II</th>
<th>III</th>
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<th>VII</th>
<th>VIII</th>
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<td><strong>CfHlc113</strong></td>
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<td><strong>Consensus</strong></td>
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Fig. 2. A: Alignment of amino acid sequences of CfHlc113 and CfHlc64. The eight conserved motifs are boxed and numbered I to VIII. The RSD-domain and the core region are based on Aubourg et al. (1999). B: Alignment of the functional motifs of DEAD box RNA helicases with CfHlc113 and CfHlc64. The conserved amino acid sequences of the eight functional motifs are cited from Aubourg et al. (1999). The residues that match the consensus residues are bolded.
norvegicus HEL117 and other RSD domain proteins (Sukagawa and Blobel, 1995; Aubourg et al., 1999). CfrHlc113 did not have Asp-Glu-Arg-Lys (DERK) and Arg-Gly-Gly (RGG) domains or a consecutive Gly hinge, which are found in some RNA-binding proteins between the RSD domain and the RNA helicase core region (Binney et al., 1993; Gibson and Thompson, 1994).

Prediction of subcellular localization using Reinhardt's method (Reinhardt and Hubbard, 1998) revealed that CfrHlc113 was a nuclear protein (77% reliability). By using the k-NN Prediction (Horton and Nakai, 1997), the possibility for nuclear localization was 87%, much higher than the possibilities for cytoplasmic (8.7%) and cytoskeletal (4.3%) localizations, indicating the possible nuclear localization of CfrHlc113.

**Comparison of CfrHlc113 and CfrHlc64**

Two DEAD box helicases, CfrHlc113 (this report) and CfrHlc64 (Zhang et al., 2004; AY 460342), were isolated using antibodies against Rst(1)JH protein. They differ in size and belong to different groups in the structural and phylogenetic tree analysis (Fig. 1). Comparison of these two sequences (Table 1) revealed that overall identities were only 20 and 26% at the amino acid and nucleotide levels, respectively (Fig. 2A). All eight of the functional motifs that are common in RNA helicases were found in these two sequences. They were highly conserved with 68% identity in overall residues within these motifs (Fig. 2B). The major difference between these two sequences was that CfrHlc113 contained a longer N-terminus with an RSD domain and a longer C-terminal region, whereas CfrHlc64 did not have an RSD domain in its short N-terminus (Fig. 2A).

**Expression of Recombinant CfrHlc113 Protein in Baculovirus System**

In vitro expression of CfrHlc113 cDNA in a baculovirus system resulted in a recombinant protein with an apparent molecular mass of approximately 132 kDa in SDS-PAGE gel (Fig. 3), confirming that the open reading frame of the cDNA did encode a protein. The difference between apparent (132 kDa) and calculated (113 kDa) molecular mass values may be due to the high number of basic amino acids in the sequence and possible post-translational modification.

**Expression of CfrHlc113 in Tissues and Cell Lines**

Northern blotting analysis detected a 3.2-kb transcript of CfrHlc113 in six-day-old eggs, 1st instar larvae, 6th instar larvae, pupae, and adults, each at 2 days old (Fig. 4A). The embryos and 1st instar larvae had low levels of transcripts, while pupae appeared to have the highest levels. The fat body, midgut, and epidermis of 6th instar larvae had similar though low levels of the CfrHlc113 transcript. Interestingly, the three cell lines CF-203, CF-124T, and CF-70, which were derived from the midgut, neonate larvae, and ovary, respectively, con-

![Fig. 3. Recombinant protein of CfrHlc113 expressed in recombinant AcMNPV infected SF21 cells. The protein was isolated from SF21 cells 24 h post infection with AcMNPV+CfrHlc113 virus or the control virus AcMNPV. The protein was separated in a 8% SDS-PAGE gel and stained with Coomassie Blue B-250.](image)

**TABLE 1. Comparison of CfrHlc113 and CfrHlc64**

<table>
<thead>
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<th>Properties</th>
<th>CfrHlc113</th>
<th>CfrHlc64</th>
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</thead>
<tbody>
<tr>
<td>No. of nucleotides</td>
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tained much higher levels of the CfrHlc113 transcript than the in vivo tissues (Fig. 4A).

Western blotting analysis did not reveal detectable CfrHlc113 protein in six-day-old eggs, 1st instar larvae, 6th instar larvae, pupae, and adults, each at 2 days old (data not shown). However, high levels of CfrHlc113 protein were detected in all three cell lines tested in this study (Fig. 4B), which is in concurrence with the results of the Northern blotting analysis (Fig. 4A).

**Cellular Localization of CfrHlc113 in CF203 Cells**

Immunocytochemistry analyses showed that the CfrHlc113 protein was present exclusively in the nuclei of these three cell lines (Fig. 5). This observation confirmed its nuclear localization as predicted in the sequence analysis mentioned above.

**DISCUSSION**

CfrHlc113 was isolated by screening an expression cDNA library of the CF-203 cell line using the antibodies raised against Rst(1)JH protein (Pursley et al., 2000). It is not known if the Rst(1)JH antibodies contain non-specific antibodies to the helicase gene product or the Rst(1)JH antibody recognizes the helicase. However, only 11.6% overall amino acid identity was found between CfrHlc113 and the Rst(1)JH proteins. There is no evidence to indicate a structural or functional similarity between CfrHlc113 and the Rst(1)JH protein. Therefore, identification of this RNA helicase using the Rst(1)JH antibody may be spurious though fortuitous.

Interestingly, the same antibody identified the two RNA helicases, CfrHlc113 and CfrHlc64. Sequence comparison revealed that CfrHlc113 and

![Image](https://example.com/image.png)

**Fig. 4.** A: Northern blotting analysis of expression of CfrHlc113 mRNA in tissues and cell lines of *C. fumiferana*. Ten micrograms of total RNA hybridized with a 32P-labeled CfrHlc113 cDNA as a probe. The estimated size of CfrHlc113 mRNA was 3.2 kb. Ribosomal RNA stained with ethidium bromide indicates equivalent loading of total RNA. CF-203, CF-124T, and CF-70 were used for protein analysis after 4 days in culture. E: Six-day-old eggs; L1: two-day-old 1st instar larvae; P: two-day-old pupae; A: two-day-old adults; L6: two-day-old 6th instar larvae; FB: fat body; MG: midgut; EP: epidermis. B: Western blotting analysis of expression of CfrHlc113 in cell lines of *C. fumiferana*. CF203 cells were grown in 25-ml flasks with SF900 medium supplemented with 5% FBS. Cells of CF-124T and CF-70 were grown in modified Grace’s media. Proteins were extracted from the cells for protein analysis after 4 days in culture. M: molecular marker. Thirty micrograms of protein was loaded in each lane.
Fig. 5. Immunocytochemical localization of CfrHlc113 in CF-203, CF-124T, and CF-70 cells by immunofluorescence and DAPI staining. CF203 cells were grown in SF900 medium supplemented with 5% FBS; cells of CF-124T and CF-70 were grown in modified Grace’s media. A-D: CF-203; E-H: CF-70T; I-L: CF-124T. A, E, I: Controls stained with DAPI; B, F, J: controls stained with fluorescein; C, G, K: treated with anti-CfrHlc113 antibody and stained with DAPI; D, H, L: treated with anti-CfrHlc113 antibody and stained with fluorescein. The green fluorescence indicates the presence of CfrHlc113 protein and the blue indicates the presence of nuclear DNA. The primary antibody was rabbit anti-CfrHlc113 at a dilution of 1:1,000, and the secondary antibody was fluorescein-labeled sheep anti-rabbit IgG F(ab)2 fragment at a dilution of 1:500. The staining was examined using fluorescein and DAPI filters under an Olympus fluorescence microscope (Olympus BX50) and photographically documented. N: nuclei; C: cytoplasm. Scale bars = 60 μm.

CfrHlc64 displayed high similarity only in their helicase core regions, with the N- and C-terminal ends being dissimilar. For example, CfrHlc113 had an RSD domain in its N-terminal end, whereas CfrHlc64 did not. We have found that expression of CfrHlc64 was up-regulated by the edysone agonist tebufenozide (RH5992) (Zhang et al., 2004). However, expression of CfrHlc113 was not inducible by either 20E or RH5992 (data not shown). These features suggest that these two proteins may play different cellular functions.

In an attempt to predict its function, CfrHlc113 was aligned with other known RNA helicases. CfrHlc113 showed high identities to homologues in M. musculus (70%), A. gambiae (70%), and D. melanogaster (64%). However, the functions of these homologues have not yet been reported. CfrHlc113 was also 56% identical to rat HEL.117 (Sukegawa and Blobel, 1995) and human prp5p (Will et al., 2002). Rat HEL.117 also contains an RSD domain and a DERK region. It is co-located in the nucleus with splicing factor SC35 and may be involved in pre-mRNA splicing (Fu and Maniatis, 1990). Human prp5 is involved in pre-mRNA splicing and pre-spicosome assembly (Will et al., 2002) by enhancing an ATP-dependent structural change in the U2 small nuclear ribonucleoprotein (snRNP), facilitating the interaction of the snRNP with the complementary region of the pre-mRNA (O’Day et al., 1996; Dayeh et al., 2002).

From the above structural and sequence analyses, we hypothesize that CfrHlc113 is a pre-mRNA processing protein or splicing factor. The subcellular localization prediction based on sequence analysis and the observation that CfrHlc113 was localized in nuclei of the three cell lines also support this hypothesis. This finding is consistent with the reports on intranuclear localization of other
RS-domain RNA helicases such as HEL.117 (Fu and Maniatis, 1990; Sukagawa and Blobel, 1995).

One of the interesting findings in this study is that CfrHlc113 protein was undetectable in the stages and tissues of larvae tested using Western blotting analysis, whereas high levels of CfrHlc113 protein were detected in the three cell lines. Failure to detect the protein in larval tissues may be due to too low levels of expression to be detected by Western blotting. Another possibility is that the protein expression might have been restricted to very narrow developmental stages. High expression in the cell cultures suggests that CfrHlc113 may be necessary for in vitro growth and cell division. In addition, high expression in the cells was not dependent on the original tissues from which the cell lines were developed because all three cell lines yielded high expression levels. CfrHlc113 was also similar (30%) to the nuclear helicase, p68, that is found in dividing cells of many mammals and amphibians, but absent from quiescent cells (Ford et al., 1988). Understanding the mechanism behind the difference in CfrHlc113 expression between the in vivo tissues and the in vitro cells will help us to understand the biological roles of this RNA helicase.

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