Npw38, a novel nuclear protein possessing a WW domain capable of activating basal transcription

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ABSTRACT

We have found a novel cDNA encoding a 265 amino acid protein possessing a WW domain in our full-length cDNA bank. The WW domain was sandwiched between an acidic region and an acidic-basic amino acid repetitive region. In vitro transcription/translation of the cDNA produced a 38 kDa product that was also found in the cell lysate by western blot analysis. Thus this protein is named the nuclear protein containing a WW domain, Npw38. Immunofluorescence studies and expression of a fusion protein to a green fluorescent protein revealed that this protein is localized in the nucleus. Npw38 was shown to be capable of binding to a poly(rG) resin. Interestingly, the WW domain of Npw38 was found to function as a transcriptional activator in CHO cells using the GAL4 DNA-binding fusion system. Furthermore, the WW domains of human YAP and Pin1 were demonstrated to have a similar transcription-promoting activity. Combined mutation of the conserved first and second Trp residues and a hydrophobic triplet of TyrTyrTyr in the WW domain of Npw38 abolished the transcription-promoting activity, but single mutations of these sites did not. These results suggest that some WW domains potentially possess transcription-promoting activity in mammalian cells.

INTRODUCTION

We have been constructing a Homo•Protein cDNA Bank composed of human full-length cDNA clones (1). The full-length sequenced cDNA clones were subjected to functional analysis. To find novel proteins possessing a protein–protein interaction module in this bank, a computer-assisted search was performed. We found several novel cDNAs encoding proteins possessing modules such as WW domains, an SH2 (Src homology 2) domain, an SH3 domain, a proline-rich SH3 domain-interacting module or a cysteine-rich domain. In the present study, we focus on a cDNA clone encoding the WW domain-containing protein and its functional analysis. This protein is called the nuclear protein containing a WW domain, Npw38. During a search of proteins binding to the WW domain of Npw38 using a yeast two-hybrid system, we found that the WW domain of Npw38 has a transcription-promoting activity in the yeast cell. Furthermore, this WW domain and those of the other WW domain-possessing proteins such as human YAP and Pin1 were shown to activate basal transcription in CHO cells using the GAL4 DNA-binding fusion system. This study suggests the existence of a WW domain-mediated regulation of basal transcription in mammalian cells.
Figure 1. Structure of Npw38 cDNA. (A) The nucleotide sequence of Npw38 cDNA and the deduced amino acid sequence. The putative polyadenylation signal is underlined. (B) Schematic representation of the domains in Npw38 protein. (C) Sequence alignment among the WW domains. The identical amino acid residues are indicated by black boxes based on the sequence of Npw38. The position of the first amino acid and the accession number of each sequence in the protein or nucleotide database are indicated. Asterisks indicate the amino acid residues conserved among all sequences. (D) Alignment of repeat sequences in the acidic-basic amino acid repetitive region.

MATERIALS AND METHODS

cDNA cloning
cDNA libraries were prepared from mRNAs of human tissues and cell lines using a DNA–RNA chimeric oligo-capping method (1). All cDNA clones in the bank have already been partially or fully sequenced. Using a consensus sequence of the WW domain [LPxGWExxxxxGxx(Y/F)(Y/F)x(N/D)HxTxT(T/S)xWxxP, where x represents an arbitrary amino acid], a computer-assisted search was performed to identify the cDNA encoding WW domains from the bank. As a result, a novel cDNA clone, HP10345, encoding protein Npw38 was found. This cDNA originated from a human gastric adenocarcinoma cDNA library.

Sequencing

The sequencing reaction was performed using a dye primer cycle sequencing kit (Perkin-Elmer). The reaction samples were electrophoresed on a DNA sequencer (Perkin-Elmer ABI, model 377). The computer analysis was done using the program GENETYX-MAC (Software Development).

In vitro transcription/translation

The in vitro transcription/translation was carried out in a reaction mixture containing \([^{35}S]\)methionine (Amersham) using a TnT coupled reticulocyte lysate system (Promega) according to the manufacturer’s instructions. The translated product was analyzed by 10–20% SDS–PAGE.

Northern blot analysis

Multiple Tissue Northern Blot filters (Clontech) were hybridized with a \(^{32}P\)-labeled cDNA insert prepared from the Npw38 cDNA using a random primer labeling kit (Takara). After washing with 0.1× SSC, 0.05% SDS at 50°C, the blots were exposed to an X-ray film for 36 h at –80°C with an intensifying screen.
Localization analysis of GFP–Npw38 fusion protein

To produce an enhanced green fluorescent protein (EGFP) fusion protein with Npw38, the coding region of the Npw38 cDNA was amplified by PCR with primers containing either an EcoRI or a SalI site and subcloned into pEGFP-C1 (Clontech). COS7 cells were transfected with 0.5 µg of pEGFP-C1-Npw38 or pEGFP-C1 as a control according to the liposome-mediated method (23). After a 24 h culture, the medium was removed and the cells were washed three times with phosphate-buffered saline (PBS). The cells were fixed with 4% paraformaldehyde/PBS and viewed under a fluorescent microscope (Nikon).

Preparation of GST–Npw38 fusion protein

The entire coding region of the Npw38 cDNA was amplified by PCR using primers with an EcoRI or a SalI site. The PCR product was digested with restriction enzymes and subcloned into the EcoRI or SalI site of pGEX-5X-1 (Pharmacia). The constructed vector was introduced into Escherichia coli BL21 and expression was induced for 3 h by adding isopropyl-1-thio-ß-D-galactopyranoside to a final concentration of 0.2 mM. The proteins were affinity purified with glutathione-Sepharose beads and eluted with reduced glutathione as described in the manufacturer’s instruction manual (Pharmacia).

Antibody preparation

The anti-Npw38 antiserum was prepared in a rabbit using a bacterially expressed GST–Npw38 fusion protein. Anti-GST antibodies were removed from the serum by GST affinity chromatography using an NHS-activated column (Pharmacia) and the anti-Npw38 antibodies were purified using a GST–Npw38 affinity column. The absence of reactivity by the purified anti-Npw38 antibodies against GST was confirmed by western blot analysis.

Western blot analysis

HeLa cells were lysed in RIPA buffer (10 mM Tris–HCl, pH 7.4, 1% NP-40, 0.1% sodium deoxycholate, 0.1% SDS, 150 mM NaCl, 1 mM EDTA, 2 mM phenylmethanesulfonyl fluoride, 1 µg/ml aprotinin), separated by SDS–PAGE and transferred onto a PVDF membrane. The protein was probed with the anti-Npw38 antibody and visualized by the ECL method (Amersham).

Immunofluorescence staining

HeLa cells were grown on 4-chamber culture slides (Beckton Dickinson). The cells were briefly washed with PBS and fixed with 4% paraformaldehyde in PBS for 15 min at room temperature. The fixed cells were washed, permeabilized with 0.2% Triton X-100 in PBS, incubated with the anti-Npw38 antibody for 15 min, washed with PBS and stained with rhodamine-conjugated secondary antibody and DAPI (4’,6-diamidino-2-phenylindole) for 15 min.

Transactivation assay in yeast cells

The entire coding region of Npw38 or the coding region of the WW domain (position 39–87) was amplified by PCR with primers containing either an EcoRI or a SalI site and subcloned into the EcoRI or SalI site of vector pAS2-1 (Clontech) to express a GAL4 DNA-binding domain fusion gene. The resulting plasmids, pAS-WW and pAS-Npw38, were introduced into yeast strain Y190 and transformants were grown on Trp-deficient plates. The plasmid pAS-laminin supplied with the kit was used as the negative control. As the positive control, Y190 cells carrying pCL that expresses a complete GAL4 protein were grown on Leu-deficient plates. The plasmid pAS-laminin was used as the positive control. The cells were incubated at 30°C for 3 days and a β-galactosidase colony-lift assay was performed according to the manufacturer’s instructions (Clontech).

Transactivation assay in mammalian cells

A TransAct™ Assay kit (Clontech) was used to detect transcriptional activation in mammalian cells. An EST clone encoding a partial coding region of human YAP (GenBank accession no. N33982) and that encoding the entire coding region of human Pin1 (accession no. H18274) were obtained from Genome Systems, Inc. The coding regions containing the test sequence of the WW domain-bearing protein were amplified by PCR using primers with either an EcoRI or a SalI site. The obtained cDNA fragments were subcloned into vector pM to express a fusion protein to a GAL4 DNA-binding domain under the control of the SV40 promoter. The substitution of amino acid residues in the WW domain were carried out by using a Quick Change™ Site-Directed
Figure 3. Localization of Npw38. (A) Localization of GFP fusion protein. COS7 cells transfected with pEGFP-C1 (left) or pEGFP-C1-Npw38 (right) were visualized by fluorescence microscopy. (B) DAPI and immunofluorescence staining. HeLa cells fixed with paraformaldehyde were incubated with anti-Npw38 antibody and subsequently rhodamine-conjugated secondary antibody and DAPI. The same area was viewed.

Mutagenesis Kit (Stratagene). A pM-VP16 vector and an empty pM vector were used as the positive and negative controls, respectively. The reporter plasmid pG5CAT has a chloramphenicol acetyltransferase (CAT) gene preceded by five GAL4 binding sites and the E1b TATA box. The pM plasmid (0.5 µg) was introduced into CHO cells with pG5CAT (1 µg) and pSVGal (1 µg; Promega) by the lipofection method. The cells were harvested 24–36 h after transfection. Cell extracts were prepared and assayed for CAT activity using a CAT assay kit (Promega). The CAT activity was quantified using a BAS 1500 imaging analyzer (Fuji Film) and normalized to the β-galactosidase activity.

Nucleic acids binding assay

The Npw38 protein was prepared by in vitro transcription/translation. Nucleic acids binding was examined using 3 µl of the 35S-labeled translation product diluted in 500 µl of KHN buffer (150 mM KCl, 20 mM HEPES, pH 7.9, 0.01% NP-40). Twenty microliters of poly(rA), poly(rC), poly(rG) or poly(rU) agarose beads, calf thymus single-stranded DNA or double-stranded DNA cellulose beads (all beads purchased from Sigma) were then added and rotated for 20 min at room temperature. The beads were centrifuged and then washed five times with 1 ml of KHN buffer. Bound proteins were dissolved in the SDS sample buffer and analyzed by SDS–PAGE.

RESULTS

Primary structure of Npw38

The Npw38 cDNA has an open reading frame capable of encoding a 265 amino acid protein (Fig. 1A). A characteristic
Expression of Npw38 protein and mRNA

The cDNA was transcribed and translated in vitro in a rabbit reticulocyte lysate. As shown in Figure 2A, the translation product gave a main band of 38 kDa on SDS–PAGE that was larger than the 30.5 kDa expected from the open reading frame of the cDNA. Since a 38 kDa protein was also detected in a HeLa cell lysate by western blot analysis (Fig. 2B), the apparent molecular mass of Npw38 seems to be 38 kDa. This kind of increase in the apparent molecular mass has been observed in the case of various proteins possessing a highly charged region.

A major transcript of ~1.3 kb was detected by northern blot analysis in every human tissue tested. The expression of Npw38 mRNA was relatively high in heart, skeletal muscle, pancreas, spleen, thymus, prostate, ovary, small intestine and peripheral blood leukocytes (Fig. 2C). These results reveal that the transcripts are ubiquitously present in various tissues.

Localization of Npw38 in cells

To analyze the subcellular localization of Npw38, a GFP fusion-based method was used. COS7 cells were transfected with an expression plasmid coding for EGFP alone or for an EGFP–Npw38 fusion protein, and then localization of a transiently expressed product was determined using fluorescence microscopy. Cells transfected with a construct encoding EGFP alone displayed diffuse fluorescence throughout the cells. On the other hand, an intense nuclear fluorescent pattern was observed in the cells expressing EGFP–Npw38 (Fig. 3A).
Figure 5. Transactivation assay of the Npw38 deletion mutants. (A) Schematic representation of the Npw38 deletion mutants used in this study. Each fragment was fused to the GAL4 DNA-binding domain (1–147) in the pM plasmid. (B) CAT assays were performed as described in Figure 4B with extracts from CHO cells co-transfected with pM plasmids encoding fusion proteins. (C) The relative CAT activities normalized to β-galactosidase activity are given as the average values of three experiments. The CAT activity of CHO cells carrying pM is defined as 1.0 (control).

the manufacturer. Unexpectedly, autonomous activation of the reporter gene was observed on expression of the WW domain fusion gene (Fig. 4A) as well as a complete GAL4 construct, while not on that of a laminin construct. When the intact Npw38 was used as bait, only faint activation of the reporter gene was observed during long-term incubation, suggesting that some part of Npw38 represses the activity. These results imply that the WW domain of Npw38 can activate transcription directly or via association with the transcription machinery.

Next, activation of transcription by the WW domain in a mammalian system was examined using a reporter plasmid carrying five GAL4 binding sites followed by the adenovirus E1b TATA box upstream of the CAT gene. The cDNA fragment encoding the entire Npw38 or the WW domain was inserted into vector pM carrying the GAL4 DNA-binding domain gene under the transcriptional regulation of the SV40 early promoter. After the reporter plasmid and pM fusion vector were introduced into CHO cells, the CAT activity of the CHO cell lysate was measured. The GAL4–WW hybrid could activate transcription of the CAT gene ~50–70-fold compared with the GAL4 DNA-binding domain alone (Fig. 4B). In contrast, no activation of the reporter gene was observed when the GAL4–Npw38 hybrid expression vector was introduced into the cells.

To investigate which region in Npw38 represses the WW domain-mediated transactivation, we prepared mammalian expression vectors for fusion proteins between the GAL4 DNA-binding domain and various deletion mutants of Npw38 as described in Figure 5A. Only fusion proteins having a short region containing the WW domain (positions 1–84, 1–98, 25–98 and 43–98) could weakly activate transcription, whereas no activation was observed in other longer constructs (Fig. 5B and C). The activation seems to be repressed by both flanking regions of the WW domain and strongly by the short region of positions 84–98. These results suggest that some factor of the sequences surrounding the WW domain could be involved in the regulation of the transcriptional activity of Npw38.

Transcriptional activation by other WW domains

To examine whether the transcription-promoting activity was common to WW domains, other WW domains were tested by this assay. The WW domain-coding region of human YAP or human
transcription of the CA T gene performed. GAL4–PPWW and GAL4–Pin1WW could activate Pin1 was cloned into the pM vector and the CA T assay was compared with the GAL4 DNA-binding domain control (Fig. 6).

Furthermore, in contrast to Npw38, the entire region of Pin1 could also activate transcription.

Figure 7 shows that neither single substitution of the first Trp to Ala (W52A), the second Trp to Ala (W75A) nor the Pro to Gly (P78G) abolished the activity. Rather, the W52A mutant enhanced the activity ~2-fold compared with the wild-type WW domain. Double mutation (W52A, W75A) reduced the activity to 60% of the wild-type. Single and multiple mutations in a hydrophobic amino acid triplet (Y64A, Y65A, W66A) did not abolish the activity (data not shown about single mutation). The combination of mutations (W52A, Y64A, Y65A, W66A) reduced the activity to 20%. Furthermore, when all these residues were simultaneously substituted (W52A, Y64A, Y65A, W66A, W75A), the transcription-promoting activity was lost to the control level. These results imply that single mutation of the conserved amino acid residues did not affect the entire structure of the WW domain which may be involved in transactivation.

**Nucleic acid binding ability of Npw38**

Since the RGH repeat sequence in the acidic-basic amino acid repetitive region is similar to an RGG box sequence which is one of the RNA binding motifs (28), we examined the nucleic acid binding ability of Npw38 using a semi-specific assay which has been extensively used to study the RNA-binding properties of hnRNP and other proteins (15,29,30). When in vitro translated protein was incubated with nucleic acid resins, Npw38 bound preferentially to a poly(rG) sequence (Fig. 8), implying that Npw38 is an RNA-binding protein. The binding of Npw38 to a single-stranded or double-stranded DNA was not detected under the same conditions. The result that GST–Npw38 (position 81–265) is bound to poly(rG) (data not shown) suggests that the acidic-basic repetitive region is a putative RNA-binding site.

**DISCUSSION**

We have found a cDNA encoding a novel nuclear protein possessing a WW domain, Npw38, in the Homo•Protein cDNA bank. Interestingly, the WW domain of Npw38 was able to activate transcription in both yeast and mammalian cells in vivo, when the WW domain was recruited to the promoter region of a vector plasmid in a transactivation assay. However, intact Npw38 caused no activation because of the existence of the flanking inhibitory sequence. Thus the WW domain of Npw38 may ordinarily be masked and inactivated by a factor capable of associating with Npw38 or by the remaining part of Npw38 itself. Furthermore, we demonstrated that the WW domains of Y AP and Pin1 possess similar transcription-promoting activity. These results imply that the transcriptional machinery consists of a protein capable of interacting with these WW domains. One candidate for the interacting protein may be RNA polymerase II on which the C-terminal domain has been shown to bind to various WW domains (18). Alternatively, the interaction may be mediated by a transcriptional co-activator possessing a Pro-rich motif. At present, however, it remains unclear whether these interactions have a physiological meaning.

The mutation analysis of the amino acid residues in the WW domain of Npw38 revealed that any single mutation of the conserved sequence in the WW domain did not abolish the transactivation-promoting activity. These results are partly in agreement with the results obtained with in vitro systems such as far western blotting analysis or co-precipitation of GST–Y AP WW domain fusion protein and PY motif peptides. Chen et al. reported that a single mutation of the first Trp (W177F) or that in the hydrophobic triplet region (Y188F, F189Y) did not abolish...
Figure 7. Effects of substitution of the conserved amino acid residues in the WW domain on the transcription-promoting activity. (A) Mutant WW domains used in the assay. Selected conserved residues were substituted using site-directed mutagenesis of the WW domain of Npw38 subcloned in the pM vector. The underlining indicates the β-sheet strands expected from the solution structure of a YAP WW domain. (B) CAT assays were performed as described in Figure 4B. (C) The relative CAT activities normalized to β-galactosidase activity are given as the average values of three experiments. The CAT activity of CHO cells carrying pM is defined as 1.0 (control).

Figure 8. Binding of Npw38 to nucleic acids. The 35S-labeled product obtained by in vitro transcription/translation of Npw38 cDNA was incubated with agarose beads carrying various nucleic acids including poly(rA) (lane 2), poly(rG) (lane 3), poly(rU) (lane 4), poly(rC) (lane 5), single-stranded DNA (lane 6) and double-stranded DNA (lane 7). Bound proteins were analyzed by SDS–PAGE. An amount corresponding to 50% input of each assay was loaded on lane 1.

the binding ability to the PY peptide in solution, but mutation of the second Trp (W199F) or the Pro (P202A) did (31). The W177F mutant was bound at a slightly higher affinity than the wild-type on the filter. A similar increase in activity was observed in the corresponding W52A mutant of the Npw38 WW domain. No effect of mutation of the second Trp or the Pro on the activity in our case may be attributed to the differences in the WW domain and that of the assay systems. Our system observed an interaction between the Npw38 WW domain and an unknown ligand protein in the cell under physiological conditions. On the other hand, Chen et al. observed an interaction between the YAP WW domain and PY motif peptides on a filter or in solution. A discrepancy in the effect of the mutation was reported in an in vitro system as well. The Y188F mutant of the YAP WW domain could bind to the PY peptides on the filter but not in solution (31). The decrease in the activation by combined mutation of the conserved residues may result from a conformation change in the WW domain. According to the structure of the WW domain of YAP, the hydrophobic triplet is adjacent to the first and second Trp (16), so interactions between the sidechains of these residues may contribute to the stability of the domain structure.

The in vitro translated Npw38 was specifically bound to poly(rG) resin, suggesting that Npw38 is an RNA-binding protein. Some transcription-related factors such as WT1 (32,33) or heterogeneous nuclear ribonucleoprotein K (34,35) have been reported to bind RNAs and defined DNAs with high affinity. Since Npw38 itself could not bind to a single- or double-stranded DNA, it may bind to DNA via associated proteins and act as a transcriptional activator, if Npw38 acts as a transcriptional regulator. Alternatively, Npw38 may be involved in RNA processing such as splicing, poly(A) adenylation or capping via its RNA binding ability. Recently, McCracken et al. have proposed the concept of the ‘mRNA factory’ (36) in which the transcriptional, splicing (37,38) and cleavage–polyadenylation (39) machineries interact through contacts with the C-terminal domain of RNA polymerase II. Possibly, Npw38 may be a component of the ‘mRNA factory’. We are continuing the further characterization of Npw38 including the identification of binding proteins of Npw38 which will help elucidate its physiological role.
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