Title: MITOGEN-ACTIVATED PROTEIN KINASE PHOSPHATASE cDNAs AND THEIR BIOLOGICALLY ACTIVE EXPRESSION PRODUCTS

(57) Abstract

The invention relates to a novel mitogen-activated protein kinase phosphatase, MKP-2. The invention further relates to methods and means for preparing and to nucleic acids encoding this protein. The MKP-2 of the present invention is useful in the control of cell growth, differentiation and apoptosis.
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MITOGEN ACTIVATED PROTEIN KINASE PHOSPHATASE cDNAs AND THEIR BIOLOGICALLY ACTIVE EXPRESSION PRODUCTS

GOVERNMENT RIGHTS

This invention was made with government support under grant number ROI DK 45921 from the NIH. The government has certain rights in this invention.

FIELD OF THE INVENTION

This invention relates to mitogen activated protein kinase phosphatases and the nucleic acid sequences that encode them. The present invention comprises novel enzymes and the cDNA encoding them. The invention further provides nucleic acid hybridization probes, recombinant expression constructs capable of expressing the mitogen activated protein kinase phosphatases of the invention, homogeneous compositions of the disclosed mitogen activated protein kinase phosphatases, and antibodies against epitopes of each of the mitogen activated protein kinase phosphatases of the present invention.

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BACKGROUND OF THE INVENTION

Mitogen-activated protein kinases (MAP kinases) mediate multiple cellular pathways regulating growth (1) and differentiation (2, 3). In neuronal cells, MAP kinase activity mediates the actions of growth factors like EGF that stimulate cellular proliferation as well as factors like NGF that maintain neuronal survival and differentiation (4-6). Such ligand-activated signal transduction pathways involve activation of receptor tyrosine kinases which initiates a series of phosphorylation events that activate a cascade of serine/threonine kinases converging on the MAP kinase (also called extracellular signal regulated kinase (ERK)) isoforms, ERK1 and ERK2 (7-9).

Activation of MAP kinase involves specific phosphorylations on threonine and tyrosine residues within the Thr-Glu-Tyr motif (10) by MAP kinase kinase (MAP kinase and ERK kinase or MEK) (2, 11). Phosphorylation of both these residues is required for MAP kinase activation (11, 12). It has been suggested that the inactivation of MAP kinase is a critical event that regulates the physiological response to MAP kinase activation (13). This inactivation is mediated, in part, by dephosphorylation of MAP kinases by dual specificity phosphatases called MKPs.
(MAP kinase phosphatases) that dephosphorylate both the threonine and tyrosine residues phosphorylated by MEK (13-16). The activation of MAP kinase appears to be tightly regulated through the coordinate action of MEK and MKPs. By regulating the extent of MAP kinase activation, these MKPs may dictate the choice of differentiation or proliferation within a developing cell (17).

The prototype dual-specificity phosphatase, VH1, was identified in vaccinia and showed similarity to cdc25, a protein that controls cell entry into mitosis (18). VH1 homologues from human (PAC-1, CL100, and most recently B23), mouse [MKP-1 (3CH134 or erp)], and yeast (Yop51, MSG5) have also been isolated (19-24). All are dual-specificity phosphatases that specifically dephosphorylate MAP kinase in vitro (25) and in vivo (13, 15, 26). MKP-1 (also called 3CH134 or erp) was discovered as an immediate early gene whose rapid transcription and subsequent translation are suggested to provide a feed-back loop to terminate growth factor signals (13, 19, 26). Overexpression of mouse MKP-1 was shown to inhibit dramatically fibroblast proliferation suggesting that the inactivation of MAP kinase in vivo by MKP-1 has a profound negative effect on cellular proliferation (25, 26).

MAP kinase activation by growth factors has been extensively studied in PC12 cells (27). PC12 cells originate from a rat pheochromocytoma and retain many features of neural crest-derived cells, most notably the ability to undergo neuronal differentiation upon stimulation by NGF (28). Transfection with activated forms of the oncogenes ras, raf-1 and src into PC12 cells is sufficient for differentiation in the absence of NGF stimulation (6, 8, 29). As each of these genes has been shown to converge on MAP kinase activation, this implies that components of the MAP kinase cascade are required for neuronal differentiation. More recently it has been shown that the activation of MAP kinase kinase, MAPKK-1, is required and sufficient for PC12 cell differentiation (3). Despite our understanding of MAP kinase activation in neuronal differentiation, we know relatively little about MAP kinase inactivation.

Because of the important role of MAP kinases in controlling cell growth and differentiation, it is desirable to have molecular tools useful for inactivation of these enzymes. The present invention presents such tools.

**SUMMARY OF THE INVENTION**

An object of the present invention is the development of useful molecular tools for determining and controlling the role of MAP kinases in cell growth and differentiation. Neuronal cells are of particular interest. A further object of the
present invention is to provide novel nucleic acids and MAP kinase phosphatases useful in the study and development of drugs aimed at regulating cell growth, differentiation, metabolism, and tumor suppression.

Accordingly, the first aspect of the present invention comprises novel, active homogeneous MAP kinase phosphatase enzyme compositions and the cDNAs encoding them. These compounds are useful for developing hybridization probes and antibodies for assaying MAP kinase phosphatase activity in cells and tissues. Another aspect of the invention provides nucleic hybridization probes, epitopes and antibodies to MAP kinase phosphatase.

Yet another aspect of the invention provides recombinant expression constructs capable of expressing the MAP kinase phosphatases of the invention and homogeneous compositions of the disclosed MAP kinase phosphatases. In this manner, the present invention allows one to obtain large amounts of highly pure protein expression product without resort to the costlier and more time consuming methods involved in purifying the enzymes from tissues.

**BRIEF DESCRIPTION OF THE DRAWINGS**

The foregoing and other aspects and advantages of the invention will be apparent on consideration of the following detailed description and the accompanying drawings, wherein:

FIGURES 1A and 1B are a representation of the restriction map and sequence of the MKP-2 cDNA. A. The 4.8 kb MKP-2 cDNA was digested with various restriction enzymes and a schematic representation of some of these sites is shown. The following abbreviations are used - RI, EcoRI; Pst, PstI; A, Apal; H, HindIII; RV, EcoRV; Sma, SmaI; Bam, BamHI; 5’ UT, 5’ untranslated region; 3’ UT, 3’ untranslated region; ATG, translation start site; TAG, translation stop site. The coding region of MKP-2 is shown as a rectangular box with 2 different domains highlighted. The stippled box represents CH2 domains (cdc2 homology 2) (42) while the hatched box represents the catalytic domain. B. Nucleotide sequence and the encoded amino acid sequence of rat MKP-2 cDNA is shown. The translation start site is denoted as +1. The consensus catalytic site, the AU-sequence motifs in the 3’ untranslated region, and the putative polyadenylation signal are underlined. The 5’ and 3’ untranslated regions are depicted in lower case letters.

FIGURE 2 shows the amino acid homology between MKP-2, MKP-1, PAC-1, and B23. The amino acid sequences of rat MKP-2, mouse MKP-1, mouse PAC-1, and human B23 are aligned with each other and the areas of homology are shown
as shaded boxes. The catalytic domain is boxed. Dots represent spaces put in for alignment. The dark grey boxes represent the two CH2 domains present in all MKPs. Arrows correspond to the primers used in the RT/PCR screening strategy for cloning MKPs.

FIGURES 3A and 3B show the in vitro translation of MKP-1 and MKP-2 cDNAs. A) Autoradiogram of 35S-cysteine labeled products of a coupled in vitro transcription-translation reaction are shown using cDNAs encoding MKP-1 (lane 1) and MKP-2 (lane 2). B) Western blot analysis of PC12 cell lysates (100 μg total protein) using antisera directed to a MKP-1 peptide with significant identity to the C-terminal residues of MKP-2 (Santa Cruz Biotech). A 46 kD protein molecular weight marker is indicated.

FIGURES 4A and 4B show that MKP-2 contains phosphotyrosine phosphatase activity. A) Dephosphorylation of the synthetic peptide Raytide by bacterial lysates expressing a truncated MKP-2 protein (amino acids 163-393). U; untreated extracts from bacteria expressing MKP-2, V; vanadate-treated extracts, O; okadaic acid treated, M; microcystine treated extracts, and C; control extracts from bacteria expressing vector alone. The activity is represented as percent of maximal stimulation. B) Dephosphorylation of ERK2 by MKP-2 is shown. Activated ERK2 (pp42) (see Experimental Procedures) was incubated with phosphatase buffer alone (lane 1), bacterial lysates expressing vector alone (lane 2), bacterial lysates expressing truncated MKP-2 protein (lane 3), bacterial lysates expressing MKP-2, assayed in the presence of vanadate (lane 4). Phosphorylation was assayed by Western blotting with a phospho-tyrosine antibody as described.

FIGURE 5 shows the ability of MKP-2 to block MAP kinase-dependent gene transcription. PC12 cells were transfected with 6 μg of either RSV-globin (control) (white bars) or Raf BxB (constitutively active Raf) (grey bars) in the presence of 30 μg of vector alone (CMV), CMV-MKP-1 (MKP-1) or CMV-MKP-2 (MKP-2). In addition all cells received 3 μg of the reporter 5xGal4-E1B luciferase and 3 μg of Gal4-Elk-1. Activity is shown as light units/100 μg protein. Note the activities of cells transfected with RSV-globin and either MKP are below the limits of detection.

FIGURES 6A and 6B show the expression of MKPs in rat tissues. A filter containing 2 μg of Poly A + mRNA isolated from the indicated tissues (Clonetech Lab, Inc., Palo Alto, CA) was probed with a MKP-2 specific riboprobe (panel A), stripped, and re-probed with a MKP-1 specific riboprobe (panel B). The molecular weight markers are indicated to the right.
FIGURES 7A and 7B show the expression of MKPs in cell lines. 10 µg of total RNA was isolated from each of the cell lines indicated and probed with an MKP-2 specific riboprobe (panel A), stripped, and re-probed with a MKP-1 specific riboprobe (panel B). The migration of the ribosomal bands is indicated to the right. hMKP-2 refers to an MKP-2 specific transcript detected only in cells of human origin. Abbreviations for the cell lines include: RIN, rat insulinoma; GH3C1, rat pituitary tumor; AtT20, mouse pituitary tumor; W2, rat medullary thyroid carcinoma; LTK, mouse fibroblasts; PC12, rat adrenal medullary tumor; MiaPACa, human pancreatic carcinoma; MOLT48, human lymphoblasts; HepG2, human hepatoblastoma; SK-N-MC, human neuroblastoma.

FIGURES 8A, 8B, 8C, 8D, 8E and 8F show the localization of MKP-2 mRNA in the rat brain using in situ hybridization. Light field (A,C, and E) are thionin counterstains) and dark field (B,D, and F) photomicrographs showing representative distribution of MKP-2 mRNA (32x). DG: dentate gyrus of the hippocampus, Pir: piriform cortex, 3V: Third ventricle, and Sch: suprachiasmatic nucleus of the hypothalamus. Sense MKP-2 riboprobe did not hybridize (data not shown).

FIGURES 9A and 9B show the effect of serum and growth factors on MKP-1 and MKP-2 mRNA. A. PC12 cells were serum starved for 24h (U, untreated control) and were treated with medium containing 15% serum for the indicated times (left panel) or with NGF (100 ng/ml) or EGF (20 ng/ml) for the indicated times (right panel). 10 µg of total RNA from each treatment was analyzed by Northern blot analysis using an MKP-2 specific riboprobe, the filter was stripped and re-probed with MKP-1, stripped again and re-probed with a 18S ribosomal RNA probe. The results from each probe are shown. B. Quantitative representation of MKP-1 and -2 mRNA levels normalized to the 18S ribosomal RNA from an average of two independent experiments, one of which is shown in A. Presented values for all treatments represent fold induction compared to untreated cells (lane U) which was normalized as 1. The columns are aligned so as to be directly underneath the treatments indicated in A. Black boxes represent MKP mRNA levels in PC12 cells serum stimulated for the indicated times, grey boxes represent MKP mRNA levels from serum starved PC12 cells treated with NGF for the indicated times, and the white boxes represent MKP mRNA levels from serum starved PC12 cells treated with EGF for the indicated times.

FIGURES 10A and 10B represent the induction of MKPs in the hippocampus by Kainic acid. 10 µg of total hippocampal RNA was isolated from control rats (C)
or rats treated with 8 mg/kg Kainic acid for the indicated times (kindly provided by
Drs. James Douglass and Pastor Couceyro). A. Northern blot analysis was
performed with a MKP-2 specific riboprobe, the filter was stripped and probed with
MKP-1, stripped again, and re-probed with cyclophilin as an internal control. B. The
blot was quantitated and the results were normalized to cyclophilin and are shown
graphically. The black box is the amount of MKP-1 mRNA and the grey box
represents the levels of MKP-2 mRNA.

FIGURE 11 shows that transient transfection of MKP-1 and MKP-2 inhibits
neurite outgrowth. PC12 cells were transfected with the indicated combinations of
plasmids and where indicated were also treated with 100 ng/ml NGF for 2 days after
transfection. RasV12 transfected cells were also incubated for 2 days after
transfection. β-galactosidase assays were performed and blue cells (β-galactosidase
positive) were counted. The presented values are an average of four individual
counts from separate fields of two independent sets of transfections and are
represented as the percentage of blue cells that elaborated neurites.

FIGURES 12A and 12B show the identification of PC12 cells stably
transfected with MKP-1 or MKP-2 cDNAs. PC12 cells were transfected with CMV-
MKP1 or CMV-MKP2 respectively and selected for three weeks in the presence of
800 μg/ml G418. Neomycin resistant cells were clonally isolated and 10 μg of total
RNA was used for Northern blot analysis with an MKP-1 specific riboprobe (A) or an
MKP-2 specific riboprobe (B). Fold inductions over control untransfected cells are
indicated in A. To demonstrate equivalent amount of RNA in each lane, the 18S
ribosomal RNA bands are also shown (A and B). The endogenous sizes for MKP-1
(1.9 kb), MKP-2 (6 kb), and the exogenous MKP-2 transgene (2.4 kb) are indicated.

FIGURES 13A and 13B show ERK activation by growth factors, phorbol
esters, and hormones is reduced in MKP-1 and MKP-2 overexpressing cells. A.
PC12, MKP1.10, and MKP2.3 cells were serum starved for 24 hours and treated for
10 minutes with the indicated drugs. ERK-1 immune complex assays were
performed as described using MBP as a substrate. B. Quantitative representation
of ERK activity assays from an average of either three independent experiments
(PC12) or from two independent experiments (MKP1.10 and MKP2.3). Presented
values for all treatments represent fold induction compared with untreated control
PC12 cells which were normalized as 1.

FIGURES 14A and 14B show the comparison of JNK activity in PC12 versus
MKP2.3 cells. A. PC12 and MKP2.3 cells were serum starved for 24 hours and
treated for 15 minutes with the indicated drugs. JNK-1 immune complex assays were performed in duplicate as described using gst-c-jun as a substrate. Quantitative representation of JNK activity assays from an average of two independent experiments is shown. Presented values for all treatments represent fold induction compared with untreated control PC12 cells which were normalized as 1. B. PC12 and MKP2.3 cells were transiently transfected with or without MEKK in the presence of both Ga14-c-jun and 5XGa14-E1B-luciferase as indicated. Luciferase activity was determined and represents an average of three independent experiments.

FIGURES 15A and 15B shows the comparison of the kinetics of ERK activation by growth factors in PC12 and MKP2.3 cells. PC12 and MKP2.3 cells were serum starved for 24 hours and treated with NGF (A) or EGF (B) for the indicated times. ERK-1 immune complex assays were performed and quantitated as described. Values for each treatment represent fold induction compared to untreated PC12 cells which were normalized as 1.

FIGURE 16 shows the Elk-1 independent transcription of growth factors is reduced in MKP2.3 cells. PC12 and MKP2.3 cells were transiently transfected with Ga14-Elk-1 and 5XGa14-E1B-luciferase and were treated with the indicated drugs for 6 hours. Control represents transfected cells that were not subjected to any drug treatments. The absolute luciferase activity was then normalized to amount of protein within each lysate and the presented values are the mean of three independent experiments. Note that the basal luciferase activity in MKP2.3 control cells were undetectable.

FIGURES 17A, 17B and 17C show the regulation of MKP-1 and MKP-2 mRNA expression. PC12 (panel A), MKP1.10 (panel B), and MKP2.3 (panel C) cells were serum starved for 24 hours (control) or treated with the indicated drugs for 30 minutes prior to isolation of total RNA. 10 μg of total RNA from each treatment were analyzed by Northern blot analysis using a MKP-1 specific probe as indicated. The filter was stripped and re-probed with a MKP-2 specific riboprobe as indicated. To demonstrate equivalent amount of RNA in each lane, the 18S ribosomal RNA bands are also shown. The filter was scanned and quantitated using a Phosphorimager and the results represent a fold induction over the control untreated cells.

FIGURE 18 shows the regulation of fos mRNA expression. PC12 and MKP2.3 cells were serum starved for 24 hours (control) or treated with the indicated drugs for 30 minutes prior to RNA isolation. 10 μg of total RNA was analyzed by
Northern blot analysis using a fos riboprobe. To demonstrate equivalent amount of RNA in each lane, the 18S ribosomal RNA bands are also shown. As both filters were incubated with the same riboprobe preparation, the presented numbers represent fold induction over PC12 control cells, arbitrarily defined as unity. Figure 19 shows the regulation of stromelysin mRNA expression. PC12 and MKP2.3 cells were serum starved for 24 hours (control) or treated with the indicated drugs for 24 hours prior to RNA isolation. 10 μg of total RNA was analyzed by Northern blot analysis using a stromelysin riboprobe. The 18S ribosomal RNA bands are also shown to demonstrate equivalent amounts of RNA in each lane. The presented numbers (only indicated for agents that induced expression of stromelysin) represent fold induction over the control cells that were normalized as 1.

Figures 20A and 20B show the comparison of the rate of proliferation between PC12, MKP1.10, and MKP2.3 cells. PC12, MKP1.10, and MKP2.3 cells were seeded equally and analyzed for growth rates using BrdU incorporation as an index of proliferation. A. The growth rate was determined in cells growing in serum containing media for a period up to 6 days after plating. Each value is represented as the mean+/−standard error of six independent experiments. B. PC12, MKP1.10, and MKP2.3 cells were seeded at equal density and then serum starved for 2 days. Serum containing media was then added and the cells were analyzed at the indicated days. Again, each value is represented as the mean+/−standard error of six independent experiments.

Figure 21 shows the phase contrast micrographs of morphologically differentiated PC12, MKP1.10, and MKP2.3 cells. PC12, MKP1.10, and MKP2.3 cells were serum starved for 24 hours (control) or were treated with NGF for the indicated days before fixation. Magnification 70X.

Figure 22 is a representation of the corrected sequence of the MKP-2 DNA and amino acid sequence (see description of Figures 1A and 1B). Figure 23 shows the corrected amino acid homology between MKP-2, MKP-1, PAC-1, and B23 (see description of Figure 2).

Detailed Description of the Preferred Embodiments

In one aspect, the present invention provides a method of recovering MKP-2 protein in substantially pure form comprising the steps of removing the supernatant from unlysed cells that express MKP-2 protein, introducing the supernatant to an affinity matrix containing immobilized antibody capable of binding to MKP-2 protein,
permitting the MKP-2 protein to bind to the antibody of the matrix, washing the
matrix to remove unbound contaminants, and recovering the MKP-2 protein in
substantially pure form by eluting said MKP-2 protein from said matrix.

The term "substantially pure" indicates a protein or composition that is
essentially free of contaminants similar to the protein. In the present case, the
normal contaminants associated with rat MKP-2 protein predominately include rat
proteins. Thus, rat MKP-2 protein is substantially pure if it is free of rat proteins.
"Essentially free" is determined by weight. In general, a composition containing 70%
or more by weight rat MKP-2 protein and less than 30% of other rat proteins may be
considered substantially pure. Preferably, the composition will be at least 80% rat
MKP-2 protein, more preferably at least 90%, and most preferably at least 95% rat
MKP-2 protein. The presence of dissimilar components does not affect the
determination of purity, thus a composition containing 0.7 mg/mL rat MKP-2 protein
in PBS will still be considered substantially pure if it contains less than 0.3 mg/mL
other rat proteins. In addition, further purification utilizing a lectin or wheat germ
agglutinin column may be used before or after the antibody matrix step. Other
purification steps could include, for example, sizing chromatography, ion
chromatography, and gel electrophoresis. Further purification by velocity
sedimentation through sucrose gradients may be used.

In another aspect of the invention, nucleotide sequences are provided that
encode MKP-2 protein, as well as the use of such sequences or fragments thereof
in the production of recombinant MKP-2 protein, as hybridization probes, or for other
purposes. It will be appreciated that alternate nucleic acid forms, such as genomic
DNA, cDNA, and DNA prepared by partial or total chemical synthesis from
nucleotides, as well as DNA with deletions or mutations, are also within the
contemplation of the invention. Also provided are novel messenger RNA (mRNA)
sequences corresponding to these DNA sequences.

Figure 22 sets forth the nucleic acid and deduced amino acid sequence of
MKP-2. Figure 22 differs from Figure 1B by six nucleic acid additions and
corresponding amino acid changes which are boxed in Figure 22. Figure 23 also
contains the corrected amino acid sequence for MKP-2.

It will be appreciated that the invention includes nucleic acids having
substantial sequence homology with the nucleotide sequence shown in Figure 22 or
encoding proteins having substantial homology to the amino acid sequence shown
in Figure 22. Homology refers to sequence similarity between sequences and can
be determined by comparing a position in each sequence which may be aligned for purposes of comparison. When a position in the compared sequence is occupied by the same nucleotide base or amino acid, then the molecules are homologous at that position. A degree of homology between sequences is a function of the number of matching or homologous positions shared by the sequences.

The term "sequences having substantial sequence homology" means those nucleotide and amino acid sequences which have slight or inconsequential sequence variations from the sequences specifically set forth herein, i.e., the homologous nucleic acids function in substantially the same manner to produce substantially the same polypeptides as the actual sequences. The variations may be attributable to local mutations or structural modifications. It is expected that substitutions or alterations can be made in various regions of the nucleotide or amino acid sequence without affecting protein function, particularly if they lie outside the regions predicted to be of functional significance.

Proteins comprising an amino acid sequence which is 90% homologous with the amino acid sequence shown in Figure 22 may provide proteins having MKP-2 activity. The biological activity of MKP-2 is discussed in greater detail in Examples 1, 2, and 3 below, and is generally defined as the ability of the protein to inactivate MAP kinase.

Isolated nucleic acids encoding a protein having the biological activity of MKP-2 and having a sequence which differs from a nucleotide sequence shown in Figure 22 due to degeneracy in the genetic code are also within the scope of the invention. Such nucleic acids encode functionally equivalent proteins (e.g., a protein having MKP-2 activity) but differ in sequence from the sequence of Figure 22 due to degeneracy in the genetic code. For example, a number of amino acids are designated by more than one triplet. Codons that specify the same amino acid, or synonyms (for example, CAU and CAC are synonyms of histidine) may occur due to degeneracy in the genetic code. As one example, DNA sequence polymorphisms within the nucleotide sequence of an MKP-2 protein (especially those within the third base of a codon) may result in "silent" mutations in the DNA which do not affect the amino acid encoded. However, it is expected that DNA sequence polymorphisms that do lead to changes in the amino acid sequences of an MKP-2 protein will exist within a population. It will be appreciated by one skilled in the art that these variations in one or more nucleotides (up to about 3-4% of the nucleotides) of the nucleic acids encoding proteins having the biological activity of MKP-2 may exist.
among individuals within a population due to natural allelic variation. Any and all such nucleotide variations and resulting amino acid polymorphisms are within the scope of the invention. Furthermore, there may be one or more isoforms or related, cross-reacting family members of MKP-2 described herein. Such isoforms or family members are defined as proteins related in biological activity and amino acid sequence to MKP-2, but encoded by genes at different loci.

In another aspect, the invention further includes a method for producing an antibody which is capable of binding to MKP-2 protein or DNA comprising the steps of preparing a peptide-protein or nucleotide-protein conjugate, said conjugate comprising at least 10, more preferably at least 14, and most preferably at least 18 consecutive amino acid or nucleic acid residues present in MKP-2 protein or DNA, immunizing an animal with said peptide-protein or nucleotide-protein conjugate, boosting the animals, and obtaining the antisera. In connection with this aspect, the present invention further includes monoclonal and polyclonal antibodies specific for MKP-2 protein and DNA, i.e., capable of binding to a MKP-2 protein or nucleic acid molecule, as well as hybridoma cell lines capable of producing such an antibody.

In other aspects, the invention includes the use of antibodies specifically directed to MKP-2 protein or nucleotides, such as to isolate MKP-2 protein from sources producing the protein, for purposes of determining the presence or amount of MKP-2 protein in a sample, and for other purposes apparent to those skilled in the art.

This invention further includes a method of diagnosis of the presence and location of an MKP-2 protein expressing cell using labelled nucleotide probe sequences or labeled antibodies of the invention.

The use of rat MKP-2 DNA, or fragments thereof, as a probe in the isolation, purification, and study of other MKP-2 proteins from other organisms is contemplated. Oligonucleotide fragments of MKP-2 DNA can also be used as primers to amplify (with specific DNA polymerase) genomic DNA, isolated, for example, from bacteria, fungi, avian, and mammalian sources. The amplified genomic DNA can then be analyzed to identify sequence variation/abnormality using the polymerase chain reaction assay (Saiki et al., Science 230:1350 (1985). See also, Mullis, K.B., U.S. Patent No. 4,683,202, July 28, 1987; and Mullis, K.B., U.S. Patent No. 4,683,195, July 28, 1987).

For analysis of mRNA for MKP-2, or mRNA for related proteins, dot hybridization and Northern hybridization analyses can be used to characterize mRNA
encoding MKP-2 protein or MKP-2 protein-like molecules quantitatively and qualitatively. From these studies valuable information can be obtained about the number of different forms of MKP-2 genes and their expression in various cell types, e.g., bacteria, fungi, avian, and mammalian.

In the nucleic acid hybridization method according to the present invention, the nucleic acid probe is conjugated with a label, for example, a fluorophore, a radioisotope, a chemiluminescent compound, etc. In the most general case, the probe would be contacted with the sample and the presence of any hybridizable nucleic acid sequence would be detected by developing in the presence of a chromogenic enzyme substrate, detection of the fluorophore by epifluorescence, by autoradiography of the radioisotopically labeled probe or by chemiluminescence. The detection of hybridizable RNA sequences can be accomplished by Northern blot analysis, RNase protection assays, dot blot methodology or an in situ hybridization methodology. Methods of these last two techniques are described by Gillespie and Bresser, "mRNA Immobilization in Nal: Quick Blots," Biotechniques, 184-192, (Nov/Dec 1983) and Lawrence and Singer, "Intracellular Localization of Messenger RNAs for Cytoskeletal Proteins," Cell 45:407-415 (1986), respectively. The readout systems capable of being employed in these assays are numerous and non-limiting examples of such systems include fluorescent and colorimetric enzyme systems, radioisotopic labeling and detection and chemiluminescent systems.

Another aspect of the invention provides a nucleic acid which hybridizes under high or low stringency conditions to a nucleic acid which encodes a protein having all or a portion of an amino acid sequence shown in Figure 22. Appropriate stringency conditions which promote DNA hybridization, for example, 6.0 x sodium chloride/sodium citrate (SSC) at about 45°C, followed by a wash of 2.0 x SSC at 50°C are known to those skilled in the art or can be found in Current Protocols in Molecular Biology, John Wiley & Sons, N.Y. (1989), 6.3.1-6.3.6. For example, the salt concentration in the wash step can be selected from a low stringency of about 2.0 x SSC at 50°C to a high stringency of about 0.2 x SSC at 50°C. In addition, the temperature in the wash step can be increased from low stringency conditions at room temperature, about 22°C to high stringency conditions, at about 65°C.

The foregoing and other aspects of the invention may be better understood in connection with the following examples, which are presented for purposes of illustration and not by way of limitation.
EXAMPLE 1

**Materials.** Restriction and modification enzymes were purchased from New England Biolabs (Beverly, MA), Boehringer Mannheim (Indianapolis, IN), and Promega Biotech. (Madison, WI). Superscript reverse transcriptase was from Gibco BRL (Gaithersburg, MD), Taq DNA polymerase from Perkin-Elmer and Sequenase from U.S. Biochemical Corp. (Cleveland, OH). All enzymes were used according to the instructions from the manufacturer. [a-32P]dATP (3000 Ci/mm), 35S-dATP, 35S-UTP (1500 Ci/mm), [g-32P]-ATP (800 Ci/mm), 35S-Cysteine (1075 Ci/mm) and [a-32P]UTP (800 Ci/mm at 40 Ci/ml) were purchased from Dupont/New England Nuclear. Oligonucleotides were synthesized by a core facility at Oregon Health Sciences University. Antisera to MKP-1 was purchased from Santa Cruz Biotechnology Inc. An antibody to phospho-tyrosine (clone 4G10) was kindly provided by Brian Drucker (OHSU, Portland, OR) (30).

**RT/PCR Amplification.** One mg of total RNA from PC12 cells was used to generate first strand cDNA after an initial annealing reaction to 0.1 mg of random hexamers at 70 °C for 10 min. Following equilibration to ambient temperatures, a buffer containing 50 mM Tris-HCl pH 8.3, 75 mM KCl, 3 mM MgCl2, 10 mM DTT, 500 mM of each of four dNTPs, and 200 units/mg of Superscript reverse transcriptase (RT) was added and the mix was incubated at 37 °C for 1 hr. The reaction was terminated by placing the tubes on ice and the cDNA was recovered by ethanol precipitation. The pellet was washed with 70% ethanol and resuspended in 100 ml of 5 mM Tris, 0.5 mM EDTA mix. Five ml of this cDNA was used as a template for PCR amplification. Initially, two degenerate oligonucleotides were synthesized that generated a 204 bp cDNA fragment. The 5′ primer corresponded to the conserved WFNEAI sequence present in MKP-1 (26) and PAC-1 (23) (5′-TGG-TC(TC)-AA(TC)-GA(GA)-GC(GA(TC)-AT-3′)), while the 3′ primer corresponded to the conserved NFSFGM sequence present in MKP-1 (26) and PAC-1 (23) (5′-CAT-(AG)AA-(GATC)(GC)(AT)-(AG)AA-(AG)TT-3′) (Figure 2). Two additional oligonucleotides were made to confirm the novelty of MKP-2. The 5′ primer was a degenerate primer corresponding to the conserved YDQGQP sequence (5′-TA(TC)-GA(TC)-CA(AG)-GG(GATC)-GG(GATC)-CC-3′), while the 3′ primer was specific for MKP-2 (5′-ATGAAAGAAGGCGTGG-3′) corresponding to MKKRVR sequence (Figs. 1B, 2). This set of primers generated a 336 bp cDNA fragment corresponding to nucleotides 628-963 (Figure 1B). The PCR reaction consisted of 50 mM KCl, 1.5 mM MgCl₂, 0.2 mM dNTPs, 15 mM Tris-HCl, pH 8.4, and 0.5 mg of each primer pair.
PCR was allowed to proceed for 30 cycles. Each cycle consisted of 1 min at 94 °C, 1 min at 55 °C, and 1 min at 72 °C in a thermocycler (Perkin-Elmer Cetus). The PCR products were purified and subcloned into pBluescript (SK-) (Stratagene) using restriction enzymes engineered at the ends of each of the primers.

**Screening of the PC12 cDNA library and isolation of the full length clone.** The 336 bp PCR fragment generated by using the specific MKP-2 primer (described above) was labeled by random primed synthesis and used to screen a PC12 oligo dT primed cDNA library in lgt10 with 5 x 10⁵ individual recombinants that had been size selected prior to ligation for clones greater than 2 kb. The library was plated onto 20 LB plates and allowed to grow at 42 °C to a concentration of 50,000 recombinants/plate. The plaques were then transferred onto nitrocellulose filters in duplicate. The filters were soaked in prehybridization/hybridization buffer (6X SSC, 5X Denhardt's, 1% SDS, 0.01 M EDTA, 50% Formamide, 100 mg/ml denatured salmon sperm DNA) at 42 °C for 1-2 hrs with gentle agitation. Random primed probe was made as described in the Boehringer Mannheim kit. One to two million cpm/ml of boiled MKP-2 specific probe was added directly to the prehybridization/hybridization mix and hybridization was allowed to proceed at 42 °C for 24 hrs. The filters were washed in 2X SSC and 1% SDS for 2 hrs at 65 °C with frequent changes in the wash solution. The final wash was in 1X SSC and 1% SDS after which the filters were air dried and put on film. After the tertiary screen, three positive plaques were obtained. Phage DNA was isolated by standard methods and digested with EcoRI to release the insert. The inserts obtained were cloned into pBluescript (SK-) and subjected to restriction enzyme mapping and sequencing.

**Sequencing.** MKP-2 cDNA inserts obtained were sequenced on both strands by the method of Sanger (31) using sequenase reagents (U.S. Biochemical Corp.) according to the protocol suggested by the manufacturer. Multiple internal primers were made to allow sequencing of the complete 800 bp insert. The 4 kb insert was also sequenced using multiple internal primers on both strands through the termination codon and partially into the 3' untranslated region. Sequences proximal to the polyadenylation signal were also obtained, as shown (Figure 1B).

**Cell culture and drug treatments.** PC12-GR5 cells (courtesy of Rae Nishi, OHSU, Portland, OR) were grown at 5% CO₂ in Dulbecco's modified Eagle's medium (DMEM), containing 5% fetal calf serum, 10% horse serum, and L-glutamine. Prior to drug treatments, the cells were serum-starved for 24 h with DMEM containing no serum and treated with either 100 ng/ml NGF or 20 ng/ml EGF.
for the indicated times.

**RNA Isolation.** Total cellular RNA was isolated using RNAzol™B (Biotecx Lab, Inc.) according to the manufacturer's protocol. Briefly, cells were grown to 30 to 50% confluency in a 100 mm plate, rinsed with cold phosphate-buffered saline, and scraped into 1 ml of RNAzol B. After vortexing, 0.1 ml of chloroform was added and incubated on ice for 15 minutes. The suspension was centrifuged and the RNA was precipitated from the upper aqueous layer with an equal volume of isopropanol. After pelleting, the RNA was resuspended in water, quantitated at OD_{260}, and used directly for RT/PCR or Northern analysis.

**Northern blot analysis.** Ten μg of total RNA was electrophoresed through a 1.2% agarose formaldehyde gel and transferred onto Magna NT filter (MSI, Westboro, MA) using standard methodology in 6X SSC. Filters were prehybridized in three ml of hybridization buffer (5% SDS, 400 mM NaPO₄ pH 7.2, 1 mM EDTA, 1 mg/ml BSA, 50% formamide) at 65 °C for 4 hours in a rotating hybridization oven. 2-5 x 10⁷ cpm/ml of the antisense riboprobe was then directly added to the hybridization buffer and hybridization was allowed to proceed for 24 hrs. The next day, filters were initially washed in 1X SSC at room temperature for 15 min. and then washed in (0.05X SSC, 0.1% SDS, 5mM EDTA pH 8) at 70 °C for 3-4 hrs. Filters were autoradiographed at -70 °C on Kodak XAR-5 film using Dupont intensifying screens. Quantitations were performed using a Molecular Dynamics phosphorimager 445 SF and all signals were normalized to the 18S and cyclophilin signals respectively.

**Riboprobe Synthesis.** The 336 bp cDNA fragment generated by using the specific MKP-2 primer (described under RT/PCR amplification in Experimental Procedures) was subcloned in pBluescript (SK-) and was used to synthesize antisense riboprobes by linearizing the plasmid with Sal I that was engineered into the 5’ primer. Full length MKP-1 cDNA (kindly provided by Nicholas Tonks, Cold Spring Harbor Laboratory, Cold Spring Harbor, N.Y.) was subcloned into pBluescript (SK-) and then linearized with Bam HI to generate a 1.9 kb MKP-1 riboprobe. Rat cyclophilin (pSP65 1B15) and 18S ribosomal RNA (18S.pSP65) plasmids (kindly provided by James Douglass, Vollum Institute, Portland, OR) were linearized using Pst I and Hind III respectively to generate linear antisense riboprobes. Antisense riboprobes were synthesized as described (32). Briefly, 1 μg of template DNA was incubated in transcription reaction mix (40 mM Tris-HCl pH 7.5, 6 mM MgCl₂, 2 mM spermidine, 10 mM NaCl, 10 mM dithiothreitol, 20 units of Rnasin, 0.5 mM each of
rATP, GTP, and CTP, 12 μM rUTP, 50 μM [α-32P]UTP; 800 Ci/mmol, and 15 units of the appropriate RNA polymerase) at 37 °C for 60 min. The reaction was stopped by the addition of 2 units of RNase-free DNase and incubated at 37 °C for 15 min. 25 mM EDTA was then added, samples were phenol-chloroform extracted, and ethanol precipitated. Antisense riboprobes were resuspended in water at a concentration of 1-2 x 10^6 cpm/μl.

In situ Hybridization. Male Sprague-Dawley rats (200-300 g) were anesthetized and perfused with 1000 ml of 4% paraformaldehyde in borate buffer, pH 9.5 at 4 °C (fixation buffer). Brains were dissected and incubated in fixation buffer for 8 hrs and then incubated overnight in fixation buffer with 10% sucrose. Brains were sectioned serially into 5 series of 30 mm slices with a sliding microtome. Sections were prepared and hybridized as described (33). The 336 bp MKP-2 specific cDNA fragment was used to synthesize antisense and sense riboprobes. Sections were hybridized with 35S-labeled riboprobes (10^7 cpm/ml) in 66% formamide, 0.26 M NaCl, 1.32X Denhardt’s solution, 13.2 mM Tris pH 8.0, 1.32 mM EDTA, 13.2% dextran sulfate pH 8.0, at 65 °C for 24 hrs. Slides were washed in 4X SSC, digested with RNase A (20 mg/ml for 30 min at 37 °C), and then rinsed in a stringent wash containing 0.1X SSC at 65 °C for 30 min. Sections were dehydrated, dipped in NTB-2 emulsion (Kodak), and developed after 21 days. Light and dark field photomicrographs were taken with a Dialux 22 EB at 32X magnification.

In vitro transcription and translation reactions. Full length MKP-1 and MKP-2 cDNAs were used in a coupled in vitro transcription and translation reaction using TNT™ coupled reticulocyte lysate system (Promega Corp., Madison, WI) as per the manufacturer’s instructions. Briefly, 1 μg of circular DNA was incubated with 27.5 μl TNT rabbit reticulocyte lysate, 2 μl TNT reaction buffer, 1 μl T7 RNA polymerase, 1 μl 1 mM amino acid mix minus cysteine, 2 μl [35S]-cysteine (1075 Ci/mmol at 11 mCi/μl), and 1 μl RNAsin at 40 units/μl in a final volume of 50 μl. The reaction was incubated at 30 °C for 1-2 hrs. The synthesized proteins were separated by SDS-PAGE and analyzed by autoradiography.

PC12 transfection assays. PC12 cells were grown to approximately 60% confluence prior to transfection. For transient transfection experiments, 3 μg of 5xGal4-E1B luciferase and 3 μg of cytomegalovirus (CMV) Gal4-Elk1 transactivation domain (Roberson et al., manuscript in preparation) were used in combination with either 6 μg of Rous sarcoma virus (RSV) promoter coupled to globin (control) or constitutively active form of Raf kinase (Raf BxB); (34) and 30 μg of either pCDNA3
(Invitrogen, Inc.) containing the CMV promoter, CMV MKP-1, or CMV MKP-2. Cells were transfected by calcium phosphate-mediated DNA transfer as described (35). Cell lysates were prepared 20-24 hrs following transfection and luciferase activity was determined as described (36).

**Western blotting of PC12 proteins.** PC12 cells were lysed in 200 μl of a 1% NP-40 lysis buffer (25 mM Tris pH 7.4, 150 mM NaCl, 1 mM EDTA, 1.5 mM MgCl₂, 10% glycerol, 1 mM EGTA, 1 mM PMSF (phenylmethylsulfonyl fluoride), 10 μg/ml leupeptin, and 2 mM vanadate). Protein concentrations were determined by the method of Bradford (37). One hundred μg of total protein was resolved on a 12% SDS-polyacrylamide gel and transferred onto Immobilon P membrane. The membranes were probed with an MKP-1 antibody (diluted at 1:2000) (Santa Cruz Biotech., Inc.). A HRP-conjugated secondary antibody was used to allow detection of the appropriate bands using enhanced chemiluminescence (Amersham, U. K.).

**Bacterial Expression of MKP-2.** The catalytic domain of MKP-2 encoded within a carboxyl-terminal 690 bp fragment (C-MKP-2), extending from amino acids 163-393, was subcloned into the PET 23b vector (Novagen) using specific PCR oligonucleotides. This vector provides an amino-terminal epitope tag derived from the T7 capsid protein (T7 tag) that can be detected using specific antibodies (T7 antibody, Novagen). The frame of the resultant cDNA was confirmed by sequencing. This plasmid and the vector alone were used to transform BL21 bacteria (Novagen). A protein of the expected size (28 kD) was induced upon incubation with 0.4 mM Isopropyl-b-β-thiogalactopyranoside (IPTG) and was detected using anti-T7 capsid antibodies. Prior to phosphatase assays, bacterial extracts were prepared in lysis buffer (50mM Tris-HCl, pH 8.0; 2mM EDTA) and sonicated. Insoluble debris was pelleted and the supernatant assayed directly.

**ERK2 dephosphorylation assay.** Activated ERK2 was prepared by incubating 10 ng of recombinant ERK2 (kindly provided by Dr. Edwin Krebs, University of Washington, Seattle, Washington) with 0.1 mg active MAP kinase kinase (MEK) (Santa Cruz Biotechnology, Inc.) in 1X MEK buffer (25 mM Hepes, pH 7.5; 10 mM MgCl₂; 1 mM DTT; and 50 mM [γ³P]-g-ATP) at 30 °C for 30 min. The activation of ERK2 was confirmed by Western blot analysis using an antibody directed against phospho-tyrosine (30). Ten ng of activated ERK2 was incubated with 10 μg of bacterial lysates from MKP-2 expressing and non-expressing cells in 1X Phosphatase buffer (50 mM Tris-HCl, pH 7.5; 1 mM EDTA; 10 mM DTT) for 15 min at 30 °C. The reaction was stopped by the addition of an equal volume of 2X
Laemmli sample buffer and the samples were separated by 13% SDS-PAGE. The dephosphorylation of ERK2 was confirmed by Western blotting with the phospho-
yrosine antibody using enhanced chemiluminescence for detection of the signal.

Phospho-tyrosine phosphatase assay. The synthetic peptide Raytide
(Oncogene Sciences) was phosphorylated on a unique tyrosine using src tyrosine
kinase activity immunoprecipitated from C3H10T1/2 cells (kindly provided by Sally
Parsons, University of Virginia) as described (38). Bacterial extracts containing 10
to 60 mg of bacterial proteins were incubated at 30 oC for 30 min with labeled
peptide (10^4 cpm) in 100 ml of 1X phosphatase buffer (described above). Additional
reagents [10 mM vanadate, 20 mM microsystin-leucine-arginine (M-LR) and 1 mM
okadaic acid] were added without prior incubation. The reaction was terminated
by the addition of 0.75 ml of Stop solution (2 mM NaHPO_4, 90 mM sodium
pyrophosphate, 0.9 M HCl, 4% (v/v) Norit A). Following brief centrifugation, 400 ml
of the supernatant was added to 2.5 ml of scintillant and the released counts from
phosphatase activity were measured on a scintillation counter. All phosphatase
assays were performed in duplicate.

RESULTS

PC12 cells express multiple MKPs; identification and cloning of a novel
MKP. To identify potential MKPs that are expressed in PC12 cells, a screening
strategy involving RT/PCR amplification was employed. Alignment of the sequences
of the known members of the MKP family [human PAC-1, mouse PAC-1, mouse
3CH134 and VH1] showed areas of high sequence homology particularly
surrounding the catalytic core consensus site (HCXAGXXR, where X=any amino
acid) ((23) and Figure 2). Degenerate primers were designed to the conserved
amino acid sequence WFNEAI (5’ primer) and the conserved amino acid sequence
NFSPMG (3’ primer) surrounding the catalytic core site (Figure 2). RT/PCR on total
RNA from PC12 cells with these two degenerate primers revealed the expected 204
bp product that contained a representative population of MKPs expressed in PC12
cells (data not shown). This PCR product was gel purified, digested with restriction
enzymes engineered at the primer ends, and subcloned into the appropriate sites
in Bluescript. Five positive clones were obtained that were analyzed by sequencing.
One clone was identified as the rat homolog of MKP-1 or 3CH134 (19). The
remaining four clones were identical and showed some unique features in
comparison to the other known MKPs. To analyze this clone further, two additional
primers were designed. The 3’ primer was directed to a unique stretch in the cloned
novel sequence (MKKRVR) (Figure 2). The 5' degenerate primer was designed to another stretch of conserved sequence between the MKP family members (YDQGGGP), 5' to the sequence already obtained (Figure 2). Subsequent amplification using these two primers resulted in a 336 bp amplicon (data not shown). Sequencing this fragment confirmed the novelty of this cDNA sequence (Figures 1B, 2). These results demonstrate that PC12 cells express at least two potential MKPs, MKP-1 and the novel cDNA that we have called MKP-2.

These RT/PCR partial fragments of the novel MKP-2 cDNA were labeled and used as probes to screen an oligo dT-primed PC12 cDNA library in λgt10. This library contained 5 x 10^6 individual recombinants that had been size selected prior to ligation for clones greater than 2 kb. The three positive clones obtained with this screen were plaque-purified and the phage DNA obtained was digested with Eco R1 to release the insert. Upon digestion, two of the three positive clones each revealed two insert fragments of approximately 4 kb and 800 bp while the other clone contained a single 3kb insert fragment. The 3 kb insert was never successfully subcloned and therefore not analyzed further. The two contiguous insert fragments (4 kb and 800 bp) were cloned separately into Bluescript and characterized by restriction analysis (Figure 1A) and sequencing (Figure 1B). The 5' untranslated region and the first 135 amino acids of the novel MKP were contained within the 800 bp fragment, while the remaining 3' end of the clone was contained in the 4 kb fragment. The cDNA encoding MKP-2 contained at least 377 bp of 5' untranslated region and a translation start site with a 11/13 match with the Kozak consensus sequence (Figure 1B). The open reading frame extends 1182 bp and encodes a protein product of 393 amino acids with a predicted molecular weight of 42.6 kD and an isoelectric point of 7.86. The open reading frame was followed by an unusually large 3' untranslated region of greater than 3.0 kb. In comparison, the size of full length mouse MKP-1 (3CH134, erp) and its human homolog (CL100) is only 2.4 kb (19, 22, 26) similar to the recently cloned member of this family, B23, which is 2.5 kb (24). The large 3' untranslated region of MKP-2 contains several AU sequence motifs (Figure 1B) that are thought to regulate mRNA stability and are also present in MKP-1 (26).

The putative 393 amino acid MKP-2 protein shares 58.8% identity to the 367 amino acid MKP-1 protein, 62.3% identity to the 314 amino acid mPAC protein, and 33 % identity to the 397 amino acid human B23 protein (Figure 2). The similarities are greatest at the 3' end near the catalytic domain. In contrast, however, the 5' end
has significant sequence differences compared to the other members of this family. The N-terminal half of MKP-2 (amino acids 1-187) shares only 33% identity to MKP-1 while the C-terminal half (188-393) shows much greater homology (76%) with nearly 100% identity around the catalytic core (Figure 2). The complete coding region of MKP-2 was subcloned into Bluescript to generate a full length cDNA (MKP2FL-6) to allow transcription from the T7 promoter. In vitro transcription and translation of MKP-1 and MKP-2 cDNA revealed the expected 39.4 kD and 42.6 kD proteins respectively (Figure 3A). Antisera directed to an MKP-1 C-terminal peptide sequence (amino acids 348-366) detected proteins of sizes similar to MKP-1 and -2 from PC12 cell lysates (Figure 3B). This cross-reactivity is likely the result of the significant homology between MKP-1 and -2 in this region (Figure 2).

**The MKP-2 protein contains phosphotyrosine phosphatase activity.** A carboxy terminal fragment of MKP-2 (amino acids 163-393) comprising the entire catalytic domain was subcloned into the bacterial expression vector PET-23b and expressed in the *E. coli* strain BL21 (LysS). Induction of bacterially expressed truncated MKP-2 by IPTG was confirmed by Western blotting using an antibody directed to the T7 capsid epitope (data not shown). Bacterial extracts expressing the truncated MKP-2 fusion protein displayed phosphotyrosine phosphatase activity that was inhibited by vanadate, but not by okadaic acid or microcystine-LR, inhibitors of serine/threonine phosphatases (39, 40) (Figure 4A). These extracts, but not extracts expressing vector alone, could dephosphorylate activated ERK2 protein in vitro as monitored by phospho-tyrosine Western blotting (Figure 4B). This dephosphorylation was blocked by vanadate (Figure 4B).

**MKP-2 blocked MAP kinase-dependent activation of the Elk-1 transactivation domain.** To demonstrate biologically significant activity towards ERKs in vivo, the full-length cDNA encoding MKP-2 (MKP-2FL-6 cDNA) was subcloned into a CMV driven expression vector (CMV-MKP-2) and expressed in PC12 cells with a Gal4-Elk-1 fusion protein used to direct expression of a Gal4-luciferase reporter. Transcriptional activation by the Elk-1 transactivation domain requires activated MAP kinase (41). In these cells, a constitutively active Raf-1 mutant, Raf BxB, induced Elk-1 dependent transcription of the luciferase reporter by greater than 100-fold (Figure 5). Co-transfection with a MKP-1 expression vector under the control of a CMV promoter (CMV-MKP-1) blocked the activation by greater than 60% of maximally stimulated levels. Co-transfection with similar amounts of CMV-MKP-2 reduced expression by greater than 90% of maximal levels (Figure 5).
These studies demonstrate that MKP-2 can potently inhibit transcriptional activation that is dependent on MAP kinase. Unstimulated activation of Elk-1 was assayed in the presence of serum. Both MKPs inhibited this "basal" expression of luciferase activity to undetectable levels, presumably by blocking serum-induced MAP kinase activity.

**MKP-2 is expressed in most tissues and cell lines examined; distinct expression compared to MKP-1.** To determine the distribution of MKP-2 in various tissues, Northern blot analysis using a 336 bp MKP-2 specific riboprobe (described in Experimental Procedures) revealed expression in most tissues obtained from 10-12 week rats (Figure 6A). MKP-2 mRNA was detected in most tissues including brain, spleen, and testes with the highest expression in the heart and lung and lower expression in skeletal muscle and kidney. No MKP-2 expression was detected in the liver. The same blot was stripped and reprobed with a riboprobe made to the entire coding region of MKP-1 (Figure 6B). This probe did not cross react with the 6 kb MKP-2 transcript. A 2.4 kb mRNA corresponding to MKP-1 was detected in all tissues but testes. The highest expression of MKP-1 was seen in the lung, as previously reported (19, 26), and in the heart (similar to MKP-2). In the testes, MKP-2 is abundantly expressed but not MKP-1. In the liver, the inverse expression pattern is found. These opposite expression patterns in two different tissues suggests different physiological roles for the members of the MKP family. These results show a fairly abundant basal expression of MKP-1 and MKP-2 in most tissues with distinct tissue distribution patterns between MKP-1 and MKP-2. MKP-2, as expected from the 4.8 kb cDNA which was missing a portion of the 5' untranslated region, encodes an approximate 6 kb transcript distinct from the 2.4 kb MKP-1 mRNA detected in the same tissues.

RNA was also isolated from cell lines of different lineages and 10 mg of total RNA from various cell lines was analyzed by Northern analysis using the 336 bp MKP-2 specific riboprobe (Figure 7A). A single 6 kb MKP-2 transcript was detected in all rat and mouse derived cell lines of different lineages while a single ~4 kb MKP-2 transcript was detected in all cells of human origin. The same blot was stripped and reprobed with an MKP-1 riboprobe (Figure 7B). No MKP-1 was detected in cells of human origin. This may reflect the inability of the mouse MKP-1 riboprobe to hybridize across species to cells of human origin under the stringent conditions used. A rat MKP-1 probe was also used with similar results (data not shown).

**Distribution of MKP-2 in the CNS.** It has been shown that ERK1 mRNA is
expressed in all areas of the brain with the strongest expression in the hippocampus and piriform cortex while there was no overlapping expression of CL100 (human MKP-1) in those same areas (42). In order to determine the distribution of MKP-2 mRNA in the brain, we performed in situ hybridization analysis on rat brain sections with an 35S-antisense and sense MKP-2 riboprobe. MKP-2 appeared to be expressed in many areas of the brain with very strong staining of the hippocampus, piriform cortex, and the suprachiasmatic nucleus (Figure 8). The sense riboprobe did not show any specific staining (data not shown). These results suggest the co-localization of an ERK isoform, ERK1, and a MKP isoform, MKP-2, in specific areas of the brain where MKP-1 is not expressed.

Regulation of MKP-1 and MKP-2 in PC12 cells. MKPs have been shown to be immediate early genes and are transcriptionally regulated by a variety of agents. For example, MKP-1 mRNA appears to be induced by bombesin, EGF, TPA, cAMP, and FGF to different extents and with different kinetics suggesting the involvement of this phosphatase in various signaling pathways (26). A recent member of the MKP family, B23, has also been shown to be regulated by serum in human skin fibroblasts (24). In order to determine the involvement of MKP-2 in different neuronal signal transduction cascades in PC12 cells, Northern blot analysis was performed to determine the levels of MKP-1 and -2 mRNA following growth factor stimulation.

To determine whether MKP-2 mRNA is serum-inducible, total RNA was isolated from PC12 cells that were serum starved overnight and then stimulated with media containing 15% serum for various times. Ten mg of RNA from each treatment was analyzed by Northern analysis using an MKP-2 specific riboprobe as described above. Both MKP-1 and MKP-2 were expressed in unstimulated PC12 cells (Figures 7A, B and Figure 9). Serum stimulation caused a biphasic stimulation of MKP-2 with a gradual increase by 1 hr followed by a transient decrease and then a subsequent increase in expression of MKP-2 by 4 hr. The same blot was stripped and reprobed with MKP-1 and the MKP-1 transcript followed a similar but less robust stimulation (Figures 9A, B, left panel). All quantitations were normalized to the 18S ribosomal RNA as an internal control.

To address whether differences in the induction of distinct MKPs may account for the differences observed in the kinetics of MAP kinase activation seen following NGF and EGF treatment of PC12 cells (5), PC12 cells were serum-starved and treated with NGF or EGF for various times. Ten mg of total RNA from each of these
treatments was analyzed by Northern blot analysis for MKP-1 and -2 expression (Figure 9A, right panel). All quantitations were normalized to 18S ribosomal RNA which was used as an internal control for amount of RNA loaded (Figure 9B, right panel). NGF induced expression of MKP-1 and MKP-2 by 1 hr. Continued stimulation by NGF resulted in a maximum of 5-fold increase in MKP-2 expression by 2 hr and a 3-fold increase in MKP-1 expression by 1 hr. Both MKP-1 and -2 were maintained at slightly elevated levels upon NGF stimulation. 1 hr of EGF also induced expression of MKP-1 by 2.7-fold and MKP-2 by 4-fold. In contrast to NGF, this induction however was inhibited after 2 hr of EGF treatment in both cases. Additional incubation with EGF resulted in a second induction of both MKP-1 (2.6-fold) and MKP-2 (5.7-fold) by 4 hr. This bi-phasic kinetic pattern was also seen following stimulation with serum, with a minimum RNA level seen at 2-2.5 hr, as well (Figure 9B). These results show that growth factors that activate members of the MAP kinase cascade also result in the transcriptional activation of MKPs. The significance of the different kinetic pattern of MKP induction by NGF and EGF has yet to be determined.

Regulation of hippocampal expression. Due to the high expression of MKP-2 mRNA in the hippocampus (Figure 8), we analyzed MKP-2 expression after adult rats (250-300 g) had been subjected to the global seizure inducing drug, kainic acid (8 mg/kg). RNA from the hippocampus of rats subjected to kainic acid was extracted 0.5 and 1 hr after drug treatment and analyzed by Northern blot analysis (Figure 10A). Quantitations were normalized to the amount of cyclophilin in each case. MKP-2 mRNA was induced 5.1-fold by 1 hr of kainic acid treatment while MKP-1 mRNA was induced 3.2-fold (Figure 10B). This kainic acid induced transcriptional stimulation of MKPs shows the involvement of these genes in the stress-induced pathways in the brain.

DISCUSSION

We have identified a second widely-expressed MAP kinase phosphatase we term MKP-2. It is co-expressed with MKP-1 in a large number of tissues but also shows distinct differences. In contrast, the MKP PAC-1, is expressed only in lymphoid cells (23). Like MKP-1 and PAC-1, MKP-2 shows homology to other tyrosine phosphatases. Its sequence is most conserved with MKP-1 and PAC-1 within the C-terminus and less so within the N-terminus (Figure 2). All three share identity within the catalytic core (VHCQAGISR), and display phosphotyrosine phosphatase activity against synthetic peptides and purified MAP kinase. The dual
specificity of MKP-2 towards a phospho-threonine has yet to be proven. However, we have shown that both MKP-2 and MKP-1 block MAP kinase-dependent gene transcription in vivo, as been shown for PAC-1 (15). All four MKPs contain the CH2 domains (cdc25 homology 2) which are regions present in members of the cdc25 family that flank the catalytic domain (42). In MKPs, the catalytic domain is situated at the C-terminus and not at the N-terminus where the CH2 domains are found (Figure 2). Whether these CH2 domains are functional has yet to be proven but they have been speculated to either increase substrate selectivity or to be involved in localizing proteins to nuclear or cytoskeletal locations (42).

The large (>3 kb) 3' untranslated region in MKP-2 differs from the shorter (697 bp) 3' untranslated region present in MKP-1 and may play a regulatory role in post-transcriptional events such as transcript stability. Several AUUUA motifs are found in the 3' untranslated region of both MKP-1 (19, 26) and MKP-2 (Figure 1B). MKP-2 also has a 25 nucleotide stretch of AU sequences which might also contribute to posttranscriptional control. These AU sequence motifs have been implicated in the short half life (~10 min) of MKP-1 mRNA (19). Such runs of AU sequences occur in the 3' untranslated regions of lymphokines, cytokines and proto-oncogenes and are thought to be recognition signals for selective mRNA degradation (43). The role of these motifs in MKP-2 regulation has yet to be determined.

In contrast to the expression of PAC-1 which is limited to lymphoid cells (23), MKP-1 and MKP-2 show expression in a broad range of tissues with distinct differences. These distinct tissue distribution profiles may dictate unique roles of the members of this family in the regulation of MAP kinases and might reflect their co-expression with certain MAP kinase isoforms. MKP-1, ERK-1 (42) and MKP-2 mRNA are expressed in discrete areas of the brain. ERK2 expression is also prominent in neuronal cell bodies and dendrites particularly within the superficial layer of the neocortex, the hippocampal CA3 region, dentate gyrus, and cerebellar Purkinje cells (44). The co-localization of MKP-2 and the ERK isoforms in certain discrete areas of the brain suggests that ERK1 and ERK2 may be physiological substrates for MKP-2. However, MKP-1 and MKP-2 also show overlapping expression which suggests that co-expression of MKPs and ERKs is not the only criterion for substrate specificity.

The function of MAP kinase in post-mitotic neuronal cells is unclear. MAP kinases within the developing and adult central nervous system (45) are activated
by both neurotrophic growth factors and neurotransmitters. For example, activation of the N-methyl-D-aspartate (NMDA) receptor leads to increased tyrosine phosphorylation of an ERK isozyme in hippocampal cultures (46). Kainate is an NMDA receptor antagonist that induces seizures and immediate-early genes within the hippocampus (47, 48). The induction of CL100 (human MKP-1) and B23 mRNA by oxidative stress and heat shock has been reported (22, 24) and the induction of MKP-1 and -2 observed with kainic acid treatment may represent a regulatory role these genes might play in response to cellular stress. It is not known whether the stress activated kinases are substrates for the MKPs (49).

Basal expression of MKPs may be important for the resting cell. Vanadate stimulates MAP kinase activity in resting PC12 cells (data not shown). This suggests that a constitutive tyrosine phosphatase activity may inhibit basal MAP kinase activity. In addition, the introduction of dominant negative mutants of MKP-1 into unstimulated COS cells activates MAP kinases in the absence of serum (13). Therefore MKPs may be active in resting cells and might function to minimize the level of basal MAP kinase activity in the resting cell.

In PC12 cells, both NGF and EGF stimulate a receptor tyrosine kinase to phosphorylate and activate similar intracellular substrates including MAP kinase, whose action is required for both proliferative and differentiating responses (3, 4, 50). It has been suggested that the duration of MAP kinase activation determines the biological response to growth factor stimulation (5, 51). Proliferation is associated with a transient MAP kinase activation (5, 51, 52), while agents that induce a differentiating response produce a sustained activation. It is not known whether regulation of MKPs establishes the time courses of MAP kinase inactivation. We demonstrate that both NGF and EGF induced a rapid increase in MKP-1 and MKP-2 mRNA levels with a more substantial increase in MKP-2 compared to MKP-1. Whether this induction of MKPs is responsible for the rapid inactivation of MAP kinase by EGF is not known. A recent report demonstrates that inactivation of MAP kinase by EGF is independent of MKP-1 induction (53). Our results would agree with their findings as NGF, which sustains the level of active MAP kinase for a longer duration than EGF, also resulted in elevated levels of MKP-1 and -2 for at least 4 hr of drug exposure. Therefore, it is possible that the MKPs are regulated post-transcriptionally by these agents. The differences in transcriptional regulation of MKP-1 and MKP-2 by NGF and EGF suggest that they have different mechanisms of regulating MKP activity.
In conclusion, PC12 cells express at least two related MKPs, MKP-1 and MKP-2. The identification of a family of MKPs that are expressed within the same cell suggests distinct roles for each member of this expanding family. Discrimination of the actions of these MKPs may occur through the divergent amino-termini of these proteins. Further studies are required to identify the physiological roles of each member of this unique family of phosphatases in order to gain a better understanding of the mechanisms involved in cellular proliferation, differentiation, and stress.

**EXAMPLE 2**

Insights into physiological roles of MKP-1 and MKP-2 have come from transient transfection studies where the extracellular signal regulated kinases (ERKs) activation was blocked by MKP overexpression (60, 68, 70, 73). For example, transient expression of MKP-1 and MKP-2 into PC12 cells inhibits ERK-dependent pathways (68). However, transient transfection, which permits transcription from multiple copies of the exogenous plasmid DNA, results in levels of expression that generally exceed those reached during physiological induction of transcription. In contrast to the action of transiently transfected MKPs, physiological induction of endogenous MKP-1 and MKP-2 in PC12 cells following NGF treatment does not correlate with the inactivation of ERKs (68). These observations suggest that the specificity of MKP’s actions may depend upon their level of expression as well as other factors. Less robust expression can be achieved using stable expression of transfected genes which requires chromosomal integration and selection and therefore may mimic more closely the levels reached during physiological stimulations. In the study set forth below, we compared the effects of transient and stable expression of transfected MKP-1 and MKP-2 on signaling pathways initiated by extracellular stimuli that activate the ERK signal transduction cascade. We demonstrate that ERK activation and neuronal differentiation are differentially sensitive to MKP expression.

**Materials.** PC12-GR5 cells were kindly provided by Rae Nishi (Oregon Health Sciences, University, Portland, Oregon). The Ga14-Elk-1, 5XGa14-E1B-luciferase, and RasV12 plasmids have been described previously (68). The Ga14-c-jun plasmid was a kind of gift from Richard Goodman (Vollum Institute, Portland, Oregon). C-fos cDNA was kindly provided by Jim Douglas (Amgen, Thousand Oaks, California) and the stromelysin-1 cDNA was provided by Gary Ciment (Oregon Health Sciences University). Agarose-coupled antibodies to JNK1(FL) and ERK1(C-
16) were from Santa Cruz Biotechnology (Santa Cruz, California). NGF was from Boehringer Mannheim, EGF was from Sigma, Forskolin and PMA were from Calbiochem.

**Cell culture.** PC12-GR5 cells were grown at 5% CO₂ in Dulbecco’s modified Eagle’s medium (DMEM) containing 5% fetal calf serum, 10% horse serum, and L-glutamine. Prior to drug treatments, the cells were serum starved for 24 h with DMEM alone and subsequently treated with 100 ng/ml NGF, 50 ng/ml EGF, 10 μM Forskolin, or 100 nM PMA.

**Plasmids.** Full length MKP-1 cDNA (1.9 kb Hind III-Bam HI fragment) was subcloned into pcDNA3 (Invitrogen) under the cytomegalovirus (CMV) promoter to generate CMV-MKP1. A truncated MKP-2 cDNA fragment containing the entire coding region (2.4 kb Eco RV fragment) was also subcloned into pcDNA3 to generate CMV-MKP2 (68). pcDNA3 contains the neomycin gene driven by the SV40 promoter.

**Transient transfections.** 60-80% confluent cells were co-transfected using the standard calcium phosphate co-precipitation method (Gibco BRL) with the indicated combinations of the following plasmids: 10 μg of RSV-β-galactosidase, 20 μg of CMV-MKP1, 20 μg of CMV-MKP2, 10 μg of RasV12 or 5 μg of Ga15-Elk-1 and 5 μg of 5XGa14-E1B-luciferase. In order to determine the transcriptional activation of c-jun, cells were transfected with or without 1 μg of MEKK and 5 μg of both Ga14-c-jun and 5XGa14-E1B-luciferase as indicated. The parent vector pcDNA3 was added to each set of transfections to equalize the amount of DNA the cells received. Four hours following transfection, cells were glycerol shocked and allowed to recover in serum containing media overnight. Cells were then starved overnight in supplemented serum free media (N2) which contained DMEM with 5 μg/ml insulin, 100 μg/ml apo-transferrin, 30 μM Sodium selenite, 100 μM Putrescine, and 20 nM Progesterone. Cells were then treated with the indicated drugs for 6 hours prior to harvesting. Briefly, cells were washed twice in phosphate buffered saline (PBS), scraped in PBS, spun at low speed to collect cells, and lysed by freeze-thawing three times in 100 mM K₂PO₄, pH 7.8. The lysate was spun at high speed and the supernatant was assayed for luciferase activity using a luminometer (AutoLumat LB953).

**Histological detection of β-galactosidase.** The expression of β-galactosidase was used to identify transfectants within the population of differentiating cells. For counting blue cells (β-galactosidase positive) with neurites,
the transfected cells were exposed to NGF for 2 days prior to fixation. PC12 cells were fixed in 2% paraformaldehyde and 0.2% glutaraldehyde for 5 minutes after which cells were washed in PBS and subjected to a β-galactosidase assay. Cells were exposed to 2 mM MgCl₂, 5 mM Ferric cyanide, 5 mM Ferrous cyanide, and 0.1% X-gal in PBS overnight at 37°C. Transfected cells were identified as those staining and were then counted to determine the percent of blue cells with neurites in each set of transfections. Each set of transfections was done in duplicate and 200 cells were counted for each experimental condition.

**Stable transfections.** PC12-GR5 cells were seeded at 3 x 10⁵ cells per 100 mm plate 48 hours prior to transfection. Cells were transfected with 20 µg of CMV-MKP1 and 20 µg of CMV-MKP2 respectively by calcium phosphate co-precipitation and were exposed to the precipitate for 4 hours. The cells were then glycerol shocked and allowed to recover in complete media. 48 hours later, cells were split and plated in complete media containing 800 µg/ml G418. Stable neomycin-resistant cells were clonally isolated using cloning rings at 3 weeks post-transfection and maintained in media containing 600 µg/ml G418.

**RNA isolation, Riboprobe synthesis, and Northern blot analysis.** RNA isolation using RNAzol B and MKP-1 and MKP-2 riboprobe synthesis has been described elsewhere (68). The c-fos transcript was detected by linearizing the 1.3 kb pGEM-c-fos plasmid with Eco RI and using SP6 RNA polymerase for antisense RNA probe synthesis in the presence of α-³²P-UTP (40μCl/μl). Stromelysin transcripts were detected by linearizing pGEM-TR1 with Hind III and using T7 RNA polymerase to make antisense RNA transcripts. The conditions for Northern blotting using cRNA probes has been described (68). All filters were scanned and quantitated using a Molecular Dynamics PhosphorImager 445SI.

**Proliferation Assay.** Equal numbers of cells (2000/well) were seeded for all three cell lines (PC12, MKP1.10, and MKP2.3) on 96-well plates. Proliferation was assessed by using the Cell Proliferation ELISA, BrdU kit (Boehringer Mannheim). Briefly, cells were labeled with 10 µM BrdU for 4 hours after which they were fixed directly on the plate. Cells were then incubated with a BrdU-antibody, washed 3 times, and incubated with substrate for 10 minutes prior to addition of the 1 M H₂SO₄ stop dye. Results were quantized immediately on a ELISA reader at 450 nm. Each day represents an average of six independent wells for each of the three cell lines. Cells were also serum deprived by exposure to N2 media for 2 days prior to stimulation with serum containing media for the days indicated. Again, each day
represents an average of six independent wells for each cell line.

**ERK immune complex assay.** Treated and untreated cells were lysed in a lysis buffer containing 10% Sucrose, 1% NP-40, 20 mM TrisCl pH 8.0, 137 mM NaCl, 10% glycerol, 2 mM EDTA, 1 mM PMSF, 1 μg/ml Leupeptin, 1 mM Sodium Vanadate, and 10 mM Sodium Fluoride. The lysates were spun at low speed to remove nuclei and the supernatant was assayed for ERK activity. 100 μg of total protein (as determined by Bradford Assay) was immunoprecipitated with an agarose-coupled antibody to ERK-1 (C-16) overnight at 4°C. The immunoprecipitated ERK-1 was washed 3 times in lysis buffer and was assayed for kinase activity by incubating with 10 μg myelin basic protein (MBP) and 10 μCi γ-32P-ATP in 50 μl of buffer containing 80 mM Hepes pH 7.4, 80 mM MgCl2, 0.1 mM ATP, 2 mM Sodium Vanadate, and 20 mM Sodium Fluoride for 30 minutes at 30°C. Reactions were terminated by the addition of 50 μl of Laemmli sample buffer and analyzed by SDS-PAGE. Quantitations were performed by scanning the gel using a PhosphorImager.

**JNK immune complex assay.** Treated and untreated cells were lysed in a lysis buffer containing 20 mM Hepes-KOH pH 7.4, 2 mM EGTA, 50 mM β-glycerophosphate, 10% Glycerol, 1% Triton X-100, 1 mM DTT, 1 mM Sodium Vanadate, 0.4 mM PMSF, 0.5 μg/ml Aprotinin, and 0.5 μg/ml Leupeptin. The lysates were spun at low speed to remove nuclei and the supernatant was assayed for JNK activity. 100 μg of total protein (as determined by Bradford Assay) was immunoprecipitated with an agarose-coupled antibody to JNK-1(FL) overnight at 4°C. The immunoprecipitated JNK-1 was washed 3 times in each of 3 buffers (Lysis buffer, LiCl buffer (500 mM LiCl, 100 mM Tris-HCl pH 7.6, 1 mM DTT, and 0.1% Triton X-100), and Assay buffer (20 mM MOPS pH 7.2, 10 mM MgCl2, 2 mM EGTA, 1 mM DTT, and 0.1% Triton X-100)) and was assayed for kinase activity by incubating with 3 μg Gst-c-jun and 1 μCi γ-32P-ATP in assay buffer for 30 minutes at 30°C. Reactions were terminated by the addition of 50 μl of Laemmli sample buffer and analyzed by SDS-PAGE. Quantitations were performed by scanning the gel using a PhosphorImager.

**Morphological determination.** Cells were grown on Primaria plates and were serum starved in N2 media for 24 hours (control) and subsequently treated with NGF in N2 media for the days indicated. Cells were washed twice with PBS and fixed in 4% paraformaldehyde and 0.1% glutaraldehyde for 5 minutes after which they were washed in PBS. Cells were photographed at 70X magnification with a Leitz Dialux 22 EB.
RESULTS

Transient transfection of MKP-1 and MKP-2 blocked neurite outgrowth in PC12 cells. To compare the action of MKP-1 and MKP-2 in governing cellular differentiation, we transiently transfected PC12 cells with expression vectors carrying the coding regions of MKP-1 and MKP-2 or with the control vector. We also co-transfected a gene encoding β-galactosidase that provided a marker for transfected cells following histological staining for β-galactosidase activity. One set of transfections contained a vector encoding active Ras (RasV12) along with the other plasmids, while the other set was treated with NGF for 48 hours prior to performing the β-galactosidase staining (Figure 11). Both NGF treated cells and cells receiving the RasV12 plasmid developed neurites in 40-60% of the β-galactosidase positive cells. However, following transient transfection of MKP-1 and MKP-2, only 10-20% of the β-galactosidase positive cells showed neurites (Figure 11). These results demonstrate that a second member of the MKP family, MKP-2, can inhibit neuronal differentiation when transiently overexpressed, as we have previously shown for MKP-1 (73). In contrast, under more physiological conditions, the induction of endogenous MKP-1 and MKP-2 mRNA expression by NGF does not block neuronal differentiation (68). It is important, therefore, to examine PC12 cells where the overexpression of MKPs is maintained at more physiological levels. Therefore, we have examined stable PC12-derived cell lines that express MKPs to levels that are similar to that achieved following induction of MKPs by physiological agents.

Generation of MKP-1 and MKP-2 overexpressing stable cell lines. To investigate the consequences of limited overexpression of MAP kinase phosphatases, we generated clonal isolates that stably expressed MKPs. PC12 cells were transfected with MKP-1 and MKP-2 cDNAs under the control of the CMV promoter and selected using neomycin. The percentage of neomycin-positive and MKP-1 and MKP-2-positive cells was low. Because the size of the transcribed MKP-1 transgene was identical to the endogenous MKP-1 transcript, positive cell lines were identified by quantitation of Northern blots (Figure 12A). The MKP-2 transgene was designed to encode a short 2.4 kb transcript lacking most of the long 3' untranslated region that is present in the endogenous transcript. Positive cell lines were identified based on the presence of the additional smaller band by Northern analysis (Figure 12B). The expression of the transgenes was only 2-3 fold over basal in every positive clone analyzed. This level of expression was similar to levels achieved following induction by NGF and EGF (68). Therefore, these clonal lines
provide a model for the action of physiological levels of MKP expression. Two clones (MKP1.10 and MKP2.30 were expanded for initial study. These cells show some morphological differences compared to wild type cells. MKP1.10 cells are smaller and rounder in shape and do not clump. In contrast, MKP2.3 cells although smaller than PC12 cells, exhibit a tendency to clump and are often multinucleated (see below).

**ERK activity is reduced in MKP-1 and MKP-2 overexpressing cells.**

Growth factor, hormone, and phorbol ester stimulation of PC12 cells has been known to activate the MAP kinase pathway and to stimulate the enzymatic activity of ERK-1 (Figures 13A and 13B) (61, 62, 65, 69, 71). We compared the enzymatic activity of ERK-1, in wild type cells and in MKP overexpressing cells treated with these agents. PC12 cells treated for 10 minutes with mitogenic agents such as EGF, differentiating agents such as NGF and forskolin, and the tumor promoter phorbol 12-myristate 13-acetate (PMA), produce a robust activation of ERK-1 as measured by an immune complex activity assay (Figure 13A). MKP-1 and MKP-2 overexpressing clones were then treated with the same agents for the identical times. Both MKP1.10 and MKP2.3 cells lines showed a dramatic reduction in the ability of growth factors and hormones to activate ERK-1 (Figure 13A). Additional positive clones were also analyzed and showed similar results (data not shown).

Quantitative of the immune complex assays shows that the modest overexpression of MKP-1 and MKP-2 mRNA in PC12 cells inhibited growth factor and hormone-induced activation of ERKs 80-90% in MKP2.3 cells and 50-80% in MKP1.0 cells compared to the fold activation seen in wild type PC12 cells (Figure 13B). The basal ERK activity also appeared to be lower in these MKP overexpression cells as compared to wild type cells (Figure 13B, insert).

MKPs have been shown to dephosphorylate other MAP kinases like the JNKs (56, 64). Therefore, we examined the stimulation of JNK activity by NGF, EGF, forskolin, and PMA, as well. In wild type PC12 cells, minimal stimulation of JNK activity was seen only by EGF (1.6-fold) (Figure 14A). In MKP2.3 cells, both basal and EGF stimulated levels were only slightly reduced. EGF appears to be a more potent activator of ERKs (12.5-fold) rather than the JNKs (1.6-fold). Therefore, following stimulation by growth factors (EGF and NGF) and hormones (forskolin), JNKs were not activated significantly in either wild type or MKP2.3 cells. In contrast, JNK activity following PMA treatment in MKP2.3 cells was slightly higher than in wild type cells. UV light, a stimulator of JNK activity, resulted in a 3-4 fold induction over
control non-irradiated PC12 cells which was similar to the fold induction observed by others (64). This fold induction of JNK activation by UV light was not reduced in both the MKP1.10 and MKP2.3 cells (data not shown). We also examined the ability of MKP2.3 cells to inhibit transcriptional activation of c-jun. Cells were transfected with or without the JNK activator, MEKK, and the chimeric reporter genes Gal4-c-jun and 5XGal4-E1B-luciferase (58). MEKK has been shown to activate c-jun through its actions on JNKs (67). MEKK activated JNKs to high levels as measured by the activation of c-jun-dependent luciferase activity (Figure 14B). The c-jun activation stimulated by MEKK was not altered in MKP2.3 cells (Figure 14B). These results demonstrate that the agents examined in this study preferentially activate ERKs rather than the JNKs and that the activated ERKs, to not JNKs, are the primary targets of MKP action in these cells. Therefore, we used these cells as a model system to examine the consequences of diminished ERK activation following MKP overexpression on signaling pathways via these agents.

Both the duration and magnitude of ERK activation have been proposed to dictate physiological response to growth factors. Therefore, we examined the kinetics of ERK activation in MKP overexpression cells. We treated PC12 and MKP2.3 cells with NGF and EGF for the indicated times and performed immune complex assays with the ERK-1 antibody. The results were quantitated and showed that the magnitude of ERK activity by both NGF and EGF was severely blunted in MKP2.3 cells. However, the kinetics of the activation profile remained blunted (Figures 15A and 15B). The effect of this reduction of ERK activation on the physiological response of these cells to growth factor stimulation was subsequently addressed.

**MKP overexpression cells exhibit reduced activation of ERK-responsive transcription.** In order to determine if the MKP-induced reduction in ERK-activation by growth factors and other agents led to changes in gene expression, we compared the ability of MKP overexpression cells to activate transcription of an ERK-dependent gene through the transcriptional activator, Elk-1. Several studies have shown that ERK phosphorylation sites in the carboxyl terminal transcriptional activation domain of Elk-1 are sufficient to allow transcription response to growth factors (63). Cells were transiently transfected with the chimeric reporter genes Gal4-Elk-1 and 5XGal4-E1B-luciferase and the next day were treated with EGF or NGF for 6 hours prior to harvesting and performing luciferase assays. These agents are thought to activate Elk-1 through their action on ERKs. The physiological activation of ERKs result in
increased luciferase activity (Figure 16). The activation of Elk-1 transcriptional activity by the ERK cascade stimulators (EGF and NGF) was reduced in MKP2.3 cells. Although Elk-1 can be activated by JNKs as well as ERKs (72), neither agent activated JNKs significantly in either wild type or MKP2.3 cells (Figures 14A and 14B). Therefore, Elk-1 activation by these agents reflects ERK activation rather than JNK activation. WE conclude that the reduced EGF and NGF-induced ERK activity in these cells was responsible for the reduction in ERK-dependent gene expression. Although PMA-induced ERK activation was inhibited in MKP2.3 cells (Figures 13A and 13B), this did not result in a reduction of Elk-1 induced transcription (Figures 16A and 16B). Since Elk-1 can also be activated by JNKs (72), the increased Elk-1 induced transcription by PMA in MKP2.3 cells may be due to the increase in JNK activation seen following PMA treatment of these cells (Figure 14A).

**Regulation of endogenous MKP-1 and MKP-2 transcription in MKP overexpressing cells.** MKP-1 mRNA is undetectable in quiescent fibroblasts (55). In the wild type PC12 cell line used in this study, serum starvation does not result in loss of MKP-1 and MKP-2 expression (Figure 17A), suggesting that the basal levels of MKP expression are not dependent on serum factors in these cells. However, stimulation of these serum-starved cells with EGF, NGF, forskolin, and PMA resulted in a modest increase in MKP-1 and MKP-2 mRNA (Figure 17A). NGF and forskolin were the strongest activators of MKP-1 and MKP-2 RNA (Figure 17A). These agents also produced a sustained activation of ERKs suggesting that the transient MKP induction by these agents does no inhibit activation (73).

To assess the role of ERKs in the regulation of MKP mRNAs, we compared the induction of MKP-1 and MKP-2 in wild type cells to those seen in both MKP overexpressing cell lines (where ERK activation by these agents is reduced significantly) (Figures 13A and 13B). The induction of MKP-2 mRNA by NGF, forskolin, and PMA in the MKP1.10 cells was diminished compared to wild type cells (Figure 17B). Because the MKP-1 transgene encodes an RNA of the same size as the endogenous transcript, the effect of these agents on the endogenous MKP-1 gene in MKP1.10 cells could not be distinguished. The robust induction of both MKP-1 and MKP-2 transgenes by forskolin and PMA in MKP overexpressing cells may be due to the presence of cAMP/PMA response elements present with the CMV promoter that was used to direct the expression of the MKP transgenes. This was most clearly seen in the transgene-specific MKP-2 band (Figure 17C).
In cells overexpressing MKP-2, the size of the exogenous transgene was distinguishable from the endogenous MKP-2 transcript (Figure 17C) making it possible to examine the regulation of the endogenous MKP-2 gene directly in MKP2.3 cells. In these cells, the induction of endogenous MKP-1 mRNA by EGF, NGF, and forskolin but not PMA was diminished compared to wild type cells. The induction of endogenous MKP-2 mRNA in MKP2.3 cells was more substantially reduced by EGF, NGF, forskolin, and by PMA as well (Figure 17C). Again, the larger induction of exogenous MKP-2 by forskolin and PMA may be due to the ability of these agents to activate the CMV promoter. Taken together, these results identify an ERK-dependent component to the regulation of both MKP-1 and MKP-2 mRNA expression. The induction of endogenous MKP-2 mRNA by agents that activate ERKs was more severely blunted in MKP-overexpressing cells, suggesting that the induction of MKP-2 mRNA may be more dependent on ERK activity than MKP-1.

Effect of MKP overexpression on the immediate early gene, c-fos.

Transient transfection of MKP-1 has been shown to reduce serum-stimulated, ERK-dependent activity of the c-fos promoter (54). Since ERK activation by growth factors and other agents was substantially reduced in the MKP2.3 cell line, the requirement of ERK activation in c-fos regulation was examined. We assayed the expression of the immediate early gene, c-fos, 30 minutes following drug treatment. As compared to wild type PC12 cells, induced c-fos expression by NGF, EGF, and forskolin was reduced in MKP2.3 cells (Figure 18). Modest overexpression of MKP-2, therefore, is partially able to block the signal-induced overexpression of the immediate early gene, c-fos, by multiple activators of the MAP kinase cascade. In contrast, PMA-induced expression of c-fos was not reduced in these cells (Figure 18). This effect of PMA may be related to the paradoxical increase in JNK activation following PMA-stimulation of these cells as compared to wild type cells.

Effect of MKP-1 and MKP-2 overexpression on the late gene, stromelysin-1 (transin). Stromelysin-1 (Transin) is a late gene that is induced in wild type PC12 cells upon treatment with NGF and PMA (Figure 19) and its expression is associated with neuronal differentiation (59). Because stromelysin may be regulated by ERK-dependent pathways (73), we analyzed the expression of stromelysin in these cells. Stromelysin induction by NGF was almost completely locked in MKP1.10 cells and was partially blocked in MKP2.3 cells (Figure 19). The induction of stromelysin was completely inhibited by PMA in both the MKP1.10 and MKP2.3 cells. Identical results were achieved using multiple clonal isolates of MKP.
overexpressing cells (data not shown). Therefore, reduction of ERK activity by these agents was associated with inhibition of the expression of the late gene stromelysin.

**Effect of MKP-1 and MKP-2 overexpression on proliferation and differentiation.** Previous reports suggest that the sustained activation of ERKs by NGF is required for PC12 cell differentiation whereas the transient activation of ERKs by EGF is required for proliferation (57, 66). In order to examine the biological consequences of MKP overexpression and reduced growth factor-inducible ERK activation, we measured the proliferation rate of MKP1.10 and MKP2.3 cells. The proliferation rate of parental PC 12 cells growing in serum was higher than that of MKP1.10 and MKP2.3 cells (Figure 20A). MKP2.3 cells had the slowest rate of proliferation. When the cells were partially synchronized by serum starvation for 2 days and then stimulated with serum, both MKP1.10 and MKP 2.3 cells were delayed in their entry into the cell cycle (Figure 20B). Flow cytometric analysis on the three cell lines confirmed these results with MKP1.10 and MKP2.3 cell lines showing a decrease in the percent of cells in S phase compared to wild type cells (data not shown). These results suggest that the reduced growth factor-stimulated ERK activity is associated with a reduction in proliferation in both MKP-1 and MKP-2 overexpressing cells.

We next examined the ability of NGF to induce differentiation in the MKP overexpressing cells. Despite the severely blunted response of MKP1.10 and MKP2.3 cells to ERK activation by NGF, these cells were able to extend neuritic process similar to their parental cell line (Figure 21). The time course of this differentiative response in MKP overexpression cells was similar to wild type cells (Figure 21). Additional MKP overexpressing clones were also treated with NGF and similar results were obtained (data not shown). These results suggest that high levels of sustained ERK activity are not absolutely required for morphological differentiation.

**EXAMPLE 3**

As discussed above, the mitogen-activated protein kinase cascade mediates signals that govern both cell growth and differentiation. The MAP kinases initially included only the extracellular signal regulated kinases (ERKs) which respond to mitogenic signals. However, another pathway is known to respond to a variety of cellular stresses by activating a related family of MAP kinases, called stress-activated protein kinases (SAP kinases) or jun N-terminal protein kinases (JNKs) (74). At least two JNKs, JNK-1 (81) and JNK-2 (75), response to a variety of Ras-
dependent and Ras-independent signals to phosphorylate the proto-oncogene c-jun (74, 75, 80, 82) and Elk-1 (83). The phosphorylation of c-jun is associated with growth arrest and stimulation of apoptosis (75-78). Because MAP kinase activation and inactivation is believed to be regulated by MKP-1 and MKP-2 of the present invention, a study was performed that demonstrated that JNK-mediated signaling pathways induce apoptosis in both neuronal and fibroblast cells and that this can be partially reversed by MKPs through a mechanism that is independent of their action on ERKs. Thus, the MKP-2 of the present invention may play a critical role in determining cell fate by selective regulation of mitogenic and stress-activated pathways, e.g., inhibiting apoptosis in response to activation by growth factors on oncogenes.

Based on the foregoing, it will be appreciated that the MKP-2 of the present invention and inhibitors thereto, may be used in therapeutic applications such as cancer therapy. Expression of MKPs in human tumors including prostate, breast, colon, lung and bladder, function to direct oncogenic signals into a proliferative pathway away from apoptosis. The use of MKP inhibitors such as an inhibitor of the MKP-2 of the present invention, will selectively act on tumor cells to redirect the oncogenic signal into apoptotic pathways. It will be appreciated that such novel chemotherapeutics may be combined with current non-surgical treatment for human cancers including radiation and chemotherapy, both of which are stimulators of the stress-activated protein kinase cascade (79, 85) that kill tumor cells by triggering apoptosis (86). Moreover, because apoptosis in neuronal cells contributes to the morbidity associated with neurodegenerative diseases, stroke and Alzheimer's dementia (84), the MKP-2 of the present invention may be used in the development of novel therapeutic strategies for neurodegenerative diseases as well.

While the preferred embodiment of the invention has been illustrated and described, it will be appreciated that various changes can be made therein without departing from the spirit and scope of the invention.

All publications cited herein are incorporated by reference to the same extent as if each individual publication was specifically and individually indicated to be incorporated by reference. The invention now being fully described, it will be apparent to one of ordinary skill in the art that many changes and modifications can be made thereto without departing from the spirit or scope of the invention.

REFERENCES

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Chem. 270, 8377-8380.


WE CLAIM:

1. An isolated nucleic acid comprising a nucleotide sequence encoding mammalian MKP-2 protein.

2. The isolated nucleic acid of Claim 1 comprising a nucleotide sequence encoding mammalian MKP-2 protein having an amino acid sequence shown in Figure 22 or homologs thereof.

3. The nucleic acid of Claim 2 comprising a nucleotide sequence encoding mammalian MKP-2 protein having an amino acid sequence greater than about 90% homologous with the amino acid sequence shown in Figure 22.

4. An isolated nucleic acid comprising a nucleotide sequence encoding mammalian MKP-2 protein, wherein said sequence encoding the mammalian MKP-2 protein is capable of hybridizing under low stringency conditions to the complement of the nucleotide sequence shown in Figure 22.

5. An isolated nucleic acid comprising the nucleotide sequence shown in Figure 1B.

6. An isolated nucleic acid comprising the nucleotide sequence shown in Figure 22.

7. A recombinant vector comprising the nucleic acid of any of Claims 1-6.

8. A transformant host cell transfected with the vector of Claim 7.

9. A method of expressing recombinant nucleic acid which comprises culturing the cell of Claim 8 under conditions which allow for the expression of the nucleic acid with which it has been transfected.

10. A protein encoded by the nucleic acid of any of Claims 1-6.
11. The protein of Claim 10 having an amino acid sequence as shown in Figure 22.

12. An antibody capable of binding the protein of any of Claims 10-11.

13. The antibody of Claim 12, wherein the antibody is a monoclonal antibody.

14. The antibody of Claim 12, wherein the antibody is a polyclonal antibody.

15. A hybridoma cell line capable of producing the antibody of any of Claims 13-14.
FIGURE 5
10/22

**U** 15% serum

<table>
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<th>15'</th>
<th>30'</th>
<th>1h</th>
<th>2.5h</th>
<th>4h</th>
<th>6h</th>
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</table>

| 30' | 1h  | 2h  | 4h  | 30' | 1h  | 2h  | 4h  |

- **MKP-2**
- **MKP-1**
- **18S**

**HiDA.**

**MKP-2**

**MKP-1**

**HiDA.**

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FIGURE 11

Treatment: NGF + + + - - -
Plasmids: RasV12 - - - + + +
MKP-1 - + - - + -
MKP-2 - - + - - +
β-gal + + + + + +

% neurites
FIGURE 15A

Fold activation of ERK activity/100 µg total protein

Time in the presence of 100 ng/ml NGF (minutes)

FIGURE 15B

Fold activation of ERK activity/100 µg total protein

Time in the presence of 50 ng/ml EGF (minutes)

FIGURE 16

Luciferase activity/µg protein (arbitrary units)

Control EGF NGF PMA
**Figure 18.**

**Figure 19.**

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FIGURE 20A

Growth in serum

BrdU-index of proliferation (arbitrary units)

Days

FIGURE 20B

Growth after serum stimulation of serum starved cells

BrdU-index of proliferation (arbitrary units)

Days

SUBSTITUTE SHEET (RULE 26)
**INTERNATIONAL SEARCH REPORT**

**A. CLASSIFICATION OF SUBJECT MATTER**
- IPC(6) - Please See Extra Sheet.
- US CL : 435/183, 194; 530/352, 388.26; 536/23.2
- According to International Patent Classification (IPC) or to both national classification and IPC

**B. FIELDS SEARCHED**
- Minimum documentation searched (classification system followed by classification symbols)
  - U.S. : 435/183, 194; 530/352, 388.26; 536/23.2
- Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched
  - Electronic database consulted during the international search (name of database and, where practicable, search terms used)
    - MEDLINE, WPIDS, APS

**C. DOCUMENTS CONSIDERED TO BE RELEVANT**

<table>
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<th>Citation of document, with indication, where appropriate, of the relevant passages</th>
<th>Relevant to claim No.</th>
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<td>X, E</td>
<td>US 5,512,434 A (AARONSON ET AL) 30 April 1996 (30.04.96), see entire document.</td>
<td>1-11</td>
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</table>

**X** Further documents are listed in the continuation of Box C. □ See patent family annex.

- * Special categories of cited documents:
  - "A" - document defining the general state of the art which is not considered to be of particular relevance
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**Date of the actual completion of the international search**
- 27 AUGUST 1996

**Date of mailing of the international search report**
- 09 SEP 1996

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<td>Y</td>
<td>SUN et al. MKP-1 (3CH13), an immediate early gene product, is a dual specificity phosphatase that dephosphorylates MAP kinase in vivo. Cell. 05 November 1993, Vol. 75, pages 487-493, see entire document.</td>
<td>1-15</td>
</tr>
</tbody>
</table>
A. CLASSIFICATION OF SUBJECT MATTER:

IPC (6):

C12N 9/00, 9/12; A61K 38/16; C07K 1/00, 14/00, 16/00, 17/00; C12P 21/08; C07H 21/04