AMP-activated Protein Kinase
Up-regulates Mitogen-activated Protein (MAP) Kinase-interacting Serine/Threonine Kinase 1a-dependent Phosphorylation of Eukaryotic Translation Initiation Factor 4E*

AMP-activated protein kinase (AMPK) is a molecular energy sensor that acts to sustain cellular energy balance. Although AMPK is implicated in the regulation of a multitude of ATP-dependent cellular processes, exactly how these processes are controlled by AMPK as well as the identity of AMPK targets and pathways continues to evolve. Here we identify MAP kinase-interacting serine/threonine protein kinase 1 (MNK1) as a novel AMPK target. Specifically, we show AMPK-dependent Ser353 phosphorylation of the human MNK1a isoform in cell-free and cellular systems. We show that AMPK and MNK1a physically interact and that in vivo MNK1a Ser353 phosphorylation requires T-loop phosphorylation, in good agreement with a recently proposed structural regulatory model of MNK1a. Our data suggest a physiological role for MNK1a-Ser353 phosphorylation in regulation of the MNK1a kinase, which correlates with increased eIF4E phosphorylation in vitro and in vivo.

Balancing catabolic and anabolic processes is fundamental to energy homeostasis and metabolic adaptation. The αβγ heterotrimeric AMP-activated protein kinase (AMPK)3 promotes ATP production and limits ATP consumption (1, 2). Shifts in the cellular AMP:ATP ratio are detected through the γ subunit, which cooperatively binds AMP. Consequential conformational changes within the heterotrimer stimulate AMPK kinase activity. Activation of AMPK further involves Thr172 phosphorylation in the activation loop of the α subunit kinase domain (3). Additional (auto-) phosphorylation events are known to regulate AMPK (4–6). Through its established roles in lipid, glucose, and protein metabolism, AMPK functions at the crossroads of energy metabolism and basic cellular processes including cell proliferation, growth, and survival (7–9). For example, AMPK-mediated regulation of acetyl CoA carboxylases ACC1 and ACC2 accelerates fatty acid synthesis and inhibits fatty acid oxidation, respectively (10). AMPK is involved in control of cell growth via phosphorylation of regulatory-associated protein of mTOR (mechanistic target of rapamycin, Raptor) and tuberous sclerosis 2 (TSC2, tumor suppressor) (11, 12).

Despite its well known role as energy sensor, a lack of knowledge on the identity of direct AMPK targets and thus of pathways involved in these processes persists. A substantial number of functional screening approaches have been developed to identify AMPK downstream targets and interaction partners (13–17). High-density protein microarrays enable the rapid identification of potentially novel human kinase substrates at a proteomic scale (18). Using this strategy, we identified MAP kinase-interacting serine/threonine protein kinase 1 (MNK1) as a putative novel AMPK target (19). The human MKNK1 gene encodes two splice variants, MNK1a and MNK1b, which differ in their C-terminal sequences. In contrast to other MNKs, MNK1a has low basal activity and is highly induced upon activation (20). MNK1a is activated by mitogen- and stress-activated protein kinases (M/SAPK), ERK (extracellular signaling-regulated kinase), and P38, via phosphorylation of two threonine residues (Thr209 and Thr214) within the activation/T-loop (21, 22). MNK1 and MNK2 isoforms phosphorylate the eukaryotic translation initiation factor and mRNA cap-binding protein 4E (eIF4E) on serine 209 (23–25). Although the exact biological relevance of this phosphorylation event is still under debate, cellular eIF4E plays an important role in the regulation of mRNA translation, in which interaction with the 5′-cap structure of mRNA appears pivotal (26). Deregulation of eIF4E has been linked to tumorigenesis (26, 27); inhibition of protein translation, e.g. via MNK, is being considered for cancer treatment (20).

Here we show that MNK1a is a genuine AMPK target in vitro and in vivo. MNK1a phosphorylation at Ser353 by AMPK increases its kinase activity toward eIF4E phosphorylation. The relevance of our findings for human metabolic and neoplastic conditions is discussed.
Results

**MNK1a Is a Novel AMPK Target in Vitro**—The online tool ScanSite predicted the presence of a putative AMPK phosphorylation motif LQRNSSSTMDL in MNK1a; this domain is absent in MNK1b. The putative AMPK target site in this motif, Ser\(^{353}\), is evolutionarily highly conserved among mammals and lower vertebrates (Fig. 1A). To validate the initial phospho-protein microarray finding, cell-free *in vitro* kinase (IKV) assays were performed using recombinant AMPK and MNK1a. Mass spectrometric analysis of *in vitro* phosphorylated recombinant MNK1a\(^{WT}\) indicated Ser\(^{353}\) as the primary phospho-site in the MNK1a-specific tryptic peptide sequence NSSTMDTLFAEEAIALNR (Ser\(^{353}\) underlined; localization probability: 74%; Fig. 1A). To further probe the specificity of this phosphoevent *in vitro*, the GST tag was also proteolytically removed from recombinant GST-MNK1a to exclude interference by GST tag phosphorylation (28). Both GST-MNK1a\(^{WT}\) and MNK1a\(^{WT}\) protein were phosphorylated specifically and only in the presence of AMPK (Fig. 1B). The absence of radiolabeled MNK1a\(^{WT}\) protein in AMPK-free control reactions suggested that MNK1a\(^{WT}\) phosphorylation *in vitro* was AMPK-dependent and that MNK1a\(^{WT}\) protein did not show auto-phosphorylation under these conditions. A commercially available AMPK substrate antiserum, which detects a common AMPK-phosphorylated LXRXXpS/P-T motif, recognized *in vitro* phosphorylated MNK1a\(^{WT}\), indicating that MNK1a phosphorylation occurred at an AMPK substrate-like motif (Fig. 1C). The absence of detectable MNK1a\(^{WT}\) phosphorylation using a kinase-dead AMPK further corroborated the notion that MNK1 phosphorylation is dependent on the kinase activity of AMPK *in vitro* (Fig. 1D).

To definitively identify the phosphorylated MNK1a residue, a recombinant MNK1a Ser\(^{353}\) to alanine mutant (MNK1a\(^{S353A}\)) was tested in cell-free assays. To exclude the possibility that the neighboring Ser\(^{352}\) residue was phosphorylated by AMPK under these conditions, we also generated a MNK1a\(^{S352A}\) mutant. Although MNK1a\(^{WT}\) and MNK1a\(^{S352A}\) were clearly detected by autoradiography, the MNK1a\(^{S353A}\) mutant protein was not radiolabeled under these conditions (Fig. 1E). Analogously, comparative analysis of MNK1a\(^{WT}\) and MNK1a\(^{S353A}\) showed that MNK1a\(^{S353A}\), in contrast to MNK1a\(^{WT}\), was not detected by the AMPK substrate antiserum (Fig. 1F). These combined data strongly support the idea that AMPK phosphorylates MNK1a\(^{WT}\) at Ser\(^{353}\) *in vitro*.

**Cellular MNK1a Is Phosphorylated at Ser\(^{353}\) upon AMPK Activation**—To establish that MNK1a phosphorylation at Ser\(^{353}\) occurs in living cells, we first compared MNK1a expression levels in different normal diploid and cancer cells. The osteosarcoma cell line U2OS was found to express endogenous MNK1a at a very low level, whereas its AMPK level was relatively high (Fig. 1G). The functional interaction between AMPK and MNK1a was studied in these cells using retrovirally expressed 2PY-MNK1a. Cells were treated with the AMPK-activating compound 5-aminoimidazole-4-carboxamide ribonucleotide (AICAR); AMPK activity was assessed by pACC1 measurement. Although relatively low under control conditions (i.e., non-stimulated or serum-starved), AICAR dramatically increased 2PY-MNK1a\(^{WT}\) phosphorylation at an AMPK substrate-like motif (Fig. 1H). Importantly, in contrast to immunoprecipitated (IP) 2PY-MNK1a\(^{WT}\), IP 2PY-MNK1a\(^{S353A}\) mutant protein did not show any detectable phosphorylation (Fig. 1H). These data are congruent with cellular AMPK-mediated phosphorylation of MNK1a at Ser\(^{353}\).

To be able to study the biological context and relevance of MNK1a phosphorylation at Ser\(^{353}\), we generated a polyclonal antiserum specifically recognizing Ser\(^{353}\)-phosphorylated MNK1a. Both phosphorylated recombinant GST-MNK1a\(^{WT}\) and proteolytically released MNK1a\(^{WT}\) were detected by the recombinant MNK1a-Ser(P)\(^{353}\) antiserum (Fig. 1I). The finding that recombinant GST-MNK1a\(^{S353A}\) was no longer detectable by the MNK1a-Ser(P)\(^{353}\) antiserum under similar assay conditions corroborated the specificity of the newly generated MNK1a-Ser(P)\(^{353}\) antiserum (Fig. 1J). To obtain direct evidence that MNK1a-Ser(P)\(^{353}\) phosphorylation occurs in living cells, U2OS cells expressing MNK1a\(^{WT}\) or MNK1a\(^{S353A}\) were treated with AICAR. As expected, AICAR increased AMPK-Thr(P)\(^{172}\) (pAMPK) and ACC1-pS79 (pACC1) in both cell types (Fig. 1K). Importantly, AMPK activation correlated well with increased MNK1a-Ser(P)\(^{353}\) signal, and the MNK1a\(^{S353A}\) mutant was not recognized by the MNK1a-Ser(P)\(^{353}\) antiserum (Fig. 1K).

**MNK1a-Ser\(^{353}\) Phosphorylation Is AMPK-dependent and Involves Direct Physical Interaction**—To obtain genetic evidence for the AMPK dependence of MNK1a-Ser\(^{353}\) phosphorylation, 2PY-MNK1a-transduced mouse embryonic fibroblasts (MEFs) derived from wild type (AMPK\(^{WT}\)) or AMPK\(^{α1/α2}\) double null mutant mice (AMPK\(^{dKO}\)) were treated with either AICAR or A769662, the latter being an AMPK activator with a distinct mode of action (29, 30). Both AICAR and A769662 enhanced pAMPK and pACC1 phosphorylation, which correlated well with enhanced MNK1a-Ser\(^{353}\) phosphorylation in AMPK\(^{WT}\), but not in AMPK\(^{dKO}\) MEFs (Fig. 2A). These data convincingly support the notion that MNK1a is phosphorylated at Ser\(^{353}\) in an AMPK-dependent fashion *in vivo*. We then asked whether AMPK physically interacts with MNK1a in living cells. Endogenous AMPK was IP from 2PY-MNK1a\(^{WT}\) U2OS cells. The interaction between MNK1a was evident under basal conditions and increased dramatically upon AMPK activation by AICAR or glucose deprivation (Fig. 2B). Accordingly, AMPK-MNK1a complex formation is induced in response to AMPK-activating stimuli.

**Metabolic Stress-induced MNK1a-Ser\(^{353}\) Phosphorylation Controls Its Kinase Activity toward eIF4E**—To chart the consequences of AMPK-mediated MNK1a-Ser\(^{353}\) phosphorylation status as a function of time, 2PY-MNK1a\(^{WT}\) U2OS cells were glucose-deprived and AMPK and MNK1 phosphorylation was monitored. MNK1a-Ser(P)\(^{353}\) and pACC1 were both induced upon glucose deprivation; both phosphorylation events correlated well with AMPK activation (Fig. 2C). Relevantly, 1 h after replenishment of glucose, cells down-regulated MNK1-Ser(P)\(^{353}\) and pACC1, in parallel with pAMPK (Fig. 2C). These results are strongly suggestive of direct dynamic control of reversible MNK1a phosphorylation at Ser\(^{353}\) by AMPK.

AMPK activation aims to conserve cellular ATP by tuning energy-consuming processes such as mRNA translation to metabolic conditions. MNK1 is known to phosphorylate the eukaryotic initiation factor 4E (eIF4E) at Ser\(^{209}\), a rate-limiting...
ACCELERATED COMMUNICATION: AMPK Augments eIF4E Phosphorylation

A

| Protein | Pos. | Gene | Score | Phospho (STY) Probability | - ATP | + ATP |
|---------|------|------|-------|---------------------------|-------|-------|
| Q9BUB5-2 | 353  | MKNK1| 257.86| NS(0.24)S(0.74)T(0.02)MDTLFAAEALN | 0 | 1.00E+08 |

B

C

D

E

F

G

H

I

J

K
component of the translation apparatus (31). To explore the potential functional relevance of AMPK-mediated MNK1a-Ser\(^{353}\) phosphorylation in regard to eIF4E Ser\(^{209}\)-Ser\(^{209}\) phosphorylation (pelF4E), we compared pelF4E phosphorylation status in the U2OS model transduced with MNK1a\(^{S353A}\) or MNK1a\(^{S353D}\) mutants. Cells expressing MNK1a\(^{S353D}\) showed almost 2-fold increased pelF4E phosphorylation at basal levels when compared with cells expressing MNK1a\(^{WT}\) or MNK1a\(^{S353A}\) (Fig. 2D). Combined, these data suggest AMPK-dependent stimulation of MNK1a toward its downstream target eIF4E, via MNK1a-Ser\(^{353}\) phosphorylation.

MNK1-Thr\(^{209}/\text{Thr}^{214}\) are phosphorylated within the activation T-loop by the M/SAPKs ERK and P38 (21, 22). The C terminus of MNK1a, harboring Ser\(^{353}\), is thought to play a dual role in the regulation of MNK1a kinase activity: the C terminus acts repressive in the basal state while allowing full stimulation upon ERK/P38-dependent phosphorylation of Thr\(^{209}/\text{Thr}^{214}\) (21). To explore the potential functional interdependence of Thr\(^{209}/\text{Thr}^{214}\) and Ser\(^{353}\) phosphorylation, we studied specific MNK1a-phospho-mutants in U2OS cells in combination with pharmacological stimulation rather than metabolic stress, as this allows separation of upstream signaling events. AICAR treatment alone or in combination with TPA (M/SAPK activator), consistently and highly induced pACC1, whereas TPA treatment alone only modestly induced pACC1 (Fig. 2E). TPA, with or without AICAR, massively increased MNK1a-Thr\(^{P}(209)/\text{Thr}^{P}(214)\) (MNK1a-pTpT) (Fig. 2E); these observations are in good agreement with reported upstream signaling events connecting M/SAPKs to MNK1a (22). Combination treatment with TPA and AICAR resulted in increased MNK1a-Ser\(^{P}(353)\), but only in the case of MNK1a\(^{WT}\) and MNK1a\(^{K^{-1}}\) (K78M; kinase-dead). Reduced Ser\(^{353}\) phosphorylation in U2OS MNK1a\(^{K^{-1}}\) cells may be explained by consistently lower expression of the MNK1a\(^{K^{-1}}\) mutant (Fig. 2E). As expected, the T209A/T214A (MNK1a\(^{T2^{-2A2}}\)) mutant was no longer phosphorylated at the mutated phospho-sites, and neither IP MNK1a\(^{T2^{-2A2}}\) nor MNK1a\(^{K^{-1}}\) mutant kinases phosphorylated pelF4E (Fig. 2E). Remarkably, MNK1a\(^{T2^{-2A2}}\) was no longer phosphorylated at Ser\(^{353}\), whereas conversely, Ser\(^{353}\) mutation (S\(\rightarrow\)A or S\(\rightarrow\)D) did not block the ability of Thr\(^{209}\)/Thr\(^{214}\) to become phosphorylated (Fig. 2E). IP MNK1a\(^{WT}\) from AICAR only-treated cells led to only marginally increased pelF4E, in support of the need for regulatory co-signaling by M/SAPKs (Fig. 2E). In contrast, and in good agreement with the observations above (cf. Fig. 2D), MNK1a\(^{S353D}\) consistently displayed enhanced pelF4E under AICAR only- or TPA only-treated conditions; moreover, eIF4E phosphorylation by MNK1a\(^{S353D}\) was already enhanced under non-stimulated conditions (Fig. 2E). Of note, pACC1 was not altered by MNK1a mutation. The combined data suggest that MNK1a selectively acts in the signaling downstream of AMPK, leading to increased pelF4E phosphorylation, and point toward a hierarchical order of regulatory events within MNK1a: M/SAPK-mediated MNK1a-Thr\(^{209}/\text{Thr}^{214}\) phosphorylation precedes AMPK-dependent MNK1a-Ser\(^{353}\) phosphorylation.

**Discussion**

In this study, we report that MNK1a is a direct AMPK target. We provide biochemical and genetic evidence that MNK1a-Ser\(^{353}\) is phosphorylated in *vitro* and *in vivo* in an AMPK-dependent manner and that AMPK and MNK1a physically interact. Finally, we show that MNK1a-Ser\(^{P}(353)\) correlates with increased eIF4E phosphorylation *in vitro* and *in vivo*.

Members of the MAPK-activated protein kinase (MK) family fulfill multiple biological roles in cellular responses to extracellular cues (e.g. mitogenic stimulation and cellular stress). MNK1a, like other MAPKAPKs (MAP kinase-activated protein kinases), is tightly regulated by M/SAPKs, including ERK and P38 (32, 33). The most well characterized biochemical connection in regard to MNK1 is eIF4E-Ser\(^{209}\)-serine phosphorylation (24), although the exact relevance of MNK1-mediated eIF4E phosphorylation remains unclear (34). MNK1a mutation appears to affect phosphorylation of eIF4E, not of ACC1, suggesting a selective role for MNK1a as a signaling intermediate in the regulation of protein synthesis, downstream of AMPK.

Based on a previously proposed regulatory model of MNK1a, our data suggest a physiological role for Ser\(^{353}\) phosphorylation in regulating MNK1a activity. MNK1a is activated upon T-loop phosphorylation, but also undergoes a conformational change to a more “open” structure (21). Our MNK1a mutant analysis indicating that Thr\(^{209}/\text{Thr}^{214}\) phosphorylation is a prerequisite for Ser\(^{353}\) phosphorylation appears congruent with this phospho-event being dependent on conformational change. Interestingly, an α-helical structure from the first part of the unique MNK1a

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**FIGURE 1.** MNK1a-Ser\(^{353}\) is a novel AMPK target. A. *Upper panel,* evolutionary conservation of putative MNK1a phosphorylation site. Sequence alignments of MNK1a proteins of human (Q9BUB5), mouse (008605), rat (Q4G050), bovine (Q58D94), pig (B8XSK1), and Western clawed frog (Q66JF3) (www.uniprot.org) are shown. *Lower panel,* identification of MNK1a-kinase-phosphate (aa 351–369: Swiss-Prot: Q9BUB5-2). Intensities of phospho-peptide M5 spectra in the presence or absence of ATP (±ATP) are shown. Probabilities for each phosphorylatable residue are indicated between brackets in the aa sequence. B. IKV analyses. Purified GST-MNK1a protein was incubated with or without active AMPK in the presence of \[^{32}P\]ATP. Protease-mediated removal of the GST tag was done as indicated. *Upper panel,* pMNK1a autoradiograph (autorad). *Lower panel,* CB staining image (used as loading control). C. IKV analyses using puriﬁed recombinant GST-MNK1a WT protein, active AMPK, and cold ATP. D. IKV analyses using purified GST-MNK1a WT, cold ATP, and active AMPK. E. IKV analyses using purified GST-MNK1a WT, cold ATP, and active AMPK. F. As in C. IKV analyses were performed with cold ATP. IB detection was done using the indicated antisera. G. MNK1a and AMPK protein levels in HeLa cervical adenocarcinoma, U2OS, ME7, TIG3 human primary fibroblasts, H629 human embryonic kidney, and HL-1 immortal murine cardiomyocytes. IB analyses of total cell extracts using the induced antisera were performed. H. U2OS cells expressing 2PY-MNK1a WT or MNK1a\(^{S353A}\) (control: empty vector (EV)) were treated as indicated (cf. control conditions; ss: serum-starved; A: ss + AICAR). MNK1a was IP using PY antisera; IB analyses of immunoprecipitation and input material were performed using the indicated antisera. J, as in C. IKV analyses using purified recombinant GST-MNK1a \(^{K^{-1}}\) protein, active AMPK, and cold ATP were performed. Protease-mediated removal of the GST tag was performed as indicated in the figure. IB of phosphorylated MNK1a were performed using the indicated antisera. J, as in H. IKV analyses using purified GST-MNK1a \(^{K^{-1}}\) or GST-MNK1a\(^{S353D}\) with active AMPK and cold ATP were performed. IB was done using the indicated antisera. K. U2OS cells expressing 2PY-MNK1a WT, 2PY-MNK1a\(^{S353A}\), or transduced with empty vector (EV) were treated as indicated (serum-starved (ss) or ss + AICAR (A)).
C-terminal region was predicted to suppress basal T-loop phosphorylation and activity (21); Ser$^{353}$ is located at the start of this region. Based on existing and current data, we propose a functional extension to the MNK1a activation model: T-loop phosphorylation by canonical M/SAPK pathway activity precedes AMPK-mediated phosphorylation of MNK1a at Ser$^{353}$ (Fig. 2F).

Our observations suggest that MNK1a-Ser$^{353}$ phosphorylation by AMPK is not required for MNK1a activation per se, but
Affects and/or possibly directs kinase activity, upon a conformational change induced by M/SAPK. MNK1-Ser(P)\textsuperscript{353} may serve to fine-tune MNK1-eIF4E-mediated protein synthesis (mRNA translation) via AMPK. This finding has potential therapeutic implications for chronic or acute human conditions, including metabolic syndrome-associated disorders and cancer. Abnormal regulation of protein synthesis is known to drive tumor cell proliferation and survival. Importantly, both AMPK and MNK1a have been independently put forward as potential druggable targets (35, 36).

**Experimental Procedures**

**Expression Vectors**—For bacterial expression of MNK1a protein, glutathione-S-transferase(GST)-tagged MNK1a (pGEX-6P1) was kindly provided by C. G. Proud (Southampton, UK). Active, hexahistidine-tagged (His\textsubscript{6}) AMPK was produced induced in 50\% active AMPK protein and 4\% antisera. Immunoprecipitations were used in IVKs with recombinant GST-eIF4E and [\textsuperscript{32}P]ribosomal protein S\textsubscript{28} (\textsuperscript{35}). AMPK and three LKB1 subunits as described (5). Expression of kinase-dead AMPK\textalpha1D\textsuperscript{157A} mutant was described earlier (37). Point mutations were introduced using QuikChange (Agilent). To generate eukaryotic expression vector, MNK1a cDNAs were subcloned in frame with an N-terminal double polyoma (2PY) tag into the BamHI and Sal sites of a retroviral PBS vector (38). Amphiphatic retrovirus was produced as described (39); transduced U2OS and MEF cells were selected in medium containing 4\% puromycin.

**Recombinant Protein Purification and Cell-free Kinase Assays**—Proteins were expressed in Rosetta 2 (DE3) Escherichia coli cells (Merck Millipore) as described (5). His\textsubscript{6}-AMPK and GST-MNK1a protein were purified using nickel-Sepharose HP (GE Healthcare) and glutathione-Sepharose 4B (GE Healthcare), respectively. AMPK protein was eluted in elution buffer (50 mM NaH\textsubscript{2}PO\textsubscript{4}, 30\% glycerol, 0.5 M sucrose, 250 mM imidazole, pH 8) MNK1a protein was eluted with freshly prepared buffer (50 mM Tris-HCl, 10 mM reduced glutathione, pH 8).

Aliquots of resin-bound MNK1a protein (10 \mu g) were incubated in 50 \mu l of kinase buffer (50 mM Hepes, pH 7.4, 5 mM MgCl\textsubscript{2}, 0.04 mM AMP, 1 mM DTT, 200 \mu M ATP) with 1 \mu g of active AMPK protein and 4 \mu g of [\textsuperscript{32}P]ATP for 15 min at 37 \degree C. Beads were collected by centrifugation (5 min, 800 \times g), washed, and suspended in cleavage buffer (50 mM Hepes, pH 7.5, 150 mM NaCl, 0.5 mM EDTA, 1 mM DTT). For proteolytic removal of GST tags, resin-bound protein was incubated with PreScission protease (GE Healthcare) for 4 h at 4 \degree C with constant mixing. Beads were pelleted (5 min, 800 \times g); supernatant was transferred to a fresh tube. Samples were heated in SDS-PAGE sample buffer (5 min, 95 \degree C). Samples were analyzed by SDS-PAGE (9\% gels) electrophoresis followed by Coomassie Brilliant Blue (CBB) staining and/or autoradiography.

**Mass Spectrometry**—GST-MNK1a and AMPK proteins were incubated with or without ATP in vitro as described above. Samples were separated by SDS-PAGE; MNK1a bands were excised and in-gel digested as described (40). Extracted tryptic peptides were desalted using homemade C18 columns, resuspended in 10\% formic acid, and analyzed by reversed phase nanoLC-MS/MS (Ultimate 3000 and Orbitrap Elite; Thermo Scientific). Peptides were trapped on a precolumn for 10 min (Acclaim PepMap100, C18, 5 \mu m, 100 \AA, 300-\mu m inner diameter \times 5 mm, Thermo Scientific) in Buffer A (0.1\% formic acid in water) and separated on an analytical column (Acclaim PepMap100, C18, 5 \mu m, 100 \AA, 75-\mu m inner diameter \times 25 cm) using a 70-min gradient (0–10 min, 5\% buffer B (80\% acetonitrile, 0.1\% formic acid); 10–45 min, 10–45\% buffer B; 45–47 min, 45–99\% buffer B; 47–53 min, 99\% buffer B; 53–70 min, 5\% buffer B) at 250 nl/min. The mass spectrometer was operated in data-dependent mode with a 20-s dynamic exclusion range. Full-scan MS spectra were acquired in the Orbitrap (range: m/z 350–1500) with a resolution of 120,000 and an automatic gain control of 16E ions. Collision-induced dissociation fragmentation in the ion trap was performed on the top five precursors of each full scan employing a collision energy of 35\%. Analysis of raw data was done by using MaxQuant (version 1.4.1.2 (41)). The spectra were searched against the human Swiss-Prot database version 06/2014 using the Andromeda search engine using default mass tolerance settings (42). Trypsin was set as the protease (two missed cleavages allowed). Fixed modification: carbamidomethylation (Cys); variable modifications: oxidation (Met), phosphorylation (Ser, Thr, Tyr), and N-terminal protein acetylation. False discovery rate was set to 0.01 for peptides, proteins, and modification sites. The minimum peptide score for modified peptides was set to 40, and the minimum peptide length was seven amino acids (aa).

**Cell Culture, Chemicals, and Antibodies**—Human osteosarcoma U2OS cells and immortal mouse embryo fibroblasts (AMPK\textsuperscript{WT} and AMPK\textsuperscript{diKO} MEFs (43)) were cultured in DMEM (25 mM glucose; Gibco), supplemented with 10\% (v/v) heat-inactivated fetal calf serum (Bodinco BV, Alkmaar, The Nether-
erlands) and penicillin/streptomycin (Invitrogen). Cellular AMPK activation was achieved with AICAR (1.5 mM; Sigma) or A769662 (100 μM; Tocris Biosciences) for 45–60 min following 16 h of serum starvation (DMEM, 25 mM glucose). For M/SAPK activation, TPA (100 μM) was used following serum starvation. For glucose deprivation, DMEM medium without glucose (Gibco) was used. Antibodies used are: total MNK1a (tMNK1a) (#2195), pMNK1 (Thr209/Thr214, #2111), AMPK-Thr(P)172 (#9411), and pERK (#9101; Cell Signaling Technology); pACC (#87-303; Merck Millipore); β-actin (bACT, #0869100; MP Biomedicals); and eIF4E (#610269; BD Biosciences). Rabbit polyclonal MNK1a Ser353 phosphorylation-specific antibodies were custom-ordered (Genosphere Biotechnologies, Paris, France).

Immunoprecipitation and Immunoblotting—Cells were lysed in mild lysis buffer (250 mM NaCl, 0.1% IGEPAL (Nonidet P-40), 5 mM EDTA, 50 mM HEPES, pH 7.0) supplemented with 5 mM benzamidine, 5 μg/ml antipain, 5 μg/ml leupeptin, 5 μg/ml aprotinin, 1 mM sodium vanadate, 10 mM sodium fluoride, 10 mM pyrophosphate, 10 mM β-glycerophosphate, 0.5 mM DTT, and 1 mM PMSF. Extracts were sonicated on ice and centrifuged; 3% of the supernatant was stored as input (−80 °C). Endogenous AMPK was IP using a mix of AMPKα1 and AMPKα2 antibodies (sheep polyclonal; kindly provided by D. G. Hardie, Dundee, Scotland, UK); 2PY-tagged MNK1α (2PY-MNK1α) was IP using PY antisera (Covance). For immunoprecipitation, 500 μg of protein lysate were incubated with primary antibody under constant mixing at 4 °C overnight, followed by incubation with protein G-Sepharose beads (4 h, 4 °C). Immune complexes were collected by centrifugation and analyzed by immunoblotting as described (5). Signals were detected using enhanced chemiluminescence (ECL; Pierce). Band intensities were quantitated with the Quantity One software (Bio-Rad). Data were statistically analyzed by performing two-tailed paired t tests using Microsoft Excel.

eIF4E Kinase Assay—350 μg of U2OS 2PY-MNK1α cell lysates were incubated with PY antibody (2 h, 4 °C); 40 μl of protein G slurry were added for 1.5 h (4 °C). Beads were pelleted (5 min, 800 × g) and washed twice with mild lysis buffer, twice with 0.5 M LiCl, and twice with kinase buffer as described previously (21). Beads were re-suspended in 50 μl of kinase buffer and incubated (15 min, 37 °C) with 1 μg of recombinant GST-eIF4E. The reaction was terminated by adding SDS-PAGE sample buffer (5 min, 95 °C). Samples were analyzed by SDS-PAGE (9% gels) and autoradiography. 200 μg of cell lysates were processed in parallel with immunoprecipitation antibody as controls.

Author Contributions—X. Z., J. W. V., and D. N. designed the study; X. Z., V. D., R. T., C. P., B. V., J. W. V., and D. N. contributed to data acquisition, analysis, and interpretation; X. Z., J. W. V., and D. N. drafted the manuscript; X. Z., and V. D., R. T., C. P., B. V., J. W. V., and D. N. provided final approval of the submitted version.

Acknowledgments—We thank R. Stead (Southampton, UK) and members of the Molecular Genetics Department (Maastricht University) for helpful comments and support. The Proteomics Facility is funded by the IZKF Aachen.

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