(54) Title: MNK KINASE HOMOLOGOUS PROTEINS INVOLVED IN THE REGULATION OF ENERGY HOMEOSTASIS AND ORGANELLE METABOLISM

(57) Abstract: The present invention discloses Mnk homologous proteins regulating the energy homeostasis, the metabolism of triglycerides, and/or is contributing to membrane stability and/or function of organelles, and polynucleotides, which identify and encode the proteins disclosed in this invention. The invention also relates to the use of these sequences in the diagnosis, study, prevention, and treatment of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs, and others.
Mnk kinase homologous proteins involved in the regulation of energy homeostasis and organelle metabolism

Description

This invention relates to the use of nucleic acid sequences of the MAP kinase-interacting kinase (Mnk) gene family and amino acid sequences encoded thereby, and to the use of these sequences or effectors of Mnk nucleic acids or polypeptides, particularly Mnk kinase inhibitors and activators, in the diagnosis, study, prevention, and treatment of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs.

There are several metabolic diseases of human and animal metabolism, eg., obesity and severe weight loss, that relate to energy imbalance where caloric intake versus energy expenditure is imbalanced. Obesity is one of the most prevalent metabolic disorder in the world. It is a still poorly understood human disease that becomes more and more relevant for western society. Obesity is defined as a body weight more than 20% in excess of the ideal body weight, frequently resulting in a significant impairment of health. It is associated with an increased risk for cardiovascular disease, hypertension, diabetes, hyperlipidemia and an increased mortality rate. Besides severe risks of illness, individuals suffering from obesity are often isolated socially.
Obesity is influenced by genetic, metabolic, biochemical, psychological, and behavioral factors. As such, it is a complex disorder that must be addressed on several fronts to achieve lasting positive clinical outcome. Since obesity is not to be considered as a single disorder but as a heterogeneous group of conditions with (potential) multiple causes, it is also characterized by elevated fasting plasma insulin and an exaggerated insulin response to oral glucose intake (Koltermann, J. Clin. Invest 65, 1980, 1272-1284). A clear involvement of obesity in type 2 diabetes mellitus can be confirmed (Kopelman, Nature 404, 2000, 635-643).

The molecular factors regulating food intake and body weight balance are incompletely understood. Even if several candidate genes have been described which are supposed to influence the homeostatic system(s) that regulate body mass/weight, like leptin, VCPI, VCPL or the peroxisome proliferator-activated receptor-gamma co-activator, the distinct molecular mechanisms and/or molecules influencing obesity or body weight/body mass regulations are not known. In addition, several single-gene mutations resulting in obesity have been described in mice, implicating genetic factors in the etiology of obesity (Friedman and Leibel, 1990, Cell 69: 217-220). In the obese mouse, a single gene mutation (obese) results in profound obesity, which is accompanied by diabetes (Friedman et. al., 1991, Genomics 11: 1054-1062).

Therefore, the technical problem underlying the present invention was to provide for means and methods for modulating (pathological) metabolic conditions influencing thermogenesis, body-weight regulation and/or energy homeostatic circuits. The solution to said technical problem is achieved by providing the embodiments characterized in the claims.

Accordingly, the present invention relates to genes with novel functions in body-weight regulation, energy homeostasis, metabolism, and obesity. The present invention provides for a specific gene involved in the regulation of
diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, cancers of the reproductive organs, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer. The present invention describes the human Mnk genes as being involved in those conditions mentioned above, in particular the human Mnk2 gene variants.

The term "GenBank Accession number" relates to National Center for Biotechnology Information (NCBI) GenBank database entries (Benson et al, Nucleic Acids Res. 28, 2000, 15-18).

Protein kinases are important molecules involved in the regulation of many cellular functions. The Drosophila melanogaster LK6 serin/threonine kinase gene has been described as a short-lived kinase that can associate with microtubules (J. Cell Sci. 1997 110(2):209-219). Genetic analysis in the development of the Drosophila compound eye suggested a role in the modulation of the RAS signalling pathway (Genetics 2000 156(3):1219-1230). As described in this invention, the closest human homologues of Drosophila LK6 kinase are the MAP kinase-interacting kinase 2 (Mnk2, for example the variants Mnk2a and Mnk2b) and MAP kinase-interacting kinase 1 (Mnk1). All three proteins are predominantly localized in the cytoplasm. Mnks are phosphorylated by the pk42 MAP kinases Erk1 and Erk2 and the p38 MAP kinases. This phosphorylation is triggered in response to growth factors, phorbol esters and oncogenes like Ras and Mos as well as by stress signaling molecules and cytokines. The phosphorylation of Mnk proteins stimulates its kinase activity towards eukaryotic initiation factor 4E (EMBO J. 16: 1909-1920 (1997), Mol Cell Biol 19:1871-1880 (1999), Mol Cell Biol 21: 743-754 (2001)).
Phosphorylation of eukaryotic initiation factor 4E (eIF4E) results in a regulation of protein translation (Mol Cell Biol 22: 5500-5511 (2001)).

There are different hypothesis describing the mode of stimulation of the protein translation by Mnk proteins. Most publications described a positive stimulatory effect on the cap-dependent protein translation upon activation of MAP kinase-interacting kinases. Thus, activation of Mnk proteins might lead to an indirect stimulation or regulation of protein translation, for example by the action on cytosolic phospholipase 2 alpha (BBA 1488:124-138, 2000).

Inhibitors of Mnk (referred to as CGP57380 and CGP052088) were described in the prior art (see, Knauf et al., 2001, Mol. Cell. Biol. 21:5500, Tschopp et al., 2000, Mol Cell Biol Res Comm 3:205 and Slentz-Kesler et al., 2000, Genomics 69:63). CGP052088 is a staurosporine derivative with an IC50 of 70 nM for inhibition of in vitro kinase activity of Mnk1. CGP57380 is a selective low-molecular weight, non cytotoxic inhibitor of Mnk2 (Mnk2a or Mnk2b) or Mnk1. The addition of CGP57380 to cell culture cells transfected with Mnk2 (Mnk2a or Mnk2b) or Mnk1 resulted in a strong reduction in phosphorylated eIF4E.

So far, it has not been described that Mnk kinases are involved in the regulation of body-weight and thermogenesis, and thus may be associated with metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs. In this application we demonstrate that the correct gene doses of Mnk kinases are essential for maintenance of energy homeostasis. A genetic screen was used to identify that mutation of Mnk kinase homologous genes causes obesity, reflected by a significant
increase of triglyceride content, the major energy storage substance. Furthermore, in this invention we relate to mutations of Mnk kinases that affect the activity of uncoupling proteins (UCPs), thereby leading to an altered mitochondrial activity. We also relate to the treatment of metabolic disorders with the Mnk-specific inhibitor CGP57380 and derivatives thereof.

In this invention we demonstrate that the correct gene dose of the Drosophila melanogaster homologue of Mnk is essential for maintenance of energy homeostasis in adult flies and for the activity of mitochondrial uncoupling protein. A genetic screen was used to identify that mutation of an Mnk homologous gene causes obesity in Drosophila melanogaster, reflected by a significant increase of triglyceride content, the major energy storage substance. In a second screen designed to identify factors that modulate activity of uncoupling protein, we discovered that mutation of this Mnk homologous gene caused a reduction the activity of uncoupling protein. Thus, the invention is also based on the finding that the Drosophila homologue of Mnk is contributing to membrane stability and/or function of organelles, preferably mitochondria. It was found that mutations in LK6 kinases affect the activity of uncoupling proteins (UCPs), thereby leading to an altered mitochondrial activity.

Further, we show that the mouse homologue of the Mnk2 gene is regulated by fasting and by genetically induced obesity. Furthermore, the Mnk2 mRNA is strongly upregulated during adipocyte differentiation in vitro (see EXAMPLES). This invention shows that Mnk2 transcripts are expressed in most mouse tissues but with highest expression levels in white (WAT) and brown adipose tissue (BAT). The expression in white adipose tissue is reduced by approx. 60% in fasted mice and in ob/ob mice. The analysis of actin-mMnk2DN transgenic mice showed that the ectopic expression of mMnk2DN transgene (see Examples) leads to an clear
increase in bodyweight. The effect seems to be diet-independent, as it can be seen on control diet as well as on high fat diet. Thus, we conclude that Mnk2 is playing an important role in the regulation of body-weight.

In addition, we found that the relative expression levels of both human Mnk2 splice variants is the same for all tissues analysed. Both Mnk2 variants show highest expression levels in human tissues relevant for metabolic disorders namely adipose and muscle tissue. Furthermore, both Mnk2 variants are upregulated during human adipocyte differentiation. Thus, we conclude that Mnk2 (or variants thereof) has a function in the metabolism of mature human adipocytes.

We also found that cellular triglyceride levels in Mnk2 overexpressing cells were significantly lower from day 4 to day 12 of adipogenesis compared to that in the control cells. Furthermore, Mnk2 overexpressing cells were less effective at synthesising lipids from exogenous glucose. Consequently, the levels of insulin stimulated lipid synthesis are significantly lower at day 12 of adipogenesis when compared to control cells. We also found that transport of exogenous fatty acids across the plasma membrane of Mnk2 overexpressing cells and hence esterification of these metabolites were considerably lower at day 12 of adipogenesis when compared to control cells.

Polynucleotides encoding a protein with homologies to proteins of the Mnk kinase family are suitable to investigate diseases and disorders as described above. Discovery of molecules related to Mnk kinases satisfies a need in the art by providing new compositions useful in diagnosis, treatment, and prognosis of diseases and disorders as described above.

Before the present proteins, nucleotide sequences, and methods are described, it is understood that this invention is not limited to the particular methodology, protocols, cell lines, vectors, and reagents described as
these may vary. It is also to be understood that the terminology used herein is for the purpose of describing particular embodiments only, and is not intended to limit the scope of the present invention, which will be limited only by the appended claims. Unless defined otherwise, all technical and scientific terms used herein have the same meanings as commonly understood by one of ordinary skill in the art to which this invention belongs. Although any methods and materials similar or equivalent to those described herein can be used in the practice or testing of the present invention, the preferred methods, devices, and materials are now described. All publications mentioned herein are incorporated herein by reference for the purpose of describing and disclosing the cell lines, vectors, and methodologies, which are reported in the publications which might be used in connection with the invention. Nothing herein is to be construed as an admission that the invention is not entitled to antedate such disclosure.

The present invention discloses that Mnk homologous proteins are regulating the energy homeostasis and fat metabolism, especially the metabolism and storage of triglycerides, and polynucleotides, which identify and encode the proteins disclosed in this invention. The present invention also discloses that Mnk homologous proteins are directly or indirectly involved in membrane stability and/or function of organelles, in particular mitochondria, and polynucleotides, which identify and encode the proteins disclosed in this invention. The invention also relates to vectors, host cells, antibodies, and recombinant methods for producing the polypeptides and polynucleotides of the invention. The invention also relates to the use of these sequences in the diagnosis, study, prevention, and treatment of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep
apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs.

Mnk homologous proteins and nucleic acid molecules coding therefore are obtainable from insect or vertebrate species, e.g. mammals or birds. Particularly preferred are human Mnk homologous polypeptides and nucleic acids encoding such polypeptides, particularly polypeptides and nucleic acids encoding a human Mnk2 protein (splice variant Mnk2a, Genbank Accession No. AF237775 as shown in Figure 3D and 3E, or splice variant Mnk2b, GenBank Accession AF237776 or No. NM_017572.1, as shown in Figure 3F and 3G, Genbank Accession No. AF237775 is identical to formerly Genbank Accession No. XM_030637 which was removed at the submitters request; see a Clustal W multiple sequence alignment in Figure 3B, see also sequences in Figures 3D-G) or a human Mnk1 protein (Genbank Accession No. AB000409.1 and NM_003684.2 as shown in Figure 3H and 3i); Genbank Accession No. AB000409 is identical to formerly Genbank Accession No. XM_001600 which was removed at the submitters request; see a Clustal W multiple sequence alignment in Figure 3C).

The invention particularly relates to a nucleic acid molecule encoding a polypeptide contributing to regulating the energy homeostasis and the metabolism of triglycerides, and/or contributing to membrane stability and/or function of organelles, wherein said nucleic acid molecule comprises

(a) the nucleotide sequences of Genbank Accession Nos. AF237775, NM_017572.1, AB000409.1, or NM_003684.2, and/or the complement thereof,

(b) a nucleotide sequence which hybridizes at 50°C in a solution containing 1 x SSC and 0.1% SDS to the nucleic acid molecule of
(a), particularly a nucleic acid encoding the amino acid sequences as shown in figure 3,

(c) a sequence corresponding to the sequences of (a) or (b) within the degeneration of the genetic code,

(d) a sequence which encodes a polypeptide which is at least 85%, preferably at least 90%, more preferably at least 95%, more preferably at least 98% and up to 99.6% identical to the amino acid sequences shown in Figure 3,

(e) a sequence which differs from the nucleic acid molecule of (a) to (d) by mutation and wherein said mutation causes an alteration, deletion, duplication or premature stop in the encoded polypeptide or a partial sequence of any of the nucleotide sequences of (a) to (e) having a length of at least 15 bases, preferably at least 20 bases, more preferably at least 25 bases and most preferably at least 50 bases.

The invention is based on the finding that Mnk homologous proteins (herein referred to as Mnk), particularly Mnk2 (Mnk2a or Mnk2b) or Mnk1, and the polynucleotides encoding these, are involved in the regulation of triglyceride storage and therefore energy homeostasis. The present invention also discloses that Mnk homologous proteins are directly or indirectly involved in membrane stability and/or function of organelles, in particular mitochondria, and polynucleotides, which identify and encode the proteins disclosed in this invention. The invention describes the use of compositions comprising the nucleotides, proteins or effectors thereof, e.g. antibodies, aptamers, anti-sense molecules, ribozymes, RNAi molecules, peptides, low-molecular weight organic molecules and other receptors recognizing the nucleic acid molecule or the polypeptide, for the diagnosis, study, prevention, or treatment of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart
disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs.

Accordingly, the present invention relates to genes with novel functions in body-weight regulation, energy homeostasis, metabolism, and obesity. To find genes with novel functions in energy homeostasis, metabolism, and obesity, a functional genetic screen was performed with the model organism Drosophila melanogaster (Meigen). Drosophila melanogaster is one of the most intensively studied organisms in biology and serves as a model system for the investigation of many developmental and cellular processes common to higher eukaryotes, including humans (see, for example, Adams et al., Science 287: 2185-2195 (2000)). The success of Drosophila melanogaster as a model organism is largely due to the power of forward genetic screens to identify the genes that are involved in a biological process (see, Johnston Nat Rev Genet 3: 176-188 (2002); Rorth, Proc Natl Acad Sci U S A 93: 12418-12422 (1996)). One resource for screening was a proprietary Drosophila melanogaster stock collection of EP-lines. The P-vector of this collection has Gal4-UAS-binding sites fused to a basal promoter that can transcribe adjacent genomic Drosophila sequences upon binding of Gal4 to UAS-sites. This enables the EP-line collection for overexpression of endogenous flanking gene sequences. In addition, without activation of the UAS-sites, integration of the EP-element into the gene is likely to cause a reduction of gene activity, and allows determining its function by evaluating the loss-of-function phenotype.

Triglycerides are the most efficient storage for energy in cells, and are significantly increased in obese patients. In this invention, we have used a genetic screen to identify, that mutations of Lk6 homologous genes cause changes in the body weight which is reflected by a significant change in the triglyceride levels. In order to isolate genes with a function in energy
homeostasis, several thousand EP-lines were tested for their triglyceride content after a prolonged feeding period. Lines with significantly changed triglyceride content were selected as positive candidates for further analysis. In this invention, the content of triglycerides of a pool of flies with the same genotype after feeding for six days was analyzed using a triglyceride assay, as, for example, but not for limiting the scope of the invention, is described below in the examples section. The change of triglyceride content due to the loss of a gene function suggests gene activities in energy homeostasis in a dose dependent manner that controls the amount of energy stored as triglycerides.

The result of the triglyceride content analysis is shown in Figure 1. Flies homozygous for EP(3)3333 and EP(3)3576 integrations were analyzed in the triglyceride assay. The average increase of triglyceride content of the homozygous viable lines EP(3)3333 and EP(3)3576 is approx. 140 % (Figure 1). Therefore, the very likely loss of a gene activity in the gene locus 86F7 (estimated, chromosomal localisation where the EP-vector of EP(3)3333 and EP(3)3576 flies is integrated) is responsible for changes in the metabolism of the energy storage triglycerides, therefore representing in both cases an obese fly model. The increase of triglyceride content due to the loss of a gene function suggests gene activities in energy homeostasis in a dose dependent manner that controls the amount of energy stored as triglycerides.

Nucleic acids encoding the Mnk protein of the present invention were identified using a plasmid-rescue technique. Genomic DNA sequences were isolated that are localised directly 3' to the EP(3)3333 and EP(3)3576 integrations. Using those isolated genomic sequences public databases like Berkeley Drosophila Genome Project (GadFly; see also FlyBase (1999) Nucleic Acids Research 27:85-88) were screened thereby confirming the integration side of EP(3)3333 in the 5' region of a 5' exon of the Mnk homologous gene and EP(3)3576 in the 5' region of an alternative 5' exon
(Figure 2). Figure 2 shows the molecular organisation of this locus. Genomic DNA sequence is represented by the assembly as a black dotted line in the middle that includes the integration site of EP(3)3333 and EP(3)3576. Numbers represent the coordinates of the genomic DNA (starting at position 7544500 on chromosome 3R). Grey bars on the two "cDNA"-lines represent the predicted genes (GadFly & Magpie), and grey symbols on the "P-Elements"-line the EP-vector integration sites. Predicted exons of gene CG17342 are shown as dark grey bars and predicted introns as light grey bars.

Lk6 (the Mnk homologous gene in Drosophila) encodes for a gene that is predicted by GadFly sequence analysis programs (GadFly Accession Number CG17342). No functional data described the regulation of obesity and metabolic diseases are available in the prior art for the genes shown in Figure 3, referred to as Mnk in the present invention.

It is also preferred that the nucleic acid molecule encodes a polypeptide contributing to membrane stability and/or function of orgnelles and represents a protein of Drosophila which has been found to be able to modify UCPs, see also appended examples. As demonstrated in the appended examples, the here described polypeptide (and encoding nucleic acid molecule) was able to modify, e.g. enhance a specific eye phenotype in Drosophila which was due to the overexpression of the Drosophila melanogaster gene dUCPy. The overexpression of dUCPy (with homology to human UCPs) in the compound eye of Drosophila led to a clearly visible eye defect which can be used as a "read-out" for a genetical "modifier screen".

In said "modifier screen" thousands of different genes are mutagenized to modify their expression in the eye. Should one of the mutagenized genes interact with dUCPy and modify its activity an enhancement or suppression of the eye defect will occur. Since such flies are easily to discern they can
be selected to isolate the interacting gene. As shown in the appended examples, a gene was deduced that can enhance the eye defect induced by the activity of dUCPy. This gene is called the LK6 gene of Drosophila with high homologies to the human Mnk proteins, as described above. It is envisaged that mutations in the herein described Mnk-polypeptides (and genes) lead to phenotypic and/or physiological changes which may comprise a modified and altered mitochondrial activity. This, in turn, may lead to, inter alia, an altered energy metabolism, altered thermogenesis and/or altered energy homeostasis. As shown in the appended examples, a gene was deduced that can enhance the eye defect induced by the activity of dUCPy.

Mnk homologous proteins and nucleic acid molecules coding therefor are obtainable from insect or vertebrate species, e.g. mammals or birds. Particularly preferred are nucleic acids encoding the human Lk6 / Mnk homologs, particularly Mnk2 variants (Mnk2a or Mnk2b) or Mnk1. The present invention is describing a polypeptide comprising the amino acid sequence of Mnk, particularly Mnk2 variants (Mnk2a or Mnk2b) or Mnk1. A comparison (Clustal X 1.8) between the Mnk proteins of different species (human and Drosophila) was conducted and is shown in Figure 3A. Based upon homology, Mnk protein of the invention and each homologous protein or peptide may share at least some activity.

In a particular embodiment, the invention encompasses the polynucleotide comprising the nucleic acid sequence of GenBank Accession Number AF237775, NM_017572.1, AB000409.1, or NM_003684.2. It will be appreciated by those skilled in the art that as a result of the degeneracy of the genetic code, a multitude of nucleotide sequences encoding Mnk, some bearing minimal homology to the nucleotide sequences of any known and naturally occurring gene, may be produced. Thus, the invention contemplates each and every possible variation of nucleotide sequence that could be made by selecting combinations based on possible codon choices.
These combinations are made in accordance with the standard triplet genetic code as applied to the nucleotide sequences of naturally occurring Mnk, and all such variations are to be considered as being specifically disclosed. Although nucleotide sequences which encode Mnk and its variants are preferably capable of hybridising to the nucleotide sequences of the naturally occurring Mnk under appropriately selected conditions of stringency, it may be advantageous to produce nucleotide sequences encoding Mnk or its derivatives possessing a substantially different codon usage. Codons may be selected to increase the rate at which expression of the peptide occurs in a particular prokaryotic or eukaryotic host in accordance with the frequency with which particular codons are utilised by the host. Other reasons for substantially altering the nucleotide sequence encoding Mnk and its derivatives without altering the encoded amino acid sequences include the production of RNA transcripts having more desirable properties, such as a greater half-life, than transcripts produced from the naturally occurring sequences. The invention also encompasses production of DNA sequences, or portions thereof, which encode Mnk and its derivatives, entirely by synthetic chemistry. After production, the synthetic sequence may be inserted into any of the many available expression vectors and cell systems using reagents that are well known in the art at the time of the filing of this application. Moreover, synthetic chemistry may be used to introduce mutations into a sequence encoding Mnk any portion thereof.

Also encompassed by the invention are polynucleotide sequences that are capable of hybridizing to the claimed nucleotide sequences, and in particular, those shown in GenBank Accession Numbers AF237775, NM_017572.1, AB000409.1, or NM_003684.2, under various conditions of stringency. Hybridization conditions are based on the melting temperature (Tm) of the nucleic acid binding complex or probe, as taught in Wahl, G. M. and S. L. Berger (1987: Methods Enzymol. 152:399-407) and Kimmel, A. R. (1987; Methods Enzymol. 152:507-511), and may be
used at a defined stringency. Preferably, hybridization under stringent conditions means that after washing for 1 h with 1 x SSC and 0.1% SDS at 50°C, preferably at 55°C, more preferably at 62°C and most preferably at 68°C, particularly for 1 h in 0.2 x SSC and 0.1% SDS at 50°C, preferably at 55°C, more preferably at 62°C and most preferably at 68°C, a positive hybridization signal is observed. Altered nucleic acid sequences encoding Mnk which are encompassed by the invention include deletions, insertions, or substitutions of different nucleotides resulting in a polynucleotide that encodes the same or a functionally equivalent Mnk.

The encoded proteins may also contain deletions, insertions, or substitutions of amino acid residues, which produce a silent change and result in a functionally equivalent Mnk. Deliberate amino acid substitutions may be made on the basis of similarity in polarity, charge, solubility, hydrophobicity, hydrophilicity, and/or the amphipathic nature of the residues as long as the biological activity of Mnk is retained. For example, negatively charged amino acids may include aspartic acid and glutamic acid; positively charged amino acids may include lysine and arginine; and amino acids with uncharged polar head groups having similar hydrophilicity values may include leucine, isoleucine, and valine; glycine and alanine; asparagine and glutamine; serine and threonine; phenylalanine and tyrosine.

Also included within the scope of the present invention are alleles of the genes encoding Mnk. As used herein, an "allele" or "allelie sequence" is an alternative form of the gene, which may result from at least one mutation in the nucleic acid sequence. Alleles may result in altered mRNAs or polypeptides whose structures or function may or may not be altered. Any given gene may have none, one, or many allelic forms. Common mutational changes, which give rise to alleles, are generally ascribed to natural deletions, additions, or substitutions of nucleotides. Each of these types of changes may occur alone, or in combination with the others, one or more
times in a given sequence. Methods for DNA sequencing which are well
known and generally available in the art may be used to practice any
embodiments of the invention. The methods may employ such enzymes as
the Klenow fragment of DNA polymerase I, SEQUENASE DNA Polymerase
(US Biochemical Corp, Cleveland Ohio), Taq polymerase (Perkin Elmer),
thermostable T7 polymerase (Amersham, Chicago, Ill.), or combinations of
recombinant polymerases and proof-reading exonucleases such as the
ELONGASE Amplification System (GIBCO/BRL, Gaithersburg, Md.).
Preferably, the process is automated with machines such as the Hamilton
MICROLAB 2200 (Hamilton, Reno Nev.), Peltier thermal cycler (PTC200;
MJ Research, Watertown, Mass.) and the ABI 377 DNA sequencers
(Perkin Elmer). The nucleic acid sequences encoding Mnk may be extended
utilising a partial nucleotide sequence and employing various methods
known in the art to detect upstream sequences such as promoters and
regulatory elements. For example, one method which may be employed,
"restriction-site" PCR, uses universal primers to retrieve unknown
sequence adjacent to a known locus (Sarkar, G. (1993) PCR Methods
Applic. 2:318-322). Inverse PCR may also be used to amplify or extend
sequences using divergent primers based on a known region (Triglia, T. et
al. (1988) Nucleic Acids Res. 16:8186). Another method which may be
used is capture PCR which involves PCR amplification of DNA fragments
adjacent to a known sequence in human and yeast artificial chromosome
DNA (Lagerstrom, M. et al. (PCR Methods Applic. 1:111-119). Another
method which may be used to retrieve unknown sequences is that of
Parker, J. D. et al. (1991; Nucleic Acids Res. 19:3055-3060). Additionally,
one may use PCR, nested primers, and PROMOTERFINDER libraries to walk
in genomic DNA (Clontech, Palo Alto, Calif.). This process avoids the need
to screen libraries and is useful in finding intron/exon junctions.

When screening for full-length cDNAs, it is preferable to use libraries that
have been size-selected to include larger cDNAs. Also, random-primed
libraries are preferable, in that they will contain more sequences, which
contain the 5’ regions of genes. Use of a randomly primed library may be especially preferable for situations in which an oligo d(T) library does not yield a full-length cDNA. Genomic libraries may be useful for extension of sequence into the 5’ and 3’ non-transcribed regulatory regions. Capillary electrophoresis systems, which are commercially available, may be used to analyse the size or confirm the nucleotide sequence of sequencing or PCR products. In particular, capillary sequencing may employ flowable polymers for electrophoretic separation, four different fluorescent dyes (one for each nucleotide) which are laser activated, and detection of the emitted wavelengths by a charge coupled devise camera. Output/light intensity may be converted to electrical signal using appropriate software (e.g. GENOTYPER and SEQUENCE NAVIGATOR, Perkin Elmer) and the entire process from loading of samples to computer analysis and electronic data display may be computer controlled. Capillary electrophoresis is especially preferable for the sequencing of small pieces of DNA, which might be present in limited amounts in a particular sample.

In another embodiment of the invention, polynucleotide sequences or fragments thereof which encode Mnk, or fusion proteins or functional equivalents thereof, may be used in recombinant DNA molecules to direct expression of Mnk in appropriate host cells. Due to the inherent degeneracy of the genetic code, other DNA sequences, which encode substantially the same, or a functionally equivalent amino acid sequence may be produced and these sequences may be used to clone and express Mnk. As will be understood by those of skill in the art, it may be advantageous to produce Mnk-encoding nucleotide sequences possessing non-naturally occurring codons. For example, codons preferred by a particular prokaryotic or eukaryotic host can be selected to increase the rate of protein expression or to produce a recombinant RNA transcript having desirable properties, such as a half-life, which is longer than that of a transcript generated from the naturally occurring sequence. The nucleotide sequences of the present invention can be engineered using
methods generally known in the art in order to alter Mnk encoding sequences for a variety of reasons, including but not limited to, alterations which modify the cloning, processing, and/or expression of the gene product. DNA shuffling by random fragmentation and PCR reassembly of gene fragments and synthetic oligonucleotides may be used to engineer the nucleotide sequences. For example, site-directed mutagenesis may be used to insert new restriction sites, alter glycosylation patterns, change codon preference, produce splice variants, or introduce mutations, and so forth.

In another embodiment of the invention, natural, modified, or recombinant nucleic acid sequences encoding Mnk may be ligated to a heterologous sequence to encode a fusion protein. For example, to screen peptide libraries for inhibitors of Mnk activities, it may be useful to construct chimeric Mnk proteins that can be recognised by a commercially available antibodies. A fusion protein may also be engineered to contain a cleavage site located between the Mnk encoding sequence and the heterologous protein sequences, so that Mnk may be cleaved and purified away from the heterologous moiety. In another embodiment, sequences encoding Mnk may be synthesised, in whole or in part, using chemical methods well known in the art (see Caruthers et al. (1980) Nucl. Acids Res. Symp. Ser. 7:215-223, Horn et al. (1980) Nucl. Acids Res. Symp. Ser. 7:225-232). Alternatively, the proteins themselves may be produced using chemical methods to synthesise the amino acid sequence of Mnk, or a portion thereof. For example, peptide synthesis can be performed using various solid-phase techniques (Roberge et al. (1995) Science 269:202-204) and automated synthesis may be achieved, for example, using the ABI 431A peptide synthesiser (Perkin Elmer). The newly synthesised peptide may be substantially purified by preparative high performance liquid chromatography (e.g., Creighton, T. (1983) Proteins, Structures and Molecular Principles, WH Freeman and Co., New York, N.Y.). The composition of the synthetic peptides may be confirmed by amino acid analysis or sequencing (e.g., the Edman degradation procedure; Creighton,
supra). Additionally, the amino acid sequences of Mnk, or any part thereof, may be altered during direct synthesis and/or combined using chemical methods with sequences from other proteins, or any part thereof, to produce a variant polypeptide.

In order to express a biologically active Mnk, the nucleotide sequences encoding Mnk functional equivalents, may be inserted into appropriate expression vectors, i.e., a vector, which contains the necessary elements for the transcription and translation of the inserted coding sequence. Methods, which are well known to those skilled in the art, may be used to construct expression vectors containing sequences encoding Mnk and appropriate transcriptional and translational control elements. These methods include in vitro recombinant DNA techniques, synthetic techniques, and in vivo genetic recombination. Such techniques are described in Sambrook, J. et al. (1989) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Press, Plainview, N.Y., and Ausubel, F. M. et al. (1989) Current Protocols in Molecular Biology, John Wiley & Sons, New York, N.Y.

Regulatory elements include for example a promoter, an initiation codon, a stop codon, a mRNA stability regulatory element, and a polyadenylation signal. Expression of a polynucleotide can be assured by (i) constitutive promoters such as the Cytomegalovirus (CMV) promoter/enhancer region, (ii) tissue specific promoters such as the insulin promoter (see, Soria et al., 2000, Diabetes 49:157), SOX2 gene promoter (see Li et al., 1998, Curr. Biol. 8:971-4), Msi-1 promoter (see Sakakibara et al., 1997, J. Neuroscience 17:8300-8312), alpha-cardia myosin heavy chain promoter or human atrial natriuretic factor promoter (Klug et al., 1996, J. clin. Invest 98:216-24; Wu et al., 1989, J. Biol. Chem. 264:6472-79) or (iii) inducible promoters such as the tetracycline inducible system. Expression vectors can also contain a selection agent or marker gene that confers antibiotic resistance such as the neomycin, hygromycin or puromycin resistance
genes. These methods include in vitro recombinant DNA techniques, synthetic techniques, and in vivo genetic recombination. Such techniques are described in Sambrook, J. et al. (1989) Molecular Cloning, A Laboratory Manual, Cold Spring Harbor Press, Plainview, N.Y. and Ausubel, F.M. et al. (1989) Current Protocols in Molecular Biology, John Wiley & Sons, New York, N.Y. In a further embodiment of the invention, natural, modified or recombinant nucleic acid sequences encoding the proteins of the invention and homologous proteins may be ligated to a heterologous sequence to encode a fusion protein.

A variety of expression vector/host systems may be utilized to contain and express sequences encoding the proteins or fusion proteins. These include, but are not limited to, micro-organisms such as bacteria transformed with recombinant bacteriophage, plasmid or cosmid DNA expression vectors; yeast transformed with yeast expression vectors; insect cell systems infected with virus expression vectors (e.g., baculovirus, adenovirus, adeno-associated virus, lentivirus, retrovirus); plant cell systems transformed with virus expression vectors (e.g., cauliflower mosaic virus, CaMV; tobacco mosaic virus, TMV) or with bacterial expression vectors (e.g., Ti or PBR322 plasmids); or animal cell systems.

The "control elements" or "regulatory sequences" are those non-translated regions of the vectors, e.g. enhancers, promoters, 5' and 3' untranslated regions, which interact with host cellular proteins to carry out transcription and translation. Such elements may vary in their strength and specificity. Depending on the vector system and host utilised, any number of suitable transcription and translation elements, including constitutive and inducible promoters, may be used. For example, when cloning in bacterial systems, inducible promoters such as the hybrid lacZ promoter of the BLUESCRIPT phagemid (Stratagene, LaJolla, Calif.) or PSPORT1 plasmid (Gibco BRL) and the like may be used. The baculovirus polyhedrin promoter may be used in insect cells. Promoters and enhancers derived from the genomes of plant
cells (e.g., heat shock, RUBISCO; and storage protein genes) or from plant viruses (e.g., viral promoters and leader sequences) may be cloned into the vector. In mammalian cell systems, promoters from mammalian genes or from mammalian viruses are preferable. If it is necessary to generate a cell line that contains multiple copies of the sequences encoding Mnk, vectors based on SV40 or EBV may be used with an appropriate selectable marker.

In bacterial systems, a number of expression vectors may be selected depending upon the use intended for Mnk. For example, when large quantities of Mnk are needed for the induction of antibodies, vectors, which direct high level expression of fusion proteins that are readily purified, may be used. Such vectors include, but are not limited to, the multifunctional E. coli cloning and expression vectors such as the BLUESCRIPT phagemid (Stratagene), in which the sequence encoding Mnk may be ligated into the vector in frame with sequences for the amino-terminal Met and the subsequent 7 residues of β-galactosidase so that a hybrid protein is produced; pIN vectors (Van Haeke, G. and S. M. Schuster (1989) J. Biol. Chem. 264:5503-5509); and the like. Vectors of the pGEX series (Amersham Biosciences, Uppsala, Sweden) may also be used to express foreign polypeptides as fusion proteins with Glutathione S-Transferase (GST). In general, such fusion proteins are soluble and can easily be purified from lysed cells by adsorption to glutathione-agarose beads followed by elution in the presence of free glutathione. Proteins made in such systems may be designed to include heparin, thrombin, or factor XA protease cleavage sites so that the cloned polypeptide of interest can be released from the GST moiety at will. In the yeast, Saccharomyces cerevisiae, a number of vectors containing constitutive or inducible promoters such as alpha factor, alcohol oxidase, and PGH may be used. For reviews, see Ausubel et al., (supra) and Grant et al. (1987) Methods Enzymol. 153:516-544.
In cases where plant expression vectors are used, the expression of sequences encoding Mnk may be driven by any of a number of promoters. For example, viral promoters such as the 35S and 19S promoters of CaMV may be used alone or in combination with the omega leader sequence from TMV (Takamatsu, N. (1987) EMBO J. 6:307-311). Alternatively, plant promoters such as the small subunit of RUBISCO or heat shock promoters may be used (Coruzzi, G. et al. (1984) EMBO J. 3:1671-1680; Broglie, R. et al. (1984) Science 224:838-843; and Winter, J. et al. (1991) Results Probl. Cell Differ. 17:85-105). These constructs can be introduced into plant cells by direct DNA transformation or pathogen-mediated transfection. Such techniques are described in a number of generally available reviews (see, for example, Hobbs, S. or Murry, L. E. in McGraw Hill Yearbook of Science and Technology (1992) McGraw Hill, New York, N.Y.; pp. 191-196).

An insect system may also be used to express Mnk. For example, in one such system, Autographa californica nuclear polyhedrosis virus (AcNPV) is used as a vector to express foreign genes in Spodoptera frugiperda cells or in Trichoplusia larvae. The sequences encoding Mnk may be cloned into a non-essential region of the virus, such as the polyhedrin gene, and placed under control of the polyhedrin promoter. Successful insertions of Mnk will render the polyhedrin gene inactive and produce recombinant virus lacking coat protein. The recombinant viruses may then be used to infect, for example, S. frugiperda cells of Trichoplusia larvae in which Mnk may be expressed (Engelhard, E. K. et al. (1994) Proc. Nat. Acad. Sci. 91:3224-3227).

In mammalian host cells, a number of viral-based expression systems may be utilised. In cases where an adenovirus is used as an expression vector, sequences encoding Mnk may be ligated into an adenovirus transcription/translation complex consisting of the late promoter and tripartite leader sequence. Insertion in a non-essential E1 or E3 region of
the viral genome may be used to obtain viable viruses which are capable of expressing Mnk in infected host cells (Logan, J. and Shenk, T. (1984) Proc. Natl. Acad. Sci. 81:3655-3659). In addition, transcription enhancers, such as the Rous sarcoma virus (RSV) enhancer, may be used to increase expression in mammalian host cells.

Specific initiation signals may also be used to achieve more efficient translation of sequences encoding Mnk. Such signals include the ATG initiation codon and adjacent sequences. In cases where sequences encoding Mnk, its initiation codons, and upstream sequences are inserted into the appropriate expression vector, no additional transcriptional or translational control signals may be needed. However, in cases where only coding sequence, or a portion thereof, is inserted, exogenous translational control signals including the ATG initiation codon should be provided. Furthermore, the initiation codon should be in the correct reading frame to ensure translation of the entire insert. Exogenous translational elements and initiation codons may be of various origins, both natural and synthetic. The efficiency of expression may be enhanced by the inclusion of enhancers which are appropriate for the particular cell system which is used, such as those described in the literature (Scharf, D. et al. (1994) Results Probl. Cell Differ. 20:125-162).

In addition, a host cell strain may be chosen for its ability to modulate the expression of the inserted sequences or to process the expressed protein in the desired fashion. Such modifications of the polypeptide include, but are not limited to, acetylation, carboxylation, glycosylation, phosphorylation, lipidation, and acylation. Post-translational processing which cleaves a "prepro" form of the protein may also be used to facilitate correct insertion, folding and/or function. Different host cells such as CHO, HeLa, MDCK, HEK293, and WI38, which have specific cellular machinery and characteristic mechanisms for such post-translational activities, may
be chosen to ensure the correct modification and processing of the foreign protein.

For long-term, high-yield production of recombinant proteins, stable expression is preferred. For example, cell lines, which stably express MnK may be transformed using expression vectors which may contain viral origins of replication and/or endogenous expression elements and a selectable marker gene on the same or on a separate vector. Following the introduction of the vector, cells may be allowed to grow for 1-2 days in an enriched media before they are switched to selective media. The purpose of the selectable marker is to confer resistance to selection, and its presence allows growth and recovery of cells, which successfully express the introduced sequences. Resistant clones of stably transformed cells may be proliferated using tissue culture techniques appropriate to the cell type. Any number of selection systems may be used to recover transformed cell lines. These include, but are not limited to, the herpes simplex virus thymidine kinase (Wigler, M. et al. (1977) Cell 11:223-32) and adenine phosphoribosyltransferase (Lowy, I. et al. (1980) Cell 22:817-23) genes, which can be employed in tk- or aprt-cells, respectively. Also, antimetabolite, antibiotic or herbicide resistance can be used as the basis for selection; for example, dhfr which confers resistance to methotrexate (Wigler, M. et al. (1980) Proc. Natl. Acad. Sci. 77:3567-70); npt, which confers resistance to the aminoglycosides neomycin and G-418 (Colbere-Garapin, F. et al (1981) J. Mol. Biol. 150:1-14) and als or pat, which confer resistance to chlorsulfuron and phosphinotricin acetyltransferase, respectively (Murry, supra). Additional selectable genes have been described, for example, trpB, which allows cells to utilise indole in place of tryptophan, or hisD, which allows cells to utilise histinol in place of histidine (Hartman, S. C. and R. C. Mulligan (1988) Proc. Natl. Acad. Sci. 85:8047-51). Recently, the use of visible markers has gained popularity with such markers as anthocyanins, β-glucuronidase and its substrate GUS, and luciferase and its substrate luciferin, being widely used
not only to identify transformants, but also to quantify the amount of transient or stable protein expression attributable to a specific vector system (Rhodes, C. A. et al. (1995) Methods Mol. Biol. 55:121-131).

In vivo, the enzymatic kinase activity of the unmodified polypeptides of Mnk towards a substrate can be enhanced by appropriate stimuli, triggering the phosphorylation of Mnk. This may be induced in the natural context by extracellular or intracellular stimuli, such as signaling molecules or environmental influences. One may generate a system containing activated Mnk, may it be an organism, a tissue, a culture of cells or cell-free environment, by exogenously applying this stimulus or by mimicking this stimulus by a variety of the techniques, some of them described further below. A system containing activated Mnk may be produced (i) for the purpose of diagnosis, study, prevention, and treatment of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs, (ii) for the purpose of identifying or validating therapeutic candidate agents, pharmaceuticals or drugs that influence the genes of the invention or their encoded polypeptides, (iii) for the purpose of generating cell lysates containing activated polypeptides encoded by the genes of the invention, (iv) for the purpose of isolating from this source activated polypeptides encoded by the genes of the invention.

In one embodiment of the invention, one may produce activated Mnk independent of the natural stimuli for the above said purposes by, for example, but not limited to, (i) an agent that mimics the natural stimulus; (ii) an agent, that acts downstream of the natural stimulus, such as
activators of the MAP kinase pathway, phorbol ester, anisomycin, constitutive active alleles of the MAP kinase kinase kinases, of the MAP kinase kinases, of the MAP kinase or Mnk itself as they are described or may be developed; (iii) by introduction of single or multiple amino acid substitutions, deletions or insertions within the sequence of Mnk to yield constitutive active forms; (iv) by the use of isolated fragments of Mnk. In addition, one may generate enzymatically active Mnk in an ectopic system, prokaryotic or eukaryotic, in vivo or in vitro, by co-transferring the activating components to this system. These could be, for example, but not limited to, components of the MAP kinase pathway such as constitutive active alleles of the MAP kinase kinases Mek1 or Mkk6, together with the MAP kinases ERK1 or ERK2 or the p38 MAPK isoforms. For example, one may activate isolated Mnk protein in solution with a mutant polypeptide of Mek1 containing the amino acid substitutions S218D and S222E together with isolated ERK2 kinase in the presence of 1.0 mM adenosine triphosphate and suitable buffer conditions such as 50 mM N-(2-Hydroxyethyl)-piperazine-N’-(2-ethanesulfonic acid)/potassium hydroxide pH 7.4, 5 mM magnesium chloride, 0.5 mM dithiothreitol (see Figure 14).

Although the presence/absence of marker gene expression suggests that the gene of interest is also present, its presence and expression may need to be confirmed. For example, if the sequences encoding Mnk are inserted within a marker gene sequence, recombinant cells containing sequences encoding Mnk can be identified by the absence of marker gene function. Alternatively, a marker gene can be placed in tandem with sequences encoding Mnk under the control of a single promoter. Expression of the marker gene in response to induction or selection usually indicates expression of the tandem gene as well. Alternatively, host cells, which contain the nucleic acid sequences encoding Mnk and express Mnk, may be identified by a variety of procedures known to those of skill in the art. These procedures include, but are not limited to, DNA-DNA, or DNA-RNA
hybridisation and protein bioassay or immunoassay techniques, which include membrane, solution, or chip based technologies for the detection and/or quantification of nucleic acid or protein.

The presence of polynucleotide sequences encoding Mnk can be detected by DNA-DNA or DNA-RNA hybridisation or amplification using probes or portions or fragments of polynucleotides encoding Mnk. Nucleic acid amplification based assays involve the use of oligonucleotides or oligomers based on the sequences encoding Mnk to detect transformants containing DNA or RNA encoding Mnk. As used herein "oligonucleotides" or "oligomers" refer to a nucleic acid sequence of at least about 10 nucleotides and as many as about 60 nucleotides, preferably about 15 to 30 nucleotides, and more preferably about 20-25 nucleotides, which can be used as a probe or amplimer.

A variety of protocols for detecting and measuring the expression of Mnk, using either polyclonal or monoclonal antibodies specific for the protein are known in the art. Examples include enzyme-linked immunosorbent assay (ELISA), radioimmunoassay (RIA), and fluorescence activated cell sorting (FACS). A two-site, monoclonal-based immunoassay utilising monoclonal antibodies reactive to two non-interfering epitopes on Mnk is preferred, but a competitive binding assay may be employed. These and other assays are described, among other places, in Hampton, R. et al. (1990; Serological Methods, a Laboratory Manual, APS Press, St Paul, Minn.) and Maddox, D. E. et al. (1983; J. Exp. Med. 158:1211-1216).

A wide variety of labels and conjugation techniques are known by those skilled in the art and may be used in various nucleic acid and amino acid assays. Means for producing labelled hybridisation or PCR probes for detecting sequences related to polynucleotides encoding Mnk include oligo-labelling, nick translation, end-labelling or PCR amplification using a labelled nucleotide.
Alternatively, the sequences encoding Mnk, or any portions thereof may be cloned into a vector for the production of an mRNA probe. Such vectors are known in the art, are commercially available, and may be used to synthesise RNA probes in vitro by addition of an appropriate RNA polymerase such as T7, T3, or SP6 and labelled nucleotides. These procedures may be conducted using a variety of commercially available kits (Pharmacia & Upjohn, (Kalamazoo, Mich.); Promega (Madison Wis.); and U.S. Biochemical Corp., (Cleveland, Ohio).

Suitable reporter molecules or labels, which may be used, include radionuclides, enzymes, fluorescent, chemiluminescent, or chromogenic agents as well as substrates, co-factors, inhibitors, magnetic particles, and the like.

Host cells transformed with nucleotide sequences encoding Mnk may be cultured under conditions suitable for the expression and recovery of the protein from cell culture. The protein produced by a recombinant cell may be secreted or contained intracellularly depending on the sequence and/or the vector used. As will be understood by those of skill in the art, expression vectors containing polynucleotides which encode Mnk may be designed to contain signal sequences, which direct secretion of Mnk through a prokaryotic or eukaryotic cell membrane. Other recombinant constructions may be used to join sequences encoding Mnk to nucleotide sequence encoding a polypeptide domain, which will facilitate purification of soluble proteins. Such purification facilitating domains include, but are not limited to, metal chelating peptides such as histidine-tryptophan modules that allow purification on immobilised metals, protein A domains that allow purification on immobilised immunoglobulin, and the domain utilised in the FLAG extension/affinity purification system (Immunex Corp., Seattle, Wash.) The inclusion of cleavable linker sequences such as those specific for Factor XA or Enterokinase (Invitrogen, San Diego, Calif.) between the purification domain and Mnk may be used to facilitate
purification. One such expression vector provides for expression of a fusion protein containing Mnk and a nucleic acid encoding 6 histidine residues preceding a Thioredoxin or an Enterokinase cleavage site. The histidine residues facilitate purification on IMAC (immobilised metal ion affinity chromatography as described in Porath, J. et al. (1992; Prot. Exp. Purif. 3: 263-281)) while the Enterokinase cleavage site provides a means for purifying Mnk from the fusion protein. A discussion of vectors which contain fusion proteins is provided in Kroll, D. J. et al. (1993; DNA Cell Biol. 12:441-453). In addition to recombinant production, fragments of Mnk may be produced by direct peptide synthesis using solid-phase techniques (Merrifield J. (1963) J. Am. Chem. Soc. 85:2149-2154). Protein synthesis may be performed using manual techniques or by automation. Automated synthesis may be achieved, for example, using Applied Biosystems 431A peptide synthesiser (Perkin Elmer). Various fragments of Mnk may be chemically synthesised separately and combined using chemical methods to produce the full length molecule.

Diagnostics and Therapeutics

The data disclosed in this invention show that the nucleic acids and proteins of the invention and effector molecules thereof are useful in diagnostic and therapeutic applications implicated, for example but not limited to, in metabolic disorders like obesity, diabetes, eating disorders, wasting syndromes (cachexia), pancreatic dysfunctions, arteriosclerosis, coronary artery disease (CAD), and other diseases and disorders as described above. Hence, diagnostic and therapeutic uses for the Mnk proteins of the invention are, for example but not limited to, the following: (i) protein therapeutic, (ii) small molecule drug target, (iii) antibody target (therapeutic, diagnostic, drug targeting/cytotoxic antibody), (iv) diagnostic and/or prognostic marker, (v) gene therapy (gene delivery/gene ablation), (vi) research tools, and (vii) tissue regeneration in vitro and in vivo
(regeneration for all these tissues and cell types composing these tissues and cell types derived from these tissues).

The nucleic acids and proteins of the invention are useful in diagnostic and therapeutic applications implicated in various diseases and disorders described above and/or other pathologies and disorders. For example, but not limited to, cDNAs encoding the Mnk proteins of the invention and particularly their human homologues may be useful in gene therapy, and the Mnk proteins of the invention and particularly their human homologues may be useful when administered to a subject in need thereof. By way of non-limiting example, the compositions of the present invention will have efficacy for treatment of patients suffering from, for example, but not limited to, in metabolic disorders like obesity, diabetes, eating disorders, wasting syndromes (cachexia), pancreatic dysfunctions, arteriosclerosis, coronary artery disease (CAD), and other diseases and disorders, particularly as described above.

The nucleic acid(s) encoding the Mnk protein(s) of the invention, or fragments thereof, may further be useful in diagnostic applications, wherein the presence or amount of the nucleic acids or the proteins are to be assessed. These materials are further useful in the generation of antibodies that bind immunospecifically to the novel substances of the invention for use in therapeutic or diagnostic methods.

For example, in one aspect, antibodies which are specific for Mnk may be used directly as an antagonist, or indirectly as a targeting or delivery mechanism for bringing a pharmaceutical agent to cells or tissue which express Mnk. The antibodies may be generated using methods that are well known in the art. Such antibodies may include, but are not limited to, polyclonal, monoclonal, chimerical, single chain, Fab fragments, and fragments produced by a Fab expression library. Neutralising antibodies,
(i.e., those which inhibit dimer formation) are especially preferred for therapeutic use.

For the production of antibodies, various hosts including goats, rabbits, rats, mice, humans, and others, may be immunised by injection with Mnk any fragment or oligopeptide thereof which has immunogenic properties. Depending on the host species, various adjuvants may be used to increase immunological response. Such adjuvants include, but are not limited to, Freund's, mineral gels such as aluminium hydroxide, and surface active substances such as lysolecithin, pluronic polyols, polyanions, peptides, oil emulsions, keyhole limpet hemocyanin, and dinitrophenol. Among adjuvants used in human, BCG (Bacille Calmette-Guerin) and Corynebacterium parvum are especially preferable. It is preferred that the peptides, fragments, or oligopeptides used to induce antibodies to Mnk have an amino acid sequence consisting of at least five amino acids, and more preferably at least 10 amino acids. It is preferable that they are identical to a portion of the amino acid sequence of the natural protein, and they may contain the entire amino acid sequence of a small, naturally occurring molecule. Short stretches of Mnk amino acids may be fused with those of another protein such as keyhole limpet hemocyanin and antibody produced against the chimeric molecule.


In addition, techniques developed for the production of "chimeric antibodies", the splicing of mouse antibody genes to human antibody


Antibody fragments, which contain specific binding sites for Mnk, may also be generated. For example, such fragments include, but are not limited to proteolytic fragments, e.g. the F(\(ab'\))\(_2\) fragments which can be produced by Pepsin digestion of the antibody molecule and the Fab fragments which can be generated by reducing the disulfide bridges of F(\(ab'\))\(_2\) fragments. Alternatively, recombinant fragments may be generated. For example, Fab expression libraries may be constructed to allow rapid and easy identification of monoclonal Fab fragments with the desired specificity (Huse, W. D. et al. (1989) Science 254:1275-1281).

Various immunoassays may be used for screening to identify antibodies having the desired specificity. Numerous protocols for competitive binding and immunoradiometric assays using either polyclonal or monoclonal antibodies with established specificities are well known in the art. Such immunoassays typically involve the measurement of complex formation between Mnk and its specific antibody. A two-site, monoclonal-based immunoassay utilising monoclonal antibodies reactive to two
non-interfering Mnk epitopes is preferred, but a competitive binding assay may also be employed (Maddox, supra).

In another embodiment of the invention, the Mnk polynucleotides or any fragment thereof, or nucleic acid effector molecules, aptamers, anti-sense molecules, ribozymes or RNAi molecules, may be used for therapeutic purposes. In one aspect, aptamers, i.e. nucleic acid molecules, which are capable of binding to a Mnk protein and modulating its activity, may be generated by a screening and selection procedure involving the use of combinational nucleic acid libraries.

In a further aspect, antisense molecules to the polynucleotide encoding Mnk may be used in situations in which it would be desirable to block the transcription of the mRNA. In particular, cells may be transformed with sequences complementary to polynucleotides encoding Mnk. Thus, antisense molecules may be used to modulate Mnk activity, or to achieve regulation of gene function. Such technology is now well know in the art, and sense or antisense oligomers or larger fragments, can be designed from various locations along the coding or control regions of sequences encoding Mnk. Expression vectors derived from retroviruses, adenovirus, herpes or vaccinia viruses, or from various bacterial plasmids may be used for delivery of nucleotide sequences to the targeted organ, tissue or cell population. Methods, which are well known to those skilled in the art, can be used to construct recombinant vectors, which will express antisense molecules complementary to the polynucleotides of the gene encoding Mnk. These techniques are described both in Sambrook et al. (supra) and in Ausubel et al. (supra). Genes encoding Mnk can be turned off by transforming a cell or tissue with expression vectors which express high levels of polynucleotide or fragment thereof which encodes Mnk. Such constructs may be used to introduce untranslatable sense or antisense sequences into a cell. Even in the absence of integration into the DNA, such vectors may continue to transcribe RNA molecules until they are
disabled by endogenous nucleases. Transient expression may last for a month or more with a non-replicating vector and even longer if appropriate replication elements are part of the vector system.

As mentioned above, modifications of gene expression can be obtained by designing antisense molecules, e.g. DNA, RNA, or nucleic acid analogues such as PNA, to the control regions of the gene encoding Mnk, i.e., the promoters, enhancers, and introns. Oligonucleotides derived from the transcription initiation site, e.g., between positions -10 and +10 from the start site, are preferred. Similarly, inhibition can be achieved using "triple helix" base-pairing methodology. Triple helix pairing is useful because it cause inhibition of the ability of the double helix to open sufficiently for the binding of polymerases, transcription factors, or regulatory molecules. Recent therapeutic advances using triplex DNA have been described in the literature (Gee, J. E. et al. (1994) In; Huber, B. E. and B. I. Carr, Molecular and Immunologic Approaches, Futura Publishing Co., Mt. Kisco, N.Y.). The antisense molecules may also be designed to block translation of mRNA by preventing the transcript from binding to ribosomes.

Ribozymes, enzymatic RNA molecules, may also be used to catalyse the specific cleavage of RNA. The mechanism of ribozyme action involves sequence-specific hybridisation of the ribozyme molecule to complementary target RNA, followed by endonucleolytic cleavage. Examples, which may be used, include engineered hammerhead motif ribozyme molecules that can be specifically and efficiently catalyse endonucleolytic cleavage of sequences encoding Mnk. Specific ribozyme cleavage sites within any potential RNA target are initially identified by scanning the target molecule for ribozyme cleavage sites which include the following sequences: GUA, GUU, and GUC. Once identified, short RNA sequences of between 15 and 20 ribonucleotides corresponding to the region of the target gene containing the cleavage site may be evaluated for secondary structural features which may render the oligonucleotide inoperable. The suitability of
candidate targets may also be evaluated by testing accessibility to hybridisation with complementary oligonucleotides using ribonuclease protection assays.

Effectors molecules, e.g. antisense molecules and ribozymes of the invention may be prepared by any method known in the art for the synthesis of nucleic acid molecules. These include techniques for chemically synthesising oligonucleotides such as solid phase phosphoramidite chemical synthesis. Alternatively, RNA molecules may be generated by in vitro and in vivo transcription of DNA sequences encoding Mnk. Such DNA sequences may be incorporated into a variety of vectors with suitable RNA polymerase promoters such as T7 or SP6. Alternatively, these cDNA constructs that synthesise antisense RNA constitutively or inducibly can be introduced into cell lines, cells, or tissues. RNA molecules may be modified to increase intracellular stability and half-life. Possible modifications include, but are not limited to, the addition of flanking sequences at the 5' and/or 3' ends of the molecule or the use of phosphorothioate or 2' O-methyl rather than phosphodiesterase linkages within the backbone of the molecule. This concept is inherent in the production of PNAs and can be extended in all of these molecules by the inclusion of non-traditional bases such as inosine, queosine, and wybutosine, as well as acetyl-, methyl-, thio-, and similarly modified forms of adenine, cytidine, guanine, thymine, and uridine which are not as easily recognised by endogenous endonucleases.

The activity of Mnk proteins can be assayed for example by in vitro kinase assays, as described by Tschopp et al., 2000, supra or any other suitable assay principle as described below. As inhibitor of Mnk in this assay, a staurosporine derivative such as CGP57380 or CGP052088 can be used, as described by Tschopp et al., 2000, supra or Knauf et al., 2001, supra. As negative control, the compound CGP52428 which is inactive against Mnk, but displays a similar cytotoxicity as CGP052088, or any other
chemical entities with kinase inhibitory activity with exception of activity against Mnk may be used. Moreover, derivatives of CGP57380 can be assayed for activity against Mnk and are substances for the treatment, prophylaxis, and diagnosis of metabolic diseases as mentioned above. Derivatives of CGP57380 could for example be generated by modification through conventional chemical, physical and biochemical means, and may be used to produce combinatorial libraries. They may be subjected to directed or random chemical modifications, such as acylation, alkylation, esterification, amidification, etc. to produce structural analogs.

Further, the invention relates to the use of Mnk kinase inhibitors or activators for the treatment, prophylaxis or diagnosis of metabolic diseases as mentioned above. Preferably, but not exclusively, the Mnk kinase inhibitors are staurosporine or pyrazole derivatives. Examples of pyrazole derivatives are described in EP-A-0 819 129 which is herein incorporated by reference. Since CGP57380 is not cytotoxic up to 30μM, this substance may be preferably used to inhibit kinase activity, preferably Mnk2, and used as substance for the treatment, prophylaxis, and diagnosis of metabolic diseases as mentioned above.

Many methods for introducing vectors into cells or tissues are available and equally suitable for use in vivo, in vitro, and ex vivo. For ex vivo therapy, vectors may be introduced into stem cells taken from the patient and clonally propagated for autologous transplant back into that same patient. Delivery by transfection and by liposome injections may be achieved using methods, which are well known in the art. Any of the therapeutic methods described above may be applied to any suitable subject including, for example, mammals such as dogs, cats, cows, horses, rabbits, monkeys, and most preferably, humans.

An additional embodiment of the invention relates to the administration of a pharmaceutical composition, in conjunction with a pharmaceutically
acceptable carrier, for any of the therapeutic effects discussed above. Such pharmaceutical compositions may consist of Mnk, antibodies to Mnk, mimetics, agonists, antagonists, or inhibitors of Mnk. The compositions may be administered alone or in combination with at least one other agent, such as stabilising compound, which may be administered in any sterile, biocompatible pharmaceutical carrier, including, but not limited to, saline, buffered saline, dextrose, and water. The compositions may be administered to a patient alone, or in combination with other agents, drugs or hormones. The pharmaceutical compositions utilised in this invention may be administered by any number of routes including, but not limited to, oral, intravenous, intramuscular, intra-arterial, intramedullary, intrathecal, intraventricular, transdermal, subcutaneous, intraperitoneal, intranasal, enteral, topical, sublingual, or rectal means.

In addition to the active ingredients, these pharmaceutical compositions may contain suitable pharmaceutically-acceptable carriers comprising excipients and auxiliaries, which facilitate processing of the active compounds into preparations which, can be used pharmaceutically. Further details on techniques for formulation and administration may be found in the latest edition of Remington's Pharmaceutical Sciences (Maack Publishing Co., Easton, Pa.). Pharmaceutical compositions for oral administration can be formulated using pharmaceutically acceptable carriers well known in the art in dosages suitable for oral administration. Such carriers enable the pharmaceutical compositions to be formulated as tablets, pills, dragees, capsules, liquids, gels, syrups, slurries, suspensions, and the like, for ingestion by the patient.

The pharmaceutical compositions of the present invention may be manufactured in a manner that is known in the art, e.g., by means of conventional mixing, dissolving, granulating, dragee-making, levigating, emulsifying, encapsulating, entrapping, or lyophilising processes. The pharmaceutical composition may be provided as a salt and can be formed
with many acids, including but not limited to, hydrochloric, sulphuric, acetic, lactic, tartaric, malic, succinic, etc. After pharmaceutical compositions have been prepared, they can be placed in an appropriate container and labelled for treatment of an indicated condition. For administration of Mnk, such labelling would include amount, frequency, and method of administration.

Pharmaceutical compositions suitable for use in the invention include compositions wherein the active ingredients are contained in an effective amount to achieve the intended purpose. The determination of an effective dose is well within the capability of those skilled in the art. For any compounds, the therapeutically effective doses can be estimated initially either in cell culture assays, e.g., of preadipocyte cell lines, or in animal models, usually mice, rabbits, dogs, or pigs. The animal model may also be used to determine the appropriate concentration range and route of administration. Such information can then be used to determine useful doses and routes for administration in humans. A therapeutically effective dose refers to that amount of active ingredient, for example Mnk fragments thereof, antibodies of Mnk, to treat a specific condition. Therapeutic efficacy and toxicity may be determined by standard pharmaceutical procedures in cell cultures or experimental animals, e.g., ED50 (the dose therapeutically effective in 50% of the population) and LD50 (the dose lethal to 50% of the population). The dose ratio between therapeutic and toxic effects is the therapeutic index, and it can be expressed as the ratio, LD50/ED50. Pharmaceutical compositions, which exhibit large therapeutic indices, are preferred. The data obtained from cell culture assays and animal studies is used in formulating a range of dosage for human use. The dosage contained in such compositions is preferably within a range of circulating concentrations that include the ED50 with little or no toxicity. The dosage varies within this range depending upon the dosage from employed, sensitivity of the patient, and the route of administration. The exact dosage will be determined by the practitioner, in light of factors
related to the subject that requires treatment. Dosage and administration are adjusted to provide sufficient levels of the active moiety or to maintain the desired effect. Factors, which may be taken into account, include the severity of the disease state, general health of the subject, age, weight, and gender of the subject, diet, time and frequency of administration, drug combination(s), reaction sensitivities, and tolerance/response to therapy. Long-acting pharmaceutical compositions may be administered every 3 to 4 days, every week, or once every two weeks depending on half-life and clearance rate of the particular formulation. Normal dosage amounts may vary from 0.1 to 100,000 micrograms, up to a total dose of about 1 g, depending upon the route of administration. Guidance as to particular dosages and methods of delivery is provided in the literature and generally available to practitioners in the art. Those skilled in the art employ different formulations for nucleotides than for proteins or their inhibitors. Similarly, delivery of polynucleotides or polypeptides will be specific to particular cells, conditions, locations, etc.

In another embodiment, antibodies which specifically bind Mnk may be used for the diagnosis of conditions or diseases characterised by or associated with over- or underexpression of Mnk, or in assays to monitor patients being treated with Mnk, agonists, antagonists or inhibitors. The antibodies useful for diagnostic purposes may be prepared in the same manner as those described above for therapeutics. Diagnostic assays for Mnk include methods, which utilise the antibody and a label to detect Mnk in human body fluids or extracts of cells or tissues. The antibodies may be used with or without modification, and may be labelled by joining them, either covalently or non-covalently, with a reporter molecule. A wide variety of reporter molecules, which are known in the art may be used several of which are described above.

A variety of protocols including ELISA, RIA, and FACS for measuring Mnk are known in the art and provide a basis for diagnosing altered or abnormal
levels of Mnk expression. Normal or standard values for Mnk expression are established by combining body fluids or cell extracts taken from normal mammalian subjects, preferably human, with antibody to Mnk under conditions suitable for complex formation. The amount of standard complex formation may be quantified by various methods, but preferably by photometric, means. Quantities of Mnk expressed in control and disease samples e.g. from biopsied tissues are compared with the standard values. Deviation between standard and subject values establishes the parameters for diagnosing disease. Analysis of Mnk expression can also be performed by determination of Mnk activity in assay formats well known in the art and described in more detail below.

In another embodiment of the invention, the polynucleotides specific for Mnk may be used for diagnostic purposes. The polynucleotides, which may be used, include oligonucleotide sequences, antisense RNA and DNA molecules, and PNAs. The polynucleotides may be used to detect and quantitate gene expression in biopsied tissues in which expression of Mnk may be correlated with disease. The diagnostic assay may be used to distinguish between absence, presence, and excess expression of Mnk, and to monitor regulation of Mnk levels during therapeutic intervention.

In one aspect, hybridisation with PCR probes which are capable of detecting polynucleotide sequences, including genomic sequences, encoding Mnk and/or closely related molecules, may be used to identify nucleic acid sequences which encode Mnk. The specificity of the probe, whether it is made from a highly specific region, e.g., unique nucleotides in the 5’ regulatory region, or a less specific region, e.g., especially in the 3’ coding region, and the stringency of the hybridisation or amplification (maximal, high, intermediate, or low) will determine whether the probe identifies only naturally occurring sequences encoding Mnk, alleles, or related sequences. Probes may also be used for the detection of related sequences, and should preferably contain at least 50% of the nucleotides
from any of the Mnk encoding sequences. The hybridisation probes of the subject invention may be DNA or RNA and derived from the nucleotide sequence of AF237775, NM_017572.1, NM_003684.2, or AB000409.1 or from a genomic sequence including promoter, enhancer elements, and introns of the naturally occurring Mnk. Means for producing specific hybridisation probes for DNAs encoding Mnk include the cloning of nucleic acid sequences encoding Mnk derivatives into vectors for the production of mRNA probes. Such vectors are known in the art, commercially available, and may be used to synthesise RNA probes in vitro by means of the addition of the appropriate RNA polymerases and the appropriate labelled nucleotides. Hybridisation probes may be labelled by a variety of reporter groups, for example, radionuclides such as $^{32}\text{P}$ or $^{35}\text{S}$, or enzymatic labels, such as alkaline phosphatase coupled to the probe via avidin/biotin coupling systems, and the like.

Polynucleotide sequences encoding Mnk may be used for the diagnosis of conditions or diseases, which are associated with expression of Mnk. Examples of such conditions or diseases include, but are not limited to, pancreatic diseases and disorders, including diabetes. Polynucleotide sequences encoding Mnk may also be used to monitor the progress of patients receiving treatment for pancreatic diseases and disorders, including diabetes. The polynucleotide sequences encoding Mnk may be used in Southern or Northern analysis, dot blot, or other membrane-based technologies; in PCR technologies; or in dip stick, pin, ELISA or chip assays utilising fluids or tissues from patient biopsies to detect altered Mnk expression. Such qualitative or quantitative methods are well known in the art.

In a particular aspect, the nucleotide sequences encoding Mnk may be useful in assays that detect activation or induction of various metabolic diseases and disorders, including obesity, diabetes, eating disorders, wasting syndromes (cachexia), pancreatic dysfunctions, arteriosclerosis,
coronary artery disease (CAD), disorders related to ROS production, and neurodegenerative diseases. The nucleotide sequences encoding Mnk may be labelled by standard methods, and added to a fluid or tissue sample from a patient under conditions suitable for the formation of hybridisation complexes. After a suitable incubation period, the sample is washed and the signal is quantitated and compared with a standard value. The presence of altered levels of nucleotide sequences encoding Mnk in the sample compared to a control sample indicates the presence of the associated disease. Such assays may also be used to evaluate the efficacy of a particular therapeutic treatment regimen in animal studies, in clinical trials, or in monitoring the treatment of an individual patient.

In order to provide a basis for the diagnosis of disease associated with expression of Mnk, a normal or standard profile for expression is established. This may be accomplished by combining body fluids or cell extracts taken from normal subjects, either animal or human, with a sequence, or a fragment thereof, which encodes Mnk, under conditions suitable for hybridisation or amplification. Standard hybridisation may be quantified by comparing the values obtained from normal subjects with those from an experiment where a known amount of a substantially purified polynucleotide is used. Standard values obtained from normal samples may be compared with values obtained from samples from patients who are symptomatic for disease. Deviation between standard and subject values is used to establish the presence of disease. Once disease is established and a treatment protocol is initiated, hybridisation assays may be repeated on a regular basis to evaluate whether the level of expression in the patient begins to approximate that, which is observed in the normal patient. The results obtained from successive assays may be used to show the efficacy of treatment over a period ranging from several days to months.
With respect to metabolic diseases and disorders, including obesity, diabetes, eating disorders, wasting syndromes (cachexia), pancreatic dysfunctions, arteriosclerosis, coronary artery disease (CAD), disorders related to ROS production, and neurodegenerative diseases presence of a relatively high amount of transcript in biopsied tissue from an individual may indicate a predisposition for the development of the disease, or may provide a means for detecting the disease prior to the appearance of actual clinical symptoms. A more definitive diagnosis of this type may allow health professionals to employ preventative measures or aggressive treatment earlier thereby preventing the development or further progression of the pancreatic diseases and disorders. Additional diagnostic uses for oligonucleotides designed from the sequences encoding Mnk may involve the use of PCR. Such oligomers may be chemically synthesised, generated enzymatically, or produced from a recombinant source. Oligomers will preferably consist of two nucleotide sequences, one with sense orientation (5’fwdarw.3’) and another with antisense (3’.rarw.5’), employed under optimised conditions for identification of a specific gene or condition. The same two oligomers, nested sets of oligomers, or even a degenerate pool of oligomers may be employed under less stringent conditions for detection and/or quantification of closely related DNA or RNA sequences.

Methods which may also be used to quantitate the expression of Mnk include radiolabelling or biotinylating nucleotides, coamplification of a control nucleic acid, and standard curves onto which the experimental results are interpolated (Melby, P. C. et al. (1993) J. Immunol. Methods, 159:235-244; Duplaa, C. et al. (1993) Anal. Biochem. 212:229-236). The speed of quantification of multiple samples may be accelerated by running the assay in an ELISA format where the oligomer of interest is presented in various dilutions and a spectrophotometric or colorimetric response gives rapid quantification.
In another embodiment of the invention, the nucleic acid Mnk sequences may also be used to generate hybridisation probes, which are useful for mapping the naturally occurring genomic sequence. The sequences may be mapped to a particular chromosome or to a specific region of the chromosome using well known techniques. Such techniques include FISH, FACS, or artificial chromosome constructions, such as yeast artificial chromosomes, bacterial artificial chromosomes, bacterial P1 constructions or single chromosome cDNA libraries as reviewed in Price, C. M. (1993) Blood Rev. 7:127-134, and Trask, B. J. (1991) Trends Genet. 7:149-154. FISH (as described in Verma et al. (1988) Human Chromosomes: A Manual of Basic Techniques, Pergamon Press, New York, N.Y.) may be correlated with other physical chromosome mapping techniques and genetic map data. Examples of genetic map data can be found in the 1994 Genome Issue of Science (265:1981f). Correlation between the location of the gene encoding Mnk on a physical chromosomal map and a specific disease, or predisposition to a specific disease, may help to delimit the region of DNA associated with that genetic disease.

The nucleotide sequences of the subject invention may be used to detect differences in gene sequences between normal, carrier or affected individuals. In situ hybridization of chromosomal preparations and physical mapping techniques such as linkage analysis using established chromosomal markers may be used for extending genetic maps. Often the placement of a gene on the chromosome of another mammalian species, such as mouse, may reveal associated markers even if the number or arm of a particular human chromosome is not known. New sequences can be assigned to chromosomal arms or parts thereof, by physical mapping. This provides valuable information to investigators searching for disease genes using positional cloning or other gene discovery techniques. Once the disease or syndrome has been crudely localized by genetic linkage to a particular genomic region, for example, AT to 11q22-23 (Gatti, R. A. et al. (1988) Nature 336:577-580), any sequences mapping to that area may
represent associated or regulatory genes for further investigation. The nucleotide sequences of the subject invention may also be used to detect differences in the chromosomal location due to translocation, inversion, etc. among normal, carrier or affected individuals.

In another embodiment of the invention, the proteins of the invention, its catalytic or immunogenic fragments or oligopeptides thereof, an in vitro model, a genetically altered cell or animal, can be used for screening libraries of compounds in any of a variety of drug screening techniques. One can identify effectors, e.g. receptors, enzymes, proteins, peptides, ligands or substrates that bind to, modulate or mimic the action of one or more of the proteins of the invention. The protein or fragment thereof employed in such screening may be free in solution, affixed to a solid support, borne on a cell surface, or located intracellularly. The formation of binding complexes, between the proteins of the invention and the agent tested, may be measured. Agents could also, either directly or indirectly, influence the activity of the proteins of the invention. Target mechanisms can for example include a kinase activity, particularly the phosphorylation of proteins or peptides, most preferably, but not limited to serine and threonine residues. Another target mechanism could include the regulation of Mnk function by posttranslational modifications such as phosphorylation, dephosphorylation, acetylation, alkylation, ubiquitination, proteolytic processing subcellular localization or degradation. Yet another target mechanism could include the interaction of Mnk with protein binding partners such as, but not limited to, phospholipase A, estrogen receptors, kinases or translation factors. Of particular interest are screening assays for agents that have a low toxicity for mammalian cells. The term "agent" as used herein describes any molecule, e.g. protein or pharmaceutical, with the capability of altering or mimicking the physiological function of one or more of the proteins of the invention. Candidate agents encompass numerous chemical classes, though typically they are organic molecules, preferably small organic compounds having a molecular weight of more
than 50 and less than about 2,500 Daltons. Candidate agents comprise functional groups necessary for structural interaction with proteins, particularly hydrogen bonding, and typically include at least an amine, carbonyl, hydroxyl or carboxyl group, preferably at least two of the functional chemical groups. The candidate agents often comprise carbocyclic or heterocyclic structures and/or aromatic or polyaromatic structures substituted with one or more of the above functional groups.

Candidate agents are also found among biomolecules including peptides, saccharides, fatty acids, steroids, purines, pyrimidines, nucleic acids and derivatives, structural analogs or combinations thereof. Candidate agents are obtained from a wide variety of sources including libraries of synthetic or natural compounds. For example, numerous means are available for random and directed synthesis of a wide variety of organic compounds and biomolecules, including expression of randomized oligonucleotides and oligopeptides. Alternatively, libraries of natural compounds in the form of bacterial, fungal, plant and animal extracts are available or readily produced. Additionally, natural or synthetically produced libraries and compounds are readily modified through conventional chemical, physical and biochemical means, and may be used to produce combinatorial libraries. Known pharmacological agents may be subjected to directed or random chemical modifications, such as acylation, alkylation, esterification, amidification, etc. to produce structural analogs. Where the screening assay is a binding assay, one or more of the molecules may be joined to a label, where the label can directly or indirectly provide a detectable signal.

Another technique for drug screening, which may be used, provides for high throughput screening of compounds having suitable binding affinity to the protein of interest as described in published PCT application WO84/03564. In this method, as applied to the proteins of the invention large numbers of different small test compounds, e.g. aptamers, peptides, low-molecular weight compounds etc., are provided or synthesized on a
solid substrate, such as plastic pins or some other surface. The test compounds are reacted with the proteins or fragments thereof, and washed. Bound proteins are then detected by methods well known in the art. Purified proteins can also be coated directly onto plates for use in the aforementioned drug screening techniques. Alternatively, non-neutralizing antibodies can be used to capture the peptide and immobilize it on a solid support. In another embodiment, one may use competitive drug screening assays in which neutralizing antibodies capable of binding the protein specifically compete with a test compound for binding the protein. In this manner, the antibodies can be used to detect the presence of any peptide, which shares one or more antigenic determinants with the protein.

Candidate agents may also be found in kinase assays where a kinase substrate such as a protein or a peptide, which may or may not include modifications as further described below, or others are phosphorylated by the proteins or protein fragments of the invention. A therapeutic candidate agent may be identified by its ability to increase or decrease the enzymatic activity of the proteins of the invention. The kinase activity may be detected by change of the chemical, physical or immunological properties of the substrate due to phosphorylation. One example could be the transfer of radioisotopically labelled phosphate groups from an appropriate donor molecule to the kinase substrate catalyzed by the polypeptides of the invention. The phosphorylation of the substrate may be followed by detection of the substrates autoradiography with techniques well known in the art.

Yet in another example, the change of mass of the substrate due to its phosphorylation may be detected by mass spectrometry techniques. One could also detect the phosphorylation status of a substrate with a reagent discriminating between the phosphorylated and unphosphorylated status of the substrate. Such a reagent may act by having different affinities for the phosphorylated and unphosphorylated forms of the substrate or by having specific affinity for phosphate groups. Such a
reagent could be, but is not limited to, an antibody or antibody derivative, a recombinant antibody-like structure, a protein, a nucleic acid, a molecule containing a complexed metal ion, an anion exchange chromatography matrix, an affinity chromatography matrix or any other molecule with phosphorylation dependent selectivity towards the substrate.

Such a reagent could be employed to detect the kinase substrate, which is immobilized on a solid support during or after an enzymatic reaction. If the reagent is an antibody, its binding to the substrate could be detected by a variety of techniques as they are described in Harlow and Lane, 1998, Antibodies, CSH Lab Press, NY. If the reagent molecule is not an antibody, it may be detected by virtue of its chemical, physical or immunological properties, being endogenously associated with it or engineered to it.

Yet in another example the kinase substrate may have features, designed or endogenous, to facilitate its binding or detection in order to generate a signal that is suitable for the analysis of the substrates phosphorylation status. These features may be, but are not limited to, a biotin molecule or derivative thereof, a glutathione-S-transferase moiety, a moiety of six or more consecutive histidine residues, an amino acid sequence or hapten to function as an epitope tag, a fluorochrome, an enzyme or enzyme fragment. The kinase substrate may be linked to these or other features with a molecular spacer arm to avoid steric hindrance.

In one example the kinase substrate may be labelled with a fluorophore. The binding of the reagent to the labelled substrate in solution may be followed by the technique of fluorescence polarization as it is described in the literature (see, for example, Deshpande, S. et al. (1999) Prog. Biomed. Optics (SPIE) 3603:261; Parker, G. J. et al. (2000) J. Biomol. Screen. 5:77-88; Wu, P. et al. (1997) Anal. Biochem. 249:29-36). In a variation of this example, a fluorescent tracer molecule may compete with the substrate for the analyte to detect kinase activity by a technique which is know to those skilled in the art as indirect fluorescence polarization.
The nucleic acids encoding the proteins of the invention can be used to
generate transgenic cell lines and animals. These transgenic animals are
useful in the study of the function and regulation of the proteins of the
invention in vivo. Transgenic animals, particularly mammalian transgenic
animals, can serve as a model system for the investigation of many
developmental and cellular processes common to humans. Transgenic
animals may be made through homologous recombination in embryonic
stem cells, where the normal locus of the gene encoding the protein of the
invention is mutated. Alternatively, a nucleic acid construct encoding the
protein is injected into oocytes and is randomly integrated into the genome.
One may also express the genes of the invention or variants thereof in
tissues where they are not normally expressed or at abnormal times of
development. Furthermore, variants of the genes of the invention like
specific constructs expressing anti-sense molecules or expression of
dominant negative mutations, which will block or alter the expression of
the proteins of the invention may be randomly integrated into the genome.
A detectable marker, such as lac Z or luciferase may be introduced into the
locus of the genes of the invention, where upregulation of expression of
the genes of the invention will result in an easily detectable change in
phenotype. Vectors for stable integration include plasmids, retroviruses and
other animal viruses, yeast artificial chromosomes (YACs), and the like.
DNA constructs for homologous recombination will contain at least
portions of the genes of the invention with the desired genetic
modification, and will include regions of homology to the target locus.
Conveniently, markers for positive and negative selection are included.
DNA constructs for random integration do not need to contain regions of
homology to mediate recombination. DNA constructs for random
integration will consist of the nucleic acids encoding the proteins of the
invention, a regulatory element (promoter), an intron and a poly-adenylation
signal. Methods for generating cells having targeted gene modifications
through homologous recombination are known in the field. For embryonic
stem (ES) cells, an ES cell line may be employed, or embryonic cells may
be obtained freshly from a host, e.g. mouse, rat, guinea pig, etc. Such cells are grown on an appropriate fibroblast-feeder layer and are grown in the presence of leukemia inhibiting factor (LIF). ES or embryonic cells may be transfected and can then be used to produce transgenic animals. After transfection, the ES cells are plated onto a feeder layer in an appropriate medium. Cells containing the construct may be selected by employing a selection medium. After sufficient time for colonies to grow, they are picked and analyzed for the occurrence of homologous recombination. Colonies that are positive may then be used for embryo manipulation and morula aggregation. Briefly, morulae are obtained from 4 to 6 week old superovulated females, the Zona Pellucida is removed and the morulae are put into small depressions of a tissue culture dish. The ES cells are trypsinized, and the modified cells are placed into the depression closely to the morulae. On the following day the aggregates are transferred into the uterine horns of pseudopregnant females. Females are then allowed to go to term. Chimeric offsprings can be readily detected by a change in coat color and are subsequently screened for the transmission of the mutation into the next generation (F1-generation). Offspring of the F1-generation are screened for the presence of the modified gene and males and females having the modification are mated to produce homozygous progeny. If the gene alterations cause lethality at some point in development, tissues or organs can be maintained as allogenic or congenic grafts or transplants, or in vitro culture. The transgenic animals may be any non-human mammal, such as laboratory animal, domestic animals, etc., for example, mouse, rat, guinea pig, sheep, cow, pig, and others. The transgenic animals may be used in functional studies, drug screening, and other applications and are useful in the study of the function and regulation of the proteins of the invention in vivo.

Finally, the invention also relates to a kit comprising at least one of (a) a Mnk2 (Mnk2a or Mnk2b) or Mnk1 nucleic acid molecule or a fragment thereof;
(b) a vector comprising the nucleic acid of (a);
(c) a host cell comprising the nucleic acid of (a) or the vector of (b);
(d) a polypeptide encoded by the nucleic acid of (a);
(e) a fusion polypeptide encoded by the nucleic acid of (a);
(f) an antibody, an aptamer or another receptor against the nucleic acid of (a) or the polypeptide of (d) or (e) and
(g) an anti-sense oligonucleotide of the nucleic acid of (a).

The kit may be used for diagnostic or therapeutic purposes or for screening applications as described above. The kit may further contain user instructions.

The Figures show:

Figure 1 shows the increase of triglyceride content of EP(3)3333 and EP(3)3576 male flies caused by homozygous viable integration of the P-vector (in comparison to EP-control males). Shown is the ratio of the triglyceride to protein content of the mutants in percent (%)).

Figure 2 shows the molecular organisation of the mutated LK6 gene locus. The dotted black line represents the position of the cDNA (from position 7544500 to 7559500 on chromosome 3R) that includes the integration sites of EP(3)3333 and EP(3)3576. Transcribed DNA sequences (ESTs) and predicted exons are shown as bars in the lower two lines. Predicted exons of gene CG17342 (GadFly, Lk6) are shown as dark grey bars and introns as light grey bars. Lk6 encodes for a gene that is predicted by GadFly sequence analysis programs as Gadfly Accession Number CG17342.

Figure 3 shows the comparison of Mnk proteins.

Figure 3A shows the comparison (CLUSTAL X 1.8) of Mnk proteins from different species, hXP_030637 refers to human Mnk2 (identical to Genbank Accession No. AF237775), hXP_001600 refers to human Mnk1
(identical to Genbank Accession No. AB000409.1), and AAB18789 refers to the protein encoded by Drosophila Lk6 gene with GadFly Accession No. CG17342.

Gaps in the alignment are represented as -.

**Figure 3B** shows the comparison (CLUSTAL W 1.82) of human Mnk2 proteins. Genbank Accession Number XM_030637.3 is identical to Genbank Accession Number AF237775, and Genbank Accession Number NM_017572.1 shows a different variant of the human Mnk2 protein.

**Figure 3C** shows the comparison (CLUSTAL W 1.82) of human Mnk1 proteins. Genbank Accession Number XM_001600.2 is identical to Genbank Accession Number AB000409.1, and Genbank Accession Number NM_003684.2 shows a different variant of the Mnk1 protein.

**FIGURE 3D.** Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 2a (SEQ.ID NO: 1; GenBank Accession Number AF237775, identical to GenBank Accession Number XM_030637)

**FIGURE 3E.** Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 2a (SEQ ID NO: 2; GenBank Accession Number AF237775, identical to GenBank Accession Number XM_030637)

**FIGURE 3F.** Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 2b (SEQ ID NO: 3; GenBank Accession Number AF237776 or NM_017572)

**FIGURE 3G.** Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 2b (SEQ ID NO: 4; GenBank Accession Number AF237776 or NM_017572)

**FIGURE 3H.** Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 1 (SEQ ID NO: 5; GenBank Accession Number AB000409 or NM_003684 or XM_001600)

**FIGURE 3I.** Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 1 (SEQ ID NO: 6; GenBank Accession Number AB000409 or NM_003684 or XM_001600)
Figure 4 shows the eye phenotype induced by overexpression of an uncoupling protein (dUCPy) that was used to discover factors modulating uncoupling protein activity. In the fly shown in the left part of the picture, dUCPy is expressed at normal levels. In the fly shown in the right part of the photograph, dUCPy is overexpressed, and the eye is reduced.

Figure 5 shows the expression of the Mnk2 gene.

Figure 5A shows the real-time PCR analysis of Mnk2 in wildtype mouse tissues

Figure 5B shows the expression of mouse Mnk2 gene in fasted and obese (ob/ob) mice

Figure 5C shows the expression of mouse Mnk2 gene in fasted and obese mice

Figure 5D shows the real-time PCR mediated comparison of Mnk2 expression during differentiation of mammalian fibroblast (3T3-L1) cells from pre-adipocytes to mature adipocytes

Figure 5E shows real-time PCR mediated comparison of Mnk2 expression during the differentiation of mammalian fibroblast 3T3-F442A cells from preadipocytes to mature adipocytes

Figure 5F shows real-time PCR mediated comparison of Mnk2 expression during the differentiation of mammalian fibroblast TA1 cells from preadipocytes to mature adipocytes

Figure 6 shows the expression of the mouse Mnk1 gene.

Figure 6A shows the real-time PCR analysis of Mnk1 in wildtype mouse tissues

Figure 6B shows the real-time PCR mediated comparison of Mnk1 expression in different mouse models

Figure 6C shows the real-time PCR mediated comparison of Mnk1 expression during differentiation of mammalian fibroblast 3T3-L1 cells from pre-adipocytes to mature adipocytes
Figure 6D shows real-time PCR mediated comparison of Mnk1 expression during the differentiation of mammalian fibroblast 3T3-F442A cells from preadipocytes to mature adipocytes.

Figure 6E shows real-time PCR mediated comparison of Mnk1 expression during the differentiation of mammalian fibroblast TA1 cells from preadipocytes to mature adipocytes.

Figure 7 shows the UCPy sequences.

Figure 7A shows the nucleic acid sequence encoding the Drosophila UCPy protein (SEQ ID NO. 7). The open reading frame is underlined.

Figure 7B shows the amino acid sequence encoding the Drosophila UCPy protein (SEQ ID NO. 8).

Figure 8: In vitro assays for the determination of triglyceride storage, synthesis and transport.

Figure 8A shows reduction in cellular triglyceride levels (μg/mg protein) in cells over expressing Mnk2 compared to control cells. All samples were analysed in duplicates (s1; sample 1, s2; sample 2). The Y-axis shows cellular triglyceride levels (μg/mg protein) and the X-axis shows days of cell differentiation.

Figure 8B shows reduction in insulin stimulated lipid synthesis (dpm/mg protein) in cells over expressing Mnk2 compared to control cells. All samples were analysed in duplicates (s1; sample 1, s2; sample 2). CB; cytochalasin B, illustrates the background synthesis in 3T3L1, O; represents the baseline or un-stimulated glucose transport and hence basal lipid synthesis in the cells, while Ins; insulin shows the stimulated glucose transport and the consequent synthesis of glucose to lipid in 3T3L1 cells. The Y-axis displays disintegrations per minutes/mg protein (dpm/mg protein) and the X-axis denotes the aforementioned proteins.

Figure 8C shows reduction in active transport (AT) of free fatty acids across the plasma membrane of cells over expressing Mnk2 compared to control cells. All samples were analysed in duplicates (as illustrated by twin
bar of identical shadings). PD; passive diffusion illustrated the baseline or non-energy dependant transportation of exogenous fatty acids across the membrane. AT; active transport represents energy dependant transportation of fatty acids across the membrane. The Y-axis shows disintegrations per minutes/mg protein (dpm/mg protein) and the X-axis displays the afore mentioned proteins.

**Figure 9** shows the expression of human Mnk2 in different human tissues.  
**Figure 9A** shows the expression of human Mnk2A and Mnk2B in different human tissues.  
**Figure 9B** shows the expression of human Mnk2A and human Mnk2B during adipocyte differentiation.

**Figure 10** shows the expression of the ectopic mouse Mnk2 (mMnk2DN) transgene in actin-mMnk2DN transgenic mice. Shown is a taqman analysis on different tissues isolated from male actin-mMnk2DN transgenic mice and male wild-type littermates. Data are expressed as fold RNA induction relative to the corresponding wild-type (wt) tissue. The number on top of each bar indicate the fold induction relative to the corresponding wt tissue. Shown is a representative experiment.

**Figure 11** shows that the ectopic mouse Mnk2 (mMnk2DN) expression in actin-mMnk2DN transgenic mice leads to increased body weight. Shown are growth curves from male actin-mMnk2DN transgenic mice (dark grey graph) and male wt littermates (light grey graph) on high fat diet over a time of 10 weeks. Data are expressed as mean body weight over time +/- SE. Shown is a representative experiment with N = 8 respectively N = 10 mice per group.

**Figure 12** shows the exon/intron boundaries of the mouse Mnk2 gene.
Exon/intron boundaries of the mouse Mnk2 gene are illustrated in this figure. Exon numbers, the position of the exons on the cDNA (GenBank accession number BC010256) and intron lengths are indicated. Intron sequences are shown in lowercase letters, exon sequences are shown in capital, bold letters.

**Figure 13** illustrates the targeted deletion of the mouse Mnk2 gene by homologous recombination.

The top line shows the wild type locus of mouse Mnk2, the graphic in the middle shows the targeting vector, and the graphic at the bottom part of the figure illustrates the targeted locus. The exons are shown as black boxes. Restriction sites, translation start site, and stop codon are indicated. The PGK-NEO cassette and the TK cassette are shown as grey boxes. 4.4 kb of genomic region of the mouse Mnk2 gene is replaced by a PGK NEO cassette. The deleted region is indicated. The outside flanking probe used for Southern blot analysis is shown by a black bar. The genomic fragments detected with this probe on EcoR1 digested DNA are shown as arrows. See examples for more detail.

**Figure 14** shows that purified Mnk2a can be activated in vitro with a preparation of the kinases Erk2 and the double point mutant Mek1 S218D S222E.

The Examples illustrate the invention:

**Example 1: Measurement of triglyceride content of homozygous flies** (Figure 1)

The change of triglyceride content of Drosophila melanogaster containing a special expression system (EP-element, Rorth P, Proc Natl Acad Sci USA 1996, 93(22):12418-22) was measured. Mutant flies are obtained from a fly mutation stock collection. The flies are grown under standard conditions
known to those skilled in the art, and in the course of the experiment, additional feedings with bakers yeast (Saccharomyces cerevisiae) are provided. Specifically, homozygous male EP(3)3333 and EP(3)3576 flies were investigated in comparison to control flies (FIGURE 1). For determination of triglyceride content, flies were incubated for 5 min at 90°C in an aqueous buffer using a waterbath, followed by hot extraction. After another 5 min incubation at 90°C and mild centrifugation, the triglyceride content of the flies extract was determined using Sigma Triglyceride (INT 336-10 or -20) assay by measuring changes in the optical density according to the manufacturer’s protocol. As a reference protein content of the same extract was measured using BIO-RAD DC Protein Assay according to the manufacturer’s protocol. The assay was repeated several times. The average triglyceride level of EP collection is shown as 100% in FIGURE 1. EP(3)3333 and EP(3)3576 homozygous flies show constantly a higher triglyceride content than the controls (approx. 140 %). Therefore, the change of gene activity in the locus 86F7 (estimated), where the EP-vector of EP(3)3333 and EP(3)3576 flies is homozygous viably integrated into the Lk6 gene locus, is responsible for changes in the metabolism of the energy storage triglycerides, therefore representing in both cases an obese fly model.

Example 2: Identification of the Drosophila gene responsible for the change in the metabolism of the energy storage triglycerides (Figure 2)

Genomic DNA sequences were isolated that are localized directly adjacent to the integration of the EP vectors (herein EP(3)3333 and EP(3)3576). Using those isolated genomic sequences, public databases like Berkeley Drosophila Genome Project (GadFly) were screened thereby confirming the homozygous viable integration site of the EP(3)3333 and EP(3)3576 vectors. FIGURE 2 shows the molecular organization of this gene locus. In Figure 2, genomic DNA sequence is represented by the assembly as a dotted black line (from position 7544500 to 7559500 on chromosome 3R)
that includes the integration sites of EP(3)3333 and EP(3)3576. Transcribed DNA sequences (expressed sequence tags, ESTs) and predicted exons are shown as bars in the lower two lines. Predicted exons of gene CG17342 (GadFly, Lk6, homologous to Mnk) are shown as dark grey bars and introns are shown as slim grey lines in the middle of the figure. Using plasmid rescue method genomic DNA sequences that are directly localised 3’ of the EP(3)3333 and EP(3)3576 integration site were isolated. Using the plasmid rescue DNA public DNA sequence databases were screened thereby identifying the integration site of EP(3)3333 and EP(3)3576 causing an increase of triglyceride content. EP(3)3333 is integrated in the 5’ region of a 5 prime exon of the gene CG17342 and EP(3)3576 in the 5’ region of an alternative 5’ exon. Mnk encodes for a gene that is predicted by GadFly sequence analysis programs as CG17342. Therefore, expression of the CG17342 could be affected by homozygous viable integration of the EP(3)3333 and EP(3)3576 leading to increase of the energy storage triglycerides and a change of uncoupling protein activity.

Example 3: Cloning of a Drosophila melanogaster gene with homology to human Uncoupling Proteins (UCPs) (Figure 7)

Sequences homologous to human UCP2 and UCP3 genes were identified using the publicly available program BLAST of the data base of the National Center for Biotechnology Information (NCBI) (see, Altschul et al., 1997, Nucleic Acids Res. 25:3389-3402). The homology search yielded sequence fragments of a family of Drosophila genes with UCP homology. They are clearly different to the next related mitochondrial proteins (oxoglutarate carrier).

Using the sequence fragment of one of this genes (herein referred to as \textit{dUCPy}'), a PCR primer pair was generated (Upper 5’-CTAAACAAAAAATTCCAACATAG (SEQ ID NO: 9), Lower 5 prime
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-AAAAGACATAGAAAATACGATAGT (SEQ ID NO: 10)) and a PCR reaction performed on Drosophila cDNA using standard PCR conditions. The amplification product was radioactively labelled and used to screen a cDNA library prepared from adult Drosophila flies (Stratagene). A full-length cDNA clone was isolated, sequenced (FIGURE 7), and used for further experiments. The nucleotide sequence of dUCPy is shown in Figure 7A (SEQ ID NO: 7), the deduced open reading is shown in Figure 7B (SEQ ID NO: 8).

Example 4: Cloning of the dUCPy cDNA into an Drosophila expression vector

In order to test the effects of dUCPy expression in Drosophila cells, the dUCPy cDNA was cloned into the expression vector pUAST (Brand A & Perrimon N, Development 1993, 118:401-415) using the restriction sites NotI and KpnI. The resulting expression construct was injected into the germline of Drosophila embryos and Drosophila strains with a stable integration of the construct were generated. Since the expression vector pUAST is activated by the yeast transcription factor Gal4 which is normally absent from Drosophila cells dUCPy is not yet expressed in these transgenic animals. If pUAST-dUCPy flies are crossed with a second Drosophila strain that expresses Gal4 in a tissue specific manner the offspring flies of this mating will express dUCPy in the Gal4 expressing tissue.

The cross of pUAST-dUCPy flies with a strain that expresses Gal4 in all cells of the body (under control of the actin promoter) showed no viable offspring. This means that dUCPy overexpression in all body cells is lethal. This finding is consistent with the assumption that dUCPy overexpression could lead to a collapse of the cellular energy production.
Expression of dUCPy in a non-vital organ like the eye (Gal4 under control of the eye-specific promoter of the "eyeless" gene) results in flies with visibly damaged eyes. This easily visible eye phenotype is the basis of a genetic screen for gene products that can modify UCP activity.

Example 5: dUCPy modifier screen (Figure 4)

Parts of the genomes of the strain with Gal4 expression in the eye and the strain carrying the pUAST-dUCPy construct were combined on one chromosome using genomic recombination. The resulting fly strain has eyes that are permanently damaged by dUCPy expression. Flies of this strain were crossed with flies of a large collection of mutagenized fly strains. In this mutant collection a special expression system (EP-element, see Rorth, 1996, supra) is integrated randomly in different genomic loci. The yeast transcription factor Gal4 can bind to the EP-element and activate the transcription of endogenous genes close the integration site of the EP-element. The activation of the genes therefore occurs in the same cells (eye) that overexpress dUCPy. Since the mutant collection contains several thousand strains with different integration sites of the EP-element it is possible to test a large number of genes whether their expression interacts with dUCPy activity. In case a gene acts as an enhancer of UCP activity the eye defect will be worsened; a suppressor will ameliorate the defect (see FIGURE 4).

Using this screen a gene with enhancing activity was discovered that was found to be the LK6 kinase in Drosophila.

Example 6: Cloning of Lk6 from Drosophila (Figure 3A)

Genomic DNA neighbouring to the eye-defect rescuing EP-element was cloned by inverse PCR and sequenced. This sequence was used for a BLAST search in a public Drosophila gene database. The amino acid
sequence of the Drosophila protein is shown in Figure 3A (referred to as dmAAB18789).

Example 7: Identification of mammalian LK6 homologous proteins (Figure 3)

Sequences homologous to Drosophila Lk6 were identified using the publicly available program BLASTP 2.2.3 of the non-redundant protein data base of the National Center for Biotechnology Information (NCBI)(see, Altschul et al., 1997, Nucleic Acids Res. 25:3389-3402).

Mnk homologous proteins and nucleic acid molecules coding therefore are obtainable from insect or vertebrate species, e.g. mammals or birds. Particularly preferred are human Mnk homologous polypeptides and nucleic acids, particularly polypeptides and nucleic acids encoding a human Mnk2 protein (Genbank Accession No. AF237775 and NM_017572.1; Genbank Accession No. AF237775 is identical to formerly Genbank Accession No. XM_030637 which was removed at the submitters request; see a Clustal W multiple sequence alignment in FIGURE 3B) or nucleic acids encoding a human Mnk1 protein (Genbank Accession No. AB000409.1 and NM_003684.2; Genbank Accession No. AB000409.1 is identical to formerly Genbank Accession No. XM_001600 which was removed at the submitters request; see a Clustal W multiple sequence alignment in Figure 3C).

Figure 3A shows the alignment of the Mnk proteins from different species, hXP_030637 refers to human Mnk2 (identical to Genbank Accession No. AF237775), hXP_001600 refers to human Mnk1 (identical to Genbank Accession No. AB000409.1), and dmAB18789 refers to the protein encoded by Drosophila gene with GadFly Accession No. CG17342.
The mouse homologous polypeptides of the invention were identified as GenBank Accession Numbers NP_067437.1 (for the mouse homolog MAP kinase-interacting serine/threonine kinase 2; Mnk2; for the cDNA GenBank Accession Number BC010256) and GenBank Accession Numbers NP_067436.1 (for the mouse homolog MAP kinase-interacting serine/threonine kinase 1; Mnk1).

Example 8: Expression of the polypeptides in mammalian (mouse) tissues (Figure 5 and Figure 6)

For analyzing the expression of the polypeptides disclosed in this invention in mammalian tissues, several mouse strains (preferably mice strains C57Bl/6J, C57Bl/6 ob/ob and C57Bl/KS db/db which are standard model systems in obesity and diabetes research) were purchased from Harlan Winkelmann (33178 Borchcn, Germany) and maintained under constant temperature (preferably 22°C), 40 per cent humidity and a light / dark cycle of preferably 14 / 10 hours. The mice were fed a standard chow (for example, from ssniff Spezialitäten GmbH, order number ssniff M-Z V1126-000). For the fasting experiment ("fasted wild type mice"), wild type mice were starved for 48 h without food, but only water supplied ad libitum. (see, for example, Schnetzler et al. J Clin Invest 1993 Jul;92(1):272-80, Mizuno et al. Proc Natl Acad Sci U S A 1996 Apr 16;93(8):3434-8). Animals were sacrificed at an age of 6 to 8 weeks. The animal tissues were isolated according to standard procedures known to those skilled in the art, snap frozen in liquid nitrogen and stored at -80°C until needed.

For analyzing the role of the proteins disclosed in this invention in the in vitro differentiation of different mammalian cell culture cells for the conversion of pre-adipocytes to adipocytes, mammalian fibroblast (3T3-L1) cells (e.g., Green & Kehinde, Cell 1: 113-116, 1974) were obtained from the American Tissue Culture Collection (ATCC, Hanassas, VA, USA;
ATCC-CL 173). 3T3-L1 cells were maintained as fibroblasts and differentiated into adipocytes as described in the prior art (e.g., Qiu et al., J. Biol. Chem. 276:11988-95, 2001; Slieker et al., BBRC 251: 225-9, 1998). In brief, cells were plated in DMEM/10% FCS (Invitrogen, Karlsruhe, Germany) at 50,000 cells/well in duplicates in 6-well plastic dishes and cultured in a humidified atmosphere of 5% CO₂ at 37°C. At confluence (defined as day 0; d0) cells were transferred to serum-free (SF) medium, containing DMEM/HamF12 (3:1; Invitrogen), Fetaun (300 μg/ml; Sigma, Munich, Germany), Transferrin (2 μg/ml; Sigma), Pantothenate (17 μM; Sigma), Biotin (1 μM; Sigma), and EGF (0.8 nM; Hoffmann-La Roche, Basel, Switzerland). Differentiation was induced by adding Dexamethasone (DEX; 1 μM; Sigma), 3-Methyl-Isobutyl-1-Methylxanthine (MIX; 0.5 mM; Sigma), and bovine Insulin (5 μg/ml; Invitrogen). Four days after confluence (d4), cells were kept in SF medium, containing bovine insulin (5 μg/ml) until differentiation was completed. At various time points of the differentiation procedure, beginning with day 0 (day of confluence) and day 2 (hormone addition; for example, dexamethason and 3-isobutyl-1-methylxanthin), up to 10 days of differentiation, suitable aliquots of cells were taken every two days.

Alternatively, mammalian fibroblast 3T3-F442A cells (e.g., Green & Kehinde, Cell 7: 105-113, 1976) were obtained from the Harvard Medical School, Department of Cell Biology (Boston, MA, USA). 3T3-F442A cells were maintained as fibroblasts and differentiated into adipocytes as described previously (Djian, P. et al., J. Cell. Physiol., 124:554-556, 1985). At various time points of the differentiation procedure, beginning with day 0 (day of confluence and hormone addition, for example, Insulin), up to 10 days of differentiation, suitable aliquots of cells were taken every two days. 3T3-F442A cells are differentiating in vitro already in the confluent stage after hormone (insulin) addition.
TaqMan Analysis of the proteins of the invention was carried out (Figure 5 and Figure 6). RNA was isolated from mouse tissues or cell culture cells using Trizol Reagent (for example, from Invitrogen, Karlsruhe, Germany) and further purified with the RNeasy Kit (for example, from Qiagen, Germany) in combination with an DNase-treatment according to the instructions of the manufacturers and as known to those skilled in the art. Total RNA was reverse transcribed (preferably using Superscript II RNaseH-Reverse Transcriptase, from Invitrogen, Karlsruhe, Germany) and subjected to Taqman analysis preferably using the Taqman 2xPCR Master Mix (from Applied Biosystems, Weiterstadt, Germany; the Mix contains according to the Manufacturer for example AmpliTaq Gold DNA Polymerase, AmpErase UNG, dNTPs with dUTP, passive reference Rox and optimized buffer components) on a GeneAmp 5700 Sequence Detection System (from Applied Biosystems, Weiterstadt, Germany).

For the analysis of the expression of Mnk2 or Mnk1, taqman analysis was performed using the following primer/probe pairs:
Mouse Mnk1 forward primer (Seq ID NO: 11) 5'-GCT GAG GGC CTC TGC TCC-3';
Mouse Mnk1 reverse primer (Seq ID NO: 12) 5'-TCG CCT TCG AGC CAG G-3';
Mouse Mnk1 Taqman probe (Seq ID NO: 13) (5/6-FAM) TGA AGC TGT CCC CTC CAT CCA AAT CTC (5/6-TAMRA)

Taqman-1856F Mnk2 forward primer (SEQ ID NO: 14): 5'-TGCACTTGTGGACCCGA-3'
Taqman-1923R Mnk2 reverse primer (SEQ ID NO: 15): 5'-TTTCTGATTGTCAACCCCTCCAA-3'
Taqman-1877T Mnk2 Taqman probe (SEQ ID NO: 16): (5/6-FAM)-CCCCATCATCCACCTGCAGTGCTCC-(5/6-TAMRA)
Taqman analysis revealed that Mnk2 is the more interesting homologue of the fly Lk6 gene. The results are shown in Figure 5 and Figure 6. In comparison to Mnk1, which is rather ubiquitously expressed, Mnk2 shows its highest expression levels in the brown and white adipose tissues (FIGURE 5A and 6A, respectively). The expression of Mnk2 in white adipose tissue is under metabolic control: In fasted as well as obese (ob/ob) mice, expression is reduced to about 40% of wildtype levels (FIGURE 5C; see also FIGURE 5B). In addition, expression of Mnk2 is strongly induced during the in vitro differentiation of 3T3-L1 (FIGURE 5D) as well as of two additional model systems for the in vitro differentiation of preadipocytes to adipocytes, the 3T3-F442A and TA1 cell lines (FIGURE 5E and FIGURE 5F, respectively). Contrary to this, the relative expression levels of Mnk1 remain unchanged during the differentiation of these cell lines (FIGURES 6D and 6E, respectively).

Example 9: Expression of the polypeptides in mammalian (human) tissues (Figure 9)

Human primary adipocytes were differentiated into mature adipocytes as described by Hauner et al. 1989 (J Clin Invest84(5):1663-70). Briefly, cells were grown in DMEM/Nutrient Mix F12, 1% PenStrep, 17 μM Biotin, 33 μM Pantothenat, 10% none heat inactivated fetal calf serum. On day 0 of differentiation, the medium was changed to DMEM/Nutrient Mix F12, 1% Pen/Strep, 17 μM Biotin, 33 μM Pantothenat, 0,01 mg/ml Transferrin, Hydrocortisone, 20 nM humanes Insulin, 0,2 nM T3, 25 nM Dexamethasone, 250 μM IBMX, 3 μM Rosiglitazone. On day 4 of differentiation, the medium was changed to DMEM/Nutrient Mix F12 1%Pen/Strep, 17 μM Biotin, 33 μM Pantothenat, 0,01 mg/ml Transferrin, 100 nM Hydrocortisone, 20 nM humanes Insulin, 0,2 nM T3. At various time points of the differentiation procedure, beginning with day 0 (day of
confluence) and day 4 (hormone addition), up to 14 days of differentiation, suitable aliquots of cells were taken every two days. RNA was isolated from human cell culture cells using Trizol Reagent (for example, from Invitrogen, Karlsruhe, Germany) and further purified with the RNeasy Kit (for example, from Qiagen, Germany) in combination with an DNase-treatment according to the instructions of the manufacturers and as known to those skilled in the art.

In addition to the RNA isolated from human adipocytes at different differentiation stage, RNAs isolated from different human tissues were obtained from Invitrogen Corp., Karlsruhe, Germany: (i) total RNA from human adult skeletal muscle (Invitrogen Corp. Order Number 735030); (ii) total RNA from human adult lung (Invitrogen Corp. Order Number 735020); (iii) total RNA from human adult liver (Invitrogen Corp. Order Number 735018); (iv) total RNA from human adult placenta (Invitrogen Corp. Order Number 735026); (v) total RNA from human adult testis (Invitrogen Corp. Order Number 64101-1); (vi) total RNA from human normal adipose tissue (Invitrogen Corp. Order Number D6005-01); (vii) total RNA from human normal pancreas (Invitrogen Corp. Order Number DG6101); (viii) total RNA from human normal brain (Invitrogen Corp. Order Number D6030-01). The RNA was treated with DNase according to the instructions of the manufacturers (for example, from Qiagen, Germany) and as known to those skilled in the art.

Total RNA was reverse transcribed (preferably using Superscript II RNaseH-Reverse Transcriptase, from Invitrogen, Karlsruhe, Germany) and subjected to Taqman analysis preferably using the ‘Taqman 2xPCR Master Mix’ (page 66, line 31: Weiterstadt, Germany; see Example 8).

Taqman analysis was performed preferably using the following primer/probe pairs:
For the amplification of human Mnk2a:

human Mnk2a forward primer (SEQ ID NO: 17): 5'- cca tct ccc cct ctg tac ata gg -3'; human Mnk2a reverse primer (SEQ ID NO: 18): 5'- ccg gct ggc gat agc tta a -3'; Taqman probe (SEQ ID NO: 19): (5/6-FAM) cac ccg tcc ccc aat caa atc taa agg (5/6-TAMRA)

For the amplification of human Mnk2b:

human Mnk2b forward primer (SEQ ID NO: 20): 5'- TTA CTG TGA ATG AGT GAA GAT CCT GG -3'; human Mnk2b reverse primer (SEQ ID NO: 21): 5'- ATG GCC GTT CAC CGT CC -3'; Taqman probe (SEQ ID NO: 22): (5/6-FAM) CCA GGC CAG CTC CCA TCG CTG (5/6-TAMRA)

As shown in Figure 9A, real time PCR (Taqman) analysis of the expression of Mnk2a and Mnk2b protein in human tissues revealed that both proteins are expressed in all tissues analysed with high levels of expression in adipose tissue, muscle, lung, testis, and placenta.

The relative expression levels of both human Mnk2 splice variants is the same for all tissues analysed. Both show highest expression levels in tissues relevant for metabolic disorders namely adipose and muscle tissue.

As shown in Figure 9B, Mnk2a as well as Mnk2b are upregulated during human adipocyte differentiation. This suggests a function of both proteins in the metabolism of mature adipocytes.

Example 10: Assays for the determination of triglyceride storage, synthesis and transport (Figure 8)

Retroviral infection of preadipocytes

Packaging cells were transfected with retroviral plasmids pLPCX carrying mouse Mnk2 transgene and a selection marker using calcium phosphate procedure. Control cells were infected with pLPCX carrying no transgene.
Briefly, exponentially growing packaging cells were seeded at a density of 350,000 cells per 6-well in 2 ml DMEM + 10 % FCS one day before transfection. 10 min before transfection chloroquine was added directly to the overlaying medium (25 μM final concentration). A 250 μl transfection mix consisting of 5 μg plasmid-DNA (candidate:helper-virus in a 1:1 ratio) and 250 mM CaCl₂ was prepared in a 15 ml plastic tube. The same volume of 2 x HBS (280 μM NaCl, 50 μM HEPES, 1.5 mM Na₂HPO₄, pH 7.06) was added and air bubbles were injected into the mixture for 15 sec. The transfection mix was added drop wise to the packaging cells, distributed and the cells were incubated at 37°C, 5 % CO₂ for 6 hours. The cells were washed with PBS and the medium was exchanged with 2 ml DMEM + 10 % CS per 6-well. One day after transfection the cells were washed again and incubated for 2 days of virus collection in 1 ml DMEM + 10 % CS per 6-well at 32°C, 5 % CO₂. The supernatant was then filtered through a 0.45 μm cellulose acetate filter and polybrene (final concentration 8 μg/ml) was added. Mammalian fibroblast (3T3-L1) cells in a sub-confluent state were overlaid with the prepared virus containing medium. The infected cells were selected for 1 week with 2 μg/ml puromycin. Following selection the cells were checked for transgene expression by western blot and immunofluorescence. Over expressing cells were seeded for differentiation.

3T3-L1 cells were maintained as fibroblasts and differentiated into adipocytes as described in the prior art and in example 8. For analysing the role of the proteins disclosed in this invention in the in vitro assays for the determination of triglyceride storage, synthesis and transport were performed.

Preparation of cell lysates for analysis of metabolites
Starting at confluence (D0), cell media was changed every 48 hours. Cells and media were harvested 8 hours prior to media change as follows. Media was collected, and cells were washed twice in PBS prior to lyses in 600 μl HB-buffer (0.5% Polyoxyethylene 10 tridecylethan, 1 mM EDTA, 0.01M
NaH$_2$PO$_4$, pH 7.4). After inactivation at 70°C for 5 minutes, cell lysates were prepared on Bio 101 systems lysing matrix B (0.1 mm silica beads; Q-Biogene, Carlsbad, USA) by agitation for 2 x 45 seconds at a speed of 4.5 (Fastprep FP120, Bio 101 Thermosavant, Holbrook, USA). Supernatants of lysed cells were collected after centrifugation at 3000 rpm for 2 minutes, and stored in aliquots for later analysis at -80°C.

Changes in cellular triglyceride levels during adipogenesis (Figure 8A) Cell lysates and media were simultaneously analysed in 96-well plates for total protein and triglyceride content using the Bio-Rad DC Protein assay reagent (Bio-Rad, Munich, Germany) according to the manufacturer’s instructions and a modified enzymatic triglyceride kit (GPO-Trinder; Sigma) briefly final volumes of reagents were adjusted to the 96-well format as follows: 10 µl sample was incubated with 200 µl reagent A for 5 minutes at 37°C. After determination of glycerol (initial absorbance at 540 nm), 50 µl reagent B was added followed by another incubation for 5 minutes at 37°C (final absorbance at 540 nm). Glycerol and triglyceride concentrations were calculated using a glycerol standard set (Sigma) for the standard curve included in each assay.

As shown in Figure 8A, we found that in Mnk2 overexpressing cells cellular triglyceride levels were significantly lower from day 4 to day 12 of adipogenesis compared to that in the control cells (Figure 8A). These results indicate that Mnk2 targets regulatory pathways or enzymes involved in lipid metabolism, which we analysed in more detail in the lipid synthesis and FFA transport assays described below.

Synthesis of lipids during adipogenesis (Figure 8B) During the terminal stage of adipogenesis (day 12) cells were analysed for their ability to metabolise lipids. A modified protocol to the method of Jensen et al (2000), JBC 275, 40148, for lipid synthesis was established. Cells were washed 3 times with PBS prior to serum starvation in
Krebs-Ringer-Bicarbonate-Hepes buffer (KRBH; 134 mM NaCl, 3.5 mM KCl, 1.2 mM KH$_2$PO$_4$, 0.5 mM MgSO$_4$, 1.5 mM CaCl$_2$, 5 mM NaHCO$_3$, 10 mM Hepes, pH 7.4), supplemented with 0.1% FCS for 2.5h at 37°C. For insulin-stimulated lipid synthesis, cells were incubated with 1 μM bovine insulin (Sigma; carrier: 0.005 N HCl) for 45 min at 37°C. Basal lipid synthesis was determined with carrier only. $^{14}$C(U)-D-Glucose (NEN Life Sciences) in a final activity of 1 μCi/Well/ml in the presence of 5 mM glucose was added for 30 min at 37°C. For the calculation of background radioactivity, 25 μM Cytochalasin B (Sigma) was used. All assays were performed in duplicate wells. To terminate the reaction, cells were washed 3 times with ice cold PBS, and lysed in 1 ml 0.1 N NaOH. Protein concentration of each well was assessed using the standard Biuret method (Protein assay reagent; Bio-Rad). Total lipids were separated from aqueous phase after overnight extraction in Insta-Fluor scintillation cocktail (Packard Bioscience) followed by scintillation counting.

Our results clearly show that Mnk2 overexpressing cells were less effective at synthesising lipids from exogenous glucose. Consequently, the levels of insulin stimulated lipid synthesis are significantly lower at day 12 of adipogenesis when compared to control cells (Figure 8B). The lower lipid levels observed in the experiments above therefore result most likely from a lower lipid synthesis rate and are not the result of an increased turn over of lipid stores.

Transport and metabolism of free fatty acids across during adipogenesis (Figure 8C)

During the terminal stage of adipogenesis (D12) cells were analysed for their ability to transport long chain fatty acid across the plasma membrane. A modified protocol to the method of Abumrad et al (1991) (Proc. Natl. Acad. Sci. USA, 1991: 88; 6008-12) for cellular transportation of fatty acid was established. In summary, cells were washed 3 times with PBS prior to serum starvation. This was followed by incubation in KRBH buffer,
supplemented with 0.1% FCS for 2.5h at 37°C. Uptake of exogenous free fatty acids was initiated by the addition of isotopic media containing non radioactive oleate and (³H)oleate (NEN Life Sciences) complexed to serum albumin in a final activity of 1 μCi/Well/ml in the presence of 5 mM glucose for 30 min at room temperature (RT). For the calculation of passive diffusion (PD) in the absence of active transport (AT) across the plasma membrane 20 mM of phloretin in glucose free media (Sigma) was added for 30 min at room temperature (RT). All assays were performed in duplicate wells. To terminate the active transport 20 mM of phloretin in glucose free media was added to the cells. Cells were lysed in 1 ml 0.1N NaOH and the protein concentration of each well were assessed using the standard Biuret method (Protein assay reagent; Bio-Rad). Esterified fatty acids were separated from free fatty acids using overnight extraction in Insta-Fluor scintillation cocktail (Packard Bioscience) followed by scintillation counting.

We found that transport of exogenous fatty acids across the plasma membrane of Mnk2 overexpressing cells and hence esterification of these metabolites were considerably lower at day 12 of adipogenesis when compared to control cells (Figure 8C). Taken together the overexpression of Mnk2 showed an effect on triglyceride metabolism in all three assays we performed in 3T3-L1 cells, making it a potential interesting drug target to treat metabolic disorders.

Example 11: Generation and analysis of Mnk2 transgenic animals (β-actin-mMnk2DN)

Generation of the transgenic animals
Mouse Mnk2 cDNA was isolated from mouse brown adipose tissue (BAT) using standard protocols as known to those skilled in the art. The cDNA was amplified by RT-PCR using the following primer pair:
Mnk2 forward primer (SEQ ID NO: 23): 5’ AAG TTG GCC TTC GCG TTA GAC 3’
Mnk2 reverse primer (SEQ ID NO: 24): 5’ CGA TAT GTA CAA GGA GCT AG 3’.
The resulting Mnk2 cDNA was cloned into pBluescript KS+ (Stratagene) according to standard protocols, resulting in a plasmid referred to as pKS+-mMnk2’. The cDNA of pKS+-mMnk2 c was mutated using site directed mutagenesis (Stratagene), according to the manufacturer’s instructions. Using the Mnk2 top oligo (SEQ ID NO: 25): 5’ CTC CCC CAT CTC CGC ACC AGA GCT GCT CGC CCC GTG TGG GTC AG 3’ and the Mnk2 bottom oligo (SEQ ID NO: 26): 5’ CTG ACC CAC ACG GGG CGA GCT CTG GTG CGG AGA TGG GGG AG 3’, two point mutations were introduced into the cDNA resulting in amino acid exchanges at position T197 and T202 to A197 and A202 of the Mnk2 cDNA.

The resulting mutated cDNA (referred to as mMnk2DN) was cloned into the EcoRV cloning site of the transgenic expression vector pTG-β-actin-X-hgh-bgh-polyA. The β-actin-Mnk2DN transgene was microinjected into the male pronucleus of fertilized mouse embryos (preferably strain C57/BL6/CBA F1 (Harlan Winkelmann). Injected embryos were transferred into pseudo-pregnant foster mice. Transgenic founders were detected by PCR analysis using the forward primer (SEQ ID NO: 27): 5’ GCT GCT GGT CCG AGA TGC C 3’ and reverse primer (SEQ ID NO: 28): 5’ GGG TCA TGC GCG ATC CCC 3’. Two independent transgenic mouse lines containing the β-actin-Mnk2DN construct were established and kept on a C57/BL6 background. Briefly, founder animals were backcrossed with C57/BL6 mice to generate F1 mice for analysis. Transgenic mice were continuously bred onto the C57/BL6 background.

Expression of the construct in different mouse tissues (Figure 10) using standard techniques, β-actin-Mnk2DN transgene expression was verified by Taqman analysis using forward primer (SEQ ID NO: 29): 5’ CAG CGT GGT AGT ACA GGA CGT G 3’, reverse primer (SEQ ID NO: 30): 5’ TCC CTG TGG GCG ATG C 3’ and primer (SEQ ID NO: 31): 5’ CAG TGC
CCT GGA CTT CCT GCA TAA CAA 3'. Taqman analysis was performed using a representative panel of mouse tissues.

The expression of the bactin-Mnk2DN transgene was observed in the following tissues: WAT, muscle, liver, kidney, thymus, heart, lung, and spleen. Expression levels of the transgene were 2.8 - 16.9 fold increased relative to Mnk2 expression in wild-type mice, depending on the tissue analyzed. No Mnk2 transgene expression was detected in BAT tissue (see Figure 10).

Analysis of the bodyweight of the transgenic mice (Figure 11)
After weaning, male β-actin-mMnk2DN transgenic mice and their wild-type (wt) littermates controls were placed in groups of 4 to 5 animals (N=4 up to N=5) on control diet (preferably Altromin C1057 mod control, 4.5% crude fat or high fat diet (preferably Altromin C1057mod. high fat, 23.5% crude fat). Total body weight of the animals was measured weekly over a period of 12-16 weeks.

On each diet, mean bodyweight of β-actin-mMnk2DN transgenic mice was clearly increased compared to wildtype littermates on the respective diet. Significant differences in mean bodyweight were first observed around the end of postnatal week 4 on both diets. After 10 weeks on high fat diet, the mean bodyweight of β-actin-mMnk2DN transgenic mice compared to wt littermates was increased by 8.8g (= 23% increase in mean bodyweight relative to wt littermates) (Figure 11). Similar differences in mean body weight were observed in wt and β-actin-mMnk2DN transgenic mice on control diet (data not shown). Thus, our results clearly show that the ectopic expression of mMnk2DN transgene leads to an increase in bodyweight. The effect appears independently of the diet given, as it can be seen on control diet as well as on high fat diet.

Example 11: Generation and analysis of mMnk2 -/- mice (Figure 12 and Figure 13)
A 605 base pair probe of the mMnk2 cDNA (GenBank accession number BC010256; position 61-665) was amplified from mouse white adipose tissue (WAT) cDNA by PCR using forward primer (SEQ ID NO: 32): 5′ ACA TCA GCC CAC AGT GTG A 3′ and reverse primer (SEQ ID NO: 33): 5′ TCT CCA TTG AGT TTG ATA CCA 3′. This probe was used to screen a 129SVJ genomic phage library (obtained from Stratagene). Three independent clones were isolated and subcloned into the NotI cloning site of pBluescript KS+ (Stratagene). These genomic clones were used for restriction mapping and sequencing to characterize the genomic locus of mouse Mnk2 (Figure 12). A PGK-neomycin cassette was inserted into the locus of mouse Mnk2 replacing 4.4 kb of genomic DNA thereby deleting the complete coding region of mMnk2. Briefly, an 8kb SpeI - NotI fragment was cloned into the XbaI site of pBluescript KS+ upstream of the PGK-Neomycin cassette, which was inserted into the Smal site of pBluescript. A 1kb genomic fragment was amplified by PCR using the following primer pair (non priming nucleotides / attached restriction sites are lower case letters): Mnk2-SA forward primer (SEQ ID NO: 34): EcoRI 5′ cgg aat CCA CTA GCT CCT TGT ACA TAT 3′ ; Mnk2-SA reverse primer (SEQ ID NO: 35): Clal 5′ cca tgc atG GAA CTC GTA TTG CAT AGT AG 3′. The resulting fragment was inserted into the EcoRI / Clal site of pBluescript KS+. As a negative selection marker a thymidine kinase cassette was cloned into Clal / Xhol site of the targeting construct. (Figure 13) The construct was linearized by NotI digestion and electroporated into mouse embryonic stem (ES) cells. ES cell clones were selected by G418 and Gancyclovir treatment (preferably 350 μg G418/ml and 2 μm Gancyclovir). Out of 600 neomycin resistant colonies, two independent homologous recombined ES cell clones were identified by PCR. The results were confirmed by southern blot analysis with EcoRI digested genomic DNA using a 3′ flanking probe (position 2495-3065 mMnk2 cDNA). A single integration event was confirmed by Southern blot analysis of BamHI digested DNA with a Neomycin probe. ES cell clones were aggregated with 8-cell-stage embryos from NMRI mice and developing blastocysts were
transferred into pseudo-pregnant mice to generate chimeras. Chimeras were bred with C57/B6 mice and offsprings were genotyped by PCR using the following primers: Mnk2-ES primer (SEQ ID NO: 36): 5’ AGA CTA GGG AGG AGG GTG GAG GA 3’; Mnk2-KO primer (SEQ ID NO: 37): 5’ GGT GGA TGT GGA ATG TGT GCC A 3’; Mnk2-WT 5’ GGG GTG TAG GGG TCT GTT AGG 3’. Heterozygous mice were used for further intercrosses and analyzed.

Example 11: Small molecule screening

Compounds which are suitable for the prophylaxis, treatment or diagnosis of Mnk-related metabolic disorders may be identified via a kinase assay, a binding assay or any other suitable assay to measure a function associated with the Mnk polypeptide, a Mnk polypeptide fragment or derivative thereof. This kinase assay may be based on recombinant human Mnk2 (Mnk2a or Mnk2b) or Mnk1 protein and a labelled peptide comprising the eIF4E target sequence, a labelled recombinant eIF4E target sequence or a labelled recombinant eIF4E protein as a substrate. The assay may be a radioactive kinase assay or an assay based on using an anti-phosphoserine antibody which is capable of recognizing eIF4E phosphorylation at Ser209.

For example, the kinases Mnk2a (GenBank Accession Number AF237775; see also FIGURE 3D and 3E), Erk2 (GenBank Accession Number M84489) and a double point mutant of Mek1 (GenBank Accession Number Q02750) containing the amino acid substitutions Ser218Asp and Ser222Glu (S218D S222E) were expressed in E.coli and subsequently purified using methods known to those skilled in the art. Preferably, in a kinase reaction of 50 µl, 2.0 µM Mnk2a was incubated with 200 nM Erk2 plus 20 nM Mek1 S218D S222E (labelled lanes 1 to 4 in Figure 14) or with 50 nM Erk2 plus 2.5 nM Mek1 S218D S222E (labelled lanes 5 to 8 in Figure 14) in the presence of 1.0 mM ATP, 50 mM Hepes/KOH, 5 mM magnesium chloride and 0.5 mM DTT at 30°C. At the indicated time points (0, 10, 20 and 40 minutes, see
Figure 14), samples were taken from the reaction, diluted in SDS sample buffer containing 50 mM EDTA and separated by SDS polyacrylamide gel electrophoresis (SDS-PAGE). SDS-PAGE separated reaction samples were blotted onto nitrocellulose and probed with an antibody against a phospho-epitope, essential for the activation of Mnk (anti-Phospho-Mnk Thr197/202; Cell Signaling Technology, Inc., Beverly, MA). The anti-Phospho-Mnk antibody was detected with a peroxidase-coupled anti rabbit antibody (Sigma-Aldrich, St. Louis, MO) as described elsewhere (Harlow and Lane, 1998, Antibodies, Cold Spring Harbor Laboratory Press, NY).

As the reaction progresses, the activation of Mnk2a can be visualized by Mnk2s immuno-reactivity with the anti Phospho-Mnk antibody (see upper panel in Figure 14). In addition, Mnk2a was visualized by Coomassie staining of the gel. Arrows indicate the Coomassie stained Mnk2a as its mobility is retarded with increasing phosphorylation (see lower panel in Figure 14).

The generation of the phospho-epitope, essential for the activation of Mnk2a, and the high degree of efficiency of this process (as shown by the nearly complete electrophoretic mobility shift) demonstrate the suitablility of this approach to produce enzymatically active Mnk2a.

For the validation of the assay, known Mnk inhibitors such as CGP57380 or CGP025088 may be used (see, Knauf et al., 2001, Mol. Cell. Biol. 21:5500, Tschopp et al., 2000, Mol Cell Biol Res Comm 3:205 and Slentz-Kesler et al., 2000, Genomics 69:63). As a negative control, CGP052088 may be used.

Alternatively, the screening may comprise the use of cellular based screening systems, e.g. prokaryotic or eukaryotic cells which overexpress
Mnk proteins. Furthermore, transgenic animals capable of overexpressing or underexpressing Mnk2 and/or Mnk1 may be used.

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All publications and patents mentioned in the above specification are herein incorporated by reference.

Various modifications and variations of the described method and system of the invention will be apparent to those skilled in the art without departing from the scope and spirit of the invention. Although the invention has been described in connection with specific preferred embodiments, it should be understood that the invention as claimed should not be unduly limited to such specific embodiments. Indeed, various modifications of the described modes for carrying out the invention which are obvious to those skilled in molecular biology or related fields are intended to be within the scope of the following claims.
Claims

1. A pharmaceutical composition comprising a nucleic acid molecule of the MAP kinase interacting kinase (Mnk) gene family or a polypeptide encoded thereby or a fragment or a variant of said nucleic acid molecule or said polypeptide or an antibody, an aptamer or another receptor recognizing a nucleic acid molecule of the Mnk gene family or a polypeptide encoded thereby together with pharmaceutically acceptable carriers, diluents and/or adjuvants.

2. The composition of claim 1, wherein the nucleic acid molecule is a vertebrate or insect Mnk nucleic acid, particularly a human Mnk homologous nucleic acid, particularly a nucleic acid encoding a Mnk homologous gene on human chromosome 19 (Mnk2)(Genbank Accession No. XM_030637, identical to Genbank Accession No. AF237775.1, and/or Genbank Accession Number NM_017572.1) or a human Mnk protein on chromosome 1 (Mnk1) (Genbank Accession No. XM_001600, identical to Genbank Accession No. AB00409.1, and/or Genbank Accession Number NM_003684.2) or a fragment thereof or a variant thereof.

3. The composition of claim 1 or 2, wherein said nucleic acid molecule (a) hybridizes at 50°C in a solution containing 1 x SSC and 0.1% SDS to a Mnk nucleic acid or the complementary strand thereof, particularly a nucleic acid molecule encoding the amino acid sequences shown in Figure 3; (b) it is degenerate with respect to the nucleic acid molecule of (a) (c) encodes a polypeptide which is at least 85%, preferably at least 90%, more preferably at least 95%, more preferably at
least 98% and up to 99.6% identical to polypeptides shown in figure 3;
(d) differs from the nucleic acid molecule of (a) to (c) by mutation and wherein said mutation causes an alteration, deletion, duplication or premature stop in the encoded polypeptide.

4. The composition of any one of claims 1-3, wherein the nucleic acid molecule is a DNA molecule, particularly a cDNA or a genomic DNA.

5. The composition of any one of claims 1-4, wherein said nucleic acid encodes a polypeptide contributing to regulating the energy homeostasis and/or the metabolism of triglycerides and/or to membrane stability and/or function in organelles such as mitochondria.

6. The composition of any one of claims 1-5, wherein said nucleic acid molecule is a recombinant nucleic acid molecule.

7. The composition of any one of claims 1-6, wherein the nucleic acid molecule is a vector, particularly an expression vector.

8. The composition of any one of claims 1-5, wherein the polypeptide is a recombinant polypeptide.

9. The composition of claim 8, wherein said recombinant polypeptide is a fusion polypeptide.

10. The composition of any one of claims 1-7, wherein said nucleic acid molecule is selected from hybridization probes, primers and anti-sense oligonucleotides.
11. The composition of any one of claims 1-10 which is a diagnostic composition.

12. The composition of any one of claims 1-10 which is a pharmaceutical composition.

13. The composition of any one of claims 1-12 for the manufacture of an agent for detecting and/or verifying, for the treatment, alleviation and/or prevention of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs, and others, in cells, cell masses, organs and/or subjects.

14. Use of a nucleic acid molecule of the Mnk gene family or a polynucleotide encoded thereby or a fragment or a variant of said nucleic acid molecule or said polypeptide or an antibody, an aptamer or another receptor recognizing a nucleic acid molecule of the Mnk gene family or a polypeptide encoded thereby for controlling the function of a gene and/or a gene product which is influenced and/or modified by an Mnk homologous polypeptide.

15. Use of the nucleic acid molecule of the Mnk gene family or a polynucleotide encoded thereby or a fragment or a variant of said nucleic acid molecule or said polypeptide or an antibody, an aptamer or another receptor recognizing a nucleic acid molecule of the Mnk gene family or a polypeptide encoded thereby for identifying
substances capable of interacting with an Mnk homologous polypeptide.


17. The animal of claim 16, wherein the expression of the Mnk homologous polypeptide is increased and/or reduced.

18. A recombinant host cell exhibiting a modified expression of an Mnk homologous polypeptide.

19. The cell of claim 18 which is a human cell.

20. A method of identifying a (poly)peptide involved in the regulation of energy homeostasis and/or metabolism of triglycerides in a mammal comprising the steps of
   (a) contacting a collection of (poly)peptides with an Mnk homologous polypeptide or a fragment thereof under conditions that allow binding of said (poly)peptides;
   (b) removing (poly)peptides which do not bind and
   (c) identifying (poly)peptides that bind to said Mnk homologous polypeptide.

21. A method of screening for an agent which modulates the interaction of an Mnk homologous polypeptide with a binding target/agent, comprising the steps of
   (a) incubating a mixture comprising
       (aa) an Mnk homologous polypeptide, or a fragment thereof;
       (ab) a binding target/agent of said Mnk homologous polypeptide or fragment thereof; and
(a) a candidate agent under conditions whereby said Mnk polypeptide or fragment thereof specifically binds to said binding target/agent at a reference affinity;

(b) detecting the binding affinity of said Mnk polypeptide or fragment thereof to said binding target to determine an (candidate) agent-biased affinity; and

(c) determining a difference between (candidate) agent-biased affinity and the reference affinity.

22. A method of screening for an agent which modulates the activity of an Mnk homologous polypeptide, comprising the steps of

(a) incubating a mixture comprising

   (aa) an Mnk homologous polypeptide, or a fragment thereof;

   (ab) a candidate agent, under conditions whereby said Mnk polypeptide or fragment thereof exhibits a reference activity;

(b) detecting the activity of said Mnk polypeptide or fragment thereof to determine an (candidate) agent-biased activity, and

(c) determining a difference between a (candidate) agent-biased activity and the reference activity.

23. The method of claim 21 or 22, wherein the candidate agent is selected from peptides and low-molecular weight organic compounds.

24. The method of any one of claims 20-23, wherein a known Mnk effector is used as a positive control for assay development and/or validation of candidate agents.
25. A method of producing a composition comprising the (poly)peptide identified by the method of claim 20 or the agent identified by the method of any one of claims 21-24 with a pharmaceutically acceptable carrier, diluent and/or adjuvant.

26. The method of claim 25 wherein said composition is a pharmaceutical composition for preventing, alleviating or treating diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs, and others, in cells, cell masses, organs and/or subjects.

27. Use of a polypeptide as identified by the method of claim 20 or of an agent as identified by the method of any one of claims 21-24 for the preparation of a pharmaceutical composition for the treatment, alleviation and/or prevention of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs, and others, in cells, cell masses, organs and/or subjects.
28. Use of a nucleic acid molecule of the Mrk family or of a fragment thereof for the preparation of a non-human animal which over- or under-expresses the Mrk gene product.

29. Kit comprising at least one of
   (a) an Mrk nucleic acid molecule or a fragment thereof;
   (b) a vector comprising the nucleic acid of (a);
   (c) a host cell comprising the nucleic acid of (a) or the vector of (b);
   (d) a polypeptide encoded by the nucleic acid of (a);
   (e) a fusion polypeptide encoded by the nucleic acid of (a);
   (f) an antibody, an aptamer or another receptor against the nucleic acid of (a) or the polypeptide of (d) or (e) and
   (g) an anti-sense oligonucleotide of the nucleic acid of (a).

30. Use of an effector of an Mrk polypeptide for the manufacture of an agent for the prophylaxis, treatment or diagnosis of diseases and disorders related to body-weight regulation and thermogenesis, for example, but not limited to, metabolic diseases such as obesity, as well as related disorders such as eating disorder, cachexia, diabetes mellitus, hypertension, coronary heart disease, hypercholesterolemia, dyslipidemia, osteoarthritis, gallstones, and sleep apnea, and disorders related to ROS defence, such as diabetes mellitus, neurodegenerative disorders, and cancer, e.g. cancers of the reproductive organs.

31. The use of claim 30, wherein the effector is of an Mrk2 or Mrk1 polypeptide.

32. The use of claim 30 or 31, wherein the effector is selected from staurosporine or pyrazole derivatives.
33. The use of any one of claims 30-32, wherein the effector is CGP57380 or a derivative thereof.
Figure 1. Triglyceride Content of Drosophila Lk6 Mutants
FIGURE 2. Molecular organisation of the Drosophila Lk6 gene (GadFly Accession Number CG17342)

Legend: ■ GadFly, DSC ■ Magpie, clot
FIGURE 3B. CLUSTAL W (1.82) multiple sequence alignment of human mnk2 proteins

XM_030637.3
AP237775
NM_017572.1

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XM_030637.3
AP237775
NM_017572.1

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XM_030637.3
AP237775
NM_017572.1

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XM_030637.3
AP237775
NM_017572.1

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XM_030637.3
AP237775
NM_017572.1

**********

XM_030637.3
AP237775
NM_017572.1

**********
FIGURE 3C. CLUSTAL W (1.82) multiple sequence alignment of human mkn1 proteins

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
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AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1

XM_001600.2
NM_003684.2
AB00409.1
FIGURE 3D. Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 2a

1  cggtcccccct ccccgctggc gggcgcggga cagaagatgg tgcaagaaga accacgcaag
  61  ctgcaggtt tcaccgcttc gttcagggg cagaacctct tcgaagggccc tcgttcctca
 121  gagcagcggcc accaagcggag ctcgtgcaatt ggccgtgata gcctcgccgg cctggcacttg
 181  cccggtcgcct ccacccatgta ccacccgggg ggcgaagctg gaagagagcg
 241  gggcgggcca cccgagactt ctcgggccag ttggagagct tctacacgtg cggcagagat
 301  gtggcggggg aggcgcctca tcgccgagtt gcacactgtca tcaacctgtc caccagcgcag
 361  gatgacgccc tcgaaactcat gtaaagacgc ccagggccaa ttggagagct gtttggaggag
 421  gatgagggag ctgctgttac aatgcagagg gcgcctgagt tcctgactgc tgcgggtgttc
 481  ttcgagggagg aagcctcgtt ttcggagga gtcggaggg gaccgtctcc ctgctcctcg
 541  aggcacatcc acaagagcgag gcacccctgcac gacgtggaggg cccagcgtgt ggtgagacgac
 601  gtggcgcagcct cctggtacttc tgcgaataac aagcccatag cccagcggaa cctaaagggg
 661  gaaaaactactc tetggcggct cccacagctgc ttcccccccct ggaagatctg tgttttcctg
 721  ttcgggagct gcctataact ccaacggggaac ggctcctctt tcgtccaccct gcggattgctt
 781  actcctcgcc gtcggggcgg gatcactgcc gcggaggtgg tggagccgttc gcgcagaggg
 841  gctagactct cccagcagcgc ctggcagctc tggagcctgg gcctctccttg tattctgcc
 901  ctggcgcggtt cccccgccttg gggcgccgct tggcgcgcgg acttgccgccct gcggcggcgg
 961  gatggcgcgct ctcggcctta gcacatctct ggctgcgaggg ccaagcctcgc ccgctgcgcag
1021  ttcgggagcc aaggccctgc cccctgtggt ctggcgccgag cggccgcaag tgcagcgttgg
ttcgcggaggtt ccacccagctgc tccgagggcag gcagagctgg
1081  ctcggcggtgc cggccagaag ccgccgggctg cccggcgcgg cccgggctgg gcctgcgcagc
1141  cccgggtgag cccggcgcgg aaccaccctt gcgggaggg cccgggaggg gcctgcgcgcgc
1201  gcgcagcggc ttcgggctcct cccgtggtgc gcggcgggctg gcggcggcgg gcctgcgcgcg
1261  cggcagcggc accctggcgg gcggcggcgc gcggcggcgc gcggcggcgc gcctgcgcgcg
1321  gcctgcttgcc gcctgtccct ccctggcagc ccaagctggc gcggcggcgc gcggcggcgc
1381  cacaagctggc gcctgtctcc gcggcggagc gcgtggctgg gcagcagcgg ctcacccctgc
1441  cato

FIGURE 3E. Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 2a

MVQKKPAELQGFRSFKQNSFLAPSDQPDHCDSDFGLQCSA
RPDPASQPIDIPDAKKRKGGKGRATDSFGRFEDVQLQ8DVLGEGAHARVQTCI
NLITSGBYAVKIIKQPGHRISRFREVEMYLGQGHNRNELE1EFEEFEDRFYLFTE
KMRGSISLISHHIKRPHNRELAYSUQDVASALDLHINGIARHDLKPNLCEHPNQ
VSPVKICDFDLGSGIKLNGDCSPISTPELLPGESAYMAPBEVEAFSEASEIYDKRC
DLMSLVILYIILSYPYFPGRCGSDCGWRBEACPCAQNMLFESIQBGKYEFPDKD
AHISCAAKDLISKLLVRDADKRLSAAQVQLHPPQSCAPENTLPFMVLQRNSCKDL
TSFALLAEIAMRNRQLAQHBDLAEABAAGQQGPVPLVRSATRCLSPLPSSQSKLAQRRQR
ASLSSAVFVVLGDHA
FIGURE 3F. Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 2b

gctgcccggc cccccagaca aagatgtgca gaaagaacac cggacacttc agggtttca
gtgtgcc tcacccccaga cccctccgca gctggccttc ttcctagacc acgcgcgacca
cggagacttct gacctggccc tggcagcagc agccgcccct cagcaagcgc ggccgcccct
cgctgttcg cagcgtttgg aagaagtagt gccagctgct gccagcagtt cggagatct
cggtgccgga cagcagcttgc acacggcgca cgtgcgttgg gggagatctt cggagatctt

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

ggaacccag ccccttgcag cccccgcccag cggccccccg cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

cggagactt gcagccggcc
cgggtgcgtc ctcctggttc tctggtggtgc cggagatcgc cggagatcgc

FIGURE 3G. Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 2b

MVKKPAELQGFHRSFKQNPFLAFSLLQDQPDHGDSDDFGLQCASA
RPMPASQPIDIIPADKRRKGRKKRGRATDSFSGRFEDVQLQEDVLGBAHRVQTICI
NLITSKYVAKKIKQPGHRSVREEMLYQCOQHVRNLVELEEFEEDRFYLFVFE
KMRRGSRSLHSHRRFNLBASVQQVDASLDFMLHNRKIAHRDLKPNLCEHPNQ
VSPVCKICDFDLGSGIKLNGDCSPISTPELTPCGSAEYMAPVEVVEAFSEBASTYDKRC
DLSLGVILYILSSGYPFFVRCGSDGCWDRGECACPQCNMLFESIQGKYEFPDKW
AHISCAAKDLIKLLVLRDKAQRQLSAAQVLQHPWQGCAQPENTLPMTPMVLQRWDHPLL
PFPCHRIVPGLGVRTVTVNE
FIGURE 3H. Nucleic acid sequence of human MAP kinase-interacting kinase (Mnk) 1

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1 ggcacggaggg cgacgcgtcc ccgacgcgag ccagcgaaggg ttctcatgtc agagggcgagat
61 ggagactaag aagttgcacg ttcacgcacaa agggcagtaa gacacctctcg gtacctggaa
121 gacaccaact tctcgacagg agcttttatf cattgggtat ttcagatgta cagatggatat
181 cttctcacaag gttgggaaact tatagagaga ggggccgggg cacttgactc ctggccagaa
241 atgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
301 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
361 ttcagttgct gcctgacgct taatgacttg ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
421 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
481 gacaccaact tctcgacagg agcttttatf cattgggtat ttcagatgta cagatggatat
541 cttctcacaag gttgggaaact tatagagaga ggggccgggg cacttgactc ctggccagaa
601 atgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
661 cctctcacaag gttgggaaact tatagagaga ggggccgggg cacttgactc ctggccagaa
721 ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
781 gtcacccac gcctgacgct taatgacttg ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
841 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
901 gcctgacgct taatgacttg ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
961 gcctgacgct taatgacttg ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
1021 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
1081 gcctgacgct taatgacttg ctggccagaa aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
1141 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
1201 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
1261 aatgtgagcaag gagaagagaa aagaaagcggg ggagggcgagct cacttgactc ctggccagaa
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9/34
FIGURE 3I. Amino Acid sequence of human MAP kinase-interacting kinase (Mnk) 1

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SSTMDLTLFAEATIALNLQSHAQENELAAEPEALADGLCSMKLSPCKSRLARKRAL
AQAGRGERDRESPPTAL
Figure 4. Overexpression of the Drosophila UCP homologue dUCPy in the eye.
FIGURE 5A. Expression of Mnk2 in different mouse tissues
FIGURE 5B. Expression of Mnk2 in different mouse models
Figure 5C. Expression of Mnk2 in adipose tissue of different mice strains
FIGURE 5D. Expression of Mnk2 during differentiation of 3T3-L1 cells from pre-adipocytes to mature adipocytes.
FIGURE 5E. Expression of Mnk2 during differentiation of cultured 3T3-F442A cells
FIGURE 5F. Expression of MnK2 during differentiation of cultured TAA1 cells

TA1
(MnK2
(DCT(d0) = 13.90))

Days
0  2  4  6  8  10

Relative RNA-Expression
16,00  14,00  12,00  10,00  8,00  6,00  4,00  2,00  0,00

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FIGURE 6 A. Expression of Mnk1 in different wildtype mouse tissues
FIGURE 6B. Expression of Mnk1 in different mouse models.
FIGURE 6C. Expression of Mnk1 during differentiation of cultured 3T3-L1 cells from pre-adipocytes to mature adipocytes.
FIGURE 6D. Expression of Mnk1 during differentiation of cultured 3T3-F442A cells
FIGURE 6E. Expression of Mnk1 during the differentiation of cultured TA1 cells
Figure 7. Sequences of the Drosophila UCPy

FIGURE 7A. Sequence showing the full length cDNA of Drosophila UCPy

FIGURE 7B shows the deduced amino acid sequence of Drosophila UCPy encoded by the underlined sequence shown in Figure 7A
Figure 8A shows reduction in cellular triglyceride levels (µg/mg protein) in cells over expressing Mnk2 compared to control cells.
Figure 8B shows reduction in insulin-stimulated lipid synthesis (dpm/mg protein) in cells over expressing Mnk2 compared to control cells.
Figure 8C shows reduction in active transport (AT) of free fatty acids across the plasma membrane of cells over expressing Mnk2 compared to control cells.
Figure 9A Expression of human Mnk2 in different human tissues
Figure 9B. Expression of human Mnk2a and human Mnk2b during adipocyte differentiation.
Figure 12. Intron-exon structure of the mouse MNK2 cDNA (GenBank accession number BC010256; 3080bp)

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ratio Erk2:Mnk2a

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anti-activated Mnk

Coomassie

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