MICAL-1 Is a Negative Regulator of MST-NDR Kinase Signaling and Apoptosis

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MICALs (molecules interacting with CasL) are atypical multidomain flavoenzymes with diverse cellular functions. The molecular pathways employed by MICAL proteins to exert their cellular effects remain largely uncharacterized. Via an unbiased proteomics approach, we identify MICAL-1 as a binding partner of NDR (nuclear Dbf2-related) kinases. NDR1/2 kinases are known to mediate apoptosis downstream of the mammalian Ste-20-like kinase MST1, and ablation of NDR1 in mice predisposes the mice to cancer as a result of compromised apoptosis. MST1 phosphorylates NDR1/2 kinases at their hydrophobic motif, thereby facilitating full NDR kinase activity and function. However, if and how this key phosphorylation event is regulated are unknown. Here we show that MICAL-1 interacts with the hydrophobic motif of NDR1/2 and that overexpression or knockdown of MICAL-1 reduces or augments NDR kinase activation or activity, respectively. Surprisingly, MICAL-1 is a phosphoprotein but not an NDR or MST1 substrate. Rather, MICAL-1 competes with MST1 for NDR binding and thereby antagonizes MST1-induced NDR activation. In line with this inhibitory effect, overexpression or knockdown of MICAL-1 inhibits or enhances, respectively, NDR-dependent proapoptotic signaling induced by extrinsic stimuli. Our findings unveil a previously unknown biological role for MICAL-1 in apoptosis and define a novel negative regulatory mechanism of MST-NDR signaling.

Protein kinases are key regulatory enzymes that control important cellular processes such as growth and apoptosis by changing the properties of a substrate through phosphorylation (20). Given the importance of their biological effects, the catalytic activity of protein kinases is stringently controlled, and defects in this spatiotemporal regulation can cause major diseases such as diabetes and cancer (20). The mammalian Ste-20-like kinase MST1 is part of evolutionary conserved signal transduction pathways with functions in different biological processes of cells. In particular, Hippo signaling (the MST1 fly ortholog) has been studied extensively in Drosophila melanogaster, revealing that Hippo tumor suppressor cascades are crucial in the regulation of cell death and proliferation (45). More recently, mammalian MST1 was established as a tumor suppressor protein controlling cell proliferation and apoptosis (45). However, despite these important functions, relatively little is known about the regulation of MST1 or its downstream signaling.

Nuclear-Dbf2-related kinase 1 (NDR1; also known as STK38) and NDR2 (STK38L) can be phosphorylated by MST kinases and mediate apoptotic signaling downstream of MST1 (40). NDR kinases belong to the AGC kinase subfamily and control important cellular processes, such as mitotic exit and apoptosis, in different eukaryotic cells ranging from yeast to neurons (12). Ablation of NDR1 in mice predisposes the mice to the development of T cell lymphoma, presumably as a result of compromised apoptosis (4). The regulation of NDR1/2 kinases by MST kinases and other upstream signaling proteins is best understood in mammalian cells. Binding of MOB1A (Mps-one-binder 1A) to the N-terminal region of NDR1/2 stimulates autophosphorylation on the activation segment and activates NDR kinases (3). The human MOB protein family consists of six distinct members with MOB1 being best studied for its putative tumor suppressive functions through the regulation of NDR/LATS kinases. Additional phosphorylation on the C-terminal hydrophobic motif of NDR1/2 by MST kinases is required for full NDR kinase activation (32). It is evident that these different molecular interactions and activating events need to be tightly controlled, as deregulation of members of the MST, NDR, and MOB protein families has been implicated in diseases such as cancer. Intriguingly, another MOB family member, MOB2, competes with MOB1 for NDR1/2 binding at the N-terminal region, thereby negatively regulating NDR1/2 kinase activity and related functions in centrosome duplication and apoptosis (16). However, whether similar sophisticated regulatory mechanisms exist to control the activation of NDR1/2 by MST kinases is unknown.

Here we identify the multidomain flavoprotein monoxygenase MICAL-1 (MICAL stands for molecule interacting with CasL) as a negative regulator of MST-NDR kinase signaling and apoptosis.
with CasL (37) as an endogenous binding partner of NDR1/2 kinases. Our findings define a previously unknown biological role for MICAL-1 in apoptosis regulation and show that MICAL-1 negatively controls MST1-NDR apoptotic signaling by competing with MST1 for NDR binding, thereby revealing a novel and unique regulatory mechanism of the MST-NDR pathway. This work opens new avenues for the further molecular delineation of MST/NDR functioning in different cell biological processes and in disease.

MATERIALS AND METHODS

Antibodies and reagents.

The generation and purification of anti-T444/442-P and anti-S281/282-P antibodies has been described previously (38). Both antibodies were kindly provided by Brian Hemmings (Friedrich Miescher Institute for Biomedical Research [FMI], Basel, Switzerland). It is important to note that the anti-T444-P and anti-S281-P antibodies recognize the phosphorylated forms of both NDR isoforms, NDR1 (T444-P and S281-P) and NDR2 (T442-P and S282-P). A polyclonal rabbit anti-mouse MICAL-1 antibody was generated by immunizing rabbits with a maltose-binding protein (MBP)-mouse MICAL-1 (mMICAL-1)-amino acid 962 to 1048 fusion protein, which was purified from E. coli DH5α cells. The specificity of this antibody was confirmed using Western blotting and immunocytochemistry. We used the following commercially available antibodies: anti-Flag (M2; Stratagene), antimyc, and anti-HA (3F10; Roche), anti-NDR1 (N1-14; Santa Cruz), anti-STK38 (NDR1) monoclonal antibody (2G18-1F3; Abnova), CH1 (Millipore), anti-glutathione S-transferase (anti-GST) (Cell Signaling), anti-myHC (Cell Signaling), anti-green fluorescent protein (anti-GFP) (Invitrogen), anti-V5 (Invitrogen), anti-α-tubulin (Sigma), anti-actin (Sigma), anti-cleaved caspase-3 (Cell Signaling), and anti-poly(A)-ribosome polymerase (anti-PARP) (Asp241; BD Biosciences). Okadaic acid (OA) was purchased from Enzo Life Sciences, and etoposide, tumor necrosis factor alpha (TNF-α), and cycloheximide (CHX) were purchased from Sigma.

Construction of plasmids.

Mouse MICAL-1 and NDR1 cDNAs were amplified from embryonic whole-brain cDNA using standard molecular techniques. To construct a mouse MICAL-1-Flag/HA construct for generating stable L cell lines, mouse MICAL-1 cDNA was subcloned into a pBabeFlag-HA-puromycin destination vector using the Gateway cloning system (Invitrogen). Mouse MICAL-1 cDNA was also subcloned into pRK5-HA and pRK5-myc (N-terminal tags) using Sall and NotI restriction sites, into pEF-His/V5 vector (C-terminal tag) using the Gateway cloning system (Invitrogen), and into pEGFP (EGFP stands for enhanced green fluorescent protein). Transfection of mouse MICAL-1 was cloned via PCR. Individual PCR products were digested with Sall and NotI and cloned into pRK5-HA or pRK5-myc. Mouse MICAL-1 constructs harboring glycine-to-tryptophan mutations (G3/W3) in the first FAD stands for enhanced green fluorescent protein). Truncation mutants of mouse MICAL-1 cDNA was also subcloned into pRK5-HA and pRK5-myc (N-terminal tag) using a short hairpin RNA (shRNA) vector targeting both NDR isoforms, NDR1 (T444-P and S281-P) and NDR2 (T442-P and S282-P). A polyclonal rabbit anti-mouse MICAL-1 antibody was generated by immunizing rabbits with a maltose-binding protein (MBP)-mouse MICAL-1 (mMICAL-1)-amino acid 962 to 1048 fusion protein, which was purified from E. coli DH5α cells. The specificity of this antibody was confirmed using Western blotting and immunocytochemistry. We used the following commercially available antibodies: anti-Flag (M2; Stratagene), antimyc, and anti-HA (3F10; Roche), anti-NDR1 (N1-14; Santa Cruz), anti-STK38 (NDR1) monoclonal antibody (2G18-1F3; Abnova), CH1 (Millipore), anti-glutathione S-transferase (anti-GST) (Cell Signaling), anti-myHC (Cell Signaling), anti-green fluorescent protein (anti-GFP) (Invitogen), anti-V5 (Invitrogen), anti-α-tubulin (Sigma), anti-actin (Sigma), anti-cleaved caspase-3 (Cell Signaling), and anti-poly(A)-ribosome polymerase (anti-PARP) (Asp241; BD Biosciences). Okadaic acid (OA) was purchased from Enzo Life Sciences, and etoposide, tumor necrosis factor alpha (TNF-α), and cycloheximide (CHX) were purchased from Sigma.

Cell culture and transfection.

L. PhoNix, HEK293, and COS-7 cells were maintained in high-glucose Dulbecco’s modified Eagle’s medium (DMEM; Gibco) supplemented with 10% fetal bovine serum (FBS) (Lonza), penicillin-streptomycin, and l-glutamine (PA4) at 37°C with 5% CO2. U2OS T-Rex (tetracycline-inducible MST1 knockdown) cells were cultured as described previously (38). To induce tetracycline-controlled transcriptional activation and MST1 knockdown, 5×10⁶ cells were plated in one well of a six-well plate and 2 μg/ml tetracycline was supplied to the culture medium. For protein analysis, exponentially growing cells were plated at 3.5×10⁵ cells/cm² and transfected the next day using Lipofectamine 2000 (Invitrogen) as described by the manufacturer. For COS-7 cell transfection assays or colocalization experiments, 0.5×10⁴ COS-7 cells/cm² were seeded on glass coverslips and transfected using FuGENE transfection reagent (Roche) according to the manufacturer’s instructions. For transfection of MICAL-1 siRNA oligonucleotides, exponentially growing HEK293 cells or L cells were seeded at 5×10⁴ per well in a 6-well plate. After overnight culture, cells were washed with Opti-MEM (Invitrogen) and exposed to a mixture of 1 μl oligonucleotides (mix) (from a 20 μM stock) and 3 μl Oligo- fectamine (Invitrogen) in Opti-MEM. Four hours after transfection, the medium was replaced with standard culture medium, and the cells were incubated at 37°C with 5% CO2 for another 48 to 72 h. For double-knockdown experiments with both siRNA oligonucleotides (against MICAL-1) and sirna vector (against NDR1/2) 1×10⁶ L cells were plated in a 24-well plate. All clones of MICAL-1 oligonucleotide using Oligofectamine. After 24 to 48 h in culture, the cells were transfected with sirna vector by using Lipofectamine 2000 and cultured for another 48 h for further treatment.

Generation of stable cell lines.

A mouse MICAL-1 cDNA was introduced by Gateway cloning into a retroviral destination plasmid derived from pBabe-puro (adenovirus for transfection). HA carrying a C-terminal Flag/HA tag was transiently transfected using Lipofectamine 2000. Mouse MICAL-1-Flag/HA constructs for generating stable L cell clones expressing tagged MICAL-1 were obtained by retroviral transduction and puromycin (Clontech) selection. Positive clonal cell lines were identified by immunoblotting using antibodies directed against HA. After puromycin selection, clonal lines were maintained in culture medium containing 7.5 μg/ml puromycin. L cells with integrated empty pBabe-puro plasmid served as a control.

Pulldown from stable cell lines.

Mouse cells expressing MICAL-1-Flag/HA (MICAL-1 tagged with Flag or HA) (D1) and control (D9) L cells were cultured in 500-cm² plates (Corning) to 90% confluence. Ten plates from each cell line were harvested by scraping the cells after 2 washes with cold phosphate-buffered saline (PBS). The cells were centrifuged at 1,000 rpm for 15 min at 4°C and lysed by Dounce homogenization in 3 ml of hypotonic lysis buffer containing 10 mM HEPES (pH 8.0), 1.5 mM MgCl₂, 10 mM KCl, 0.5 mM phenylmethylsulfonyl fluoride (PMSF), 0.05 mM dithiothreitol (DTT), and protease inhibitor cocktail (Sigma). After the cells were centrifuged at 14,000 rpm for 15 min at 4°C, the supernatant was transferred to a clean Falcon tube, and an equal volume of 2× salt buffer containing 30 mM HEPES (pH 8.0), 190 mM NaCl, 0.4% glycerol, 0.4 mM EDTA, 0.5 mM PMSF, 0.5 mM DTT, and protease inhibitor cocktail (Sigma) was added. After measuring protein concentration, 100 μg of protein from each cell line was added to 100 μl prewashed M2 affinity resin (Sigma) and incubated at 4°C for 3 h. The resin was then washed 4 times with washing buffer (BC-150) containing 500 mM NaCl, 0.5 mM PMSF. Between the third and fourth wash, the sample was washed once in washing buffer (BC-300) containing 20 mM HEPES (pH 8.0), 300 mM NaCl, and 0.5 mM PMSF for 10 min. The proteins were eluted with Flag peptide (Sigma) at 16 μg/ml in BC-150 buffer. The eluted proteins were precipitated as described previously (43) and dissolved in 1× NuPAGE LDS sample buffer (Invitrogen). Finally, proteins were separated on a NuPAGE Novex 4 to 12% Bis-Tris gradient gel following the manufacturer’s description (Invitrogen). Proteins were stained using the colloidal blue staining kit (Invitrogen) and processed for mass spectrometry. In addition, 1/20 of each sample was loaded on another NuPAGE gel and processed for silver staining. Briefly, the gel was soaked twice in 50% methanol for 15 min and soaked in 5% methanol for 10 min. After 3 rinses in water, the gel was soaked in 10 μM DTT for 20 min, followed by 0.1% AgNO₃ for 20 min. The gel was then washed once in water and twice in developer containing 3% sodium carbonate and 0.0185% formaldehyde. The gel was soaked in developer until protein bands appeared. The reaction was stopped with 5% acetic acid (about 5% [w/vol]). The gel was washed with water and scanned.

Gel digestion.

Separated proteins were digested in the gel by the method of Shevchenko et al. (29). Gel lanes corresponding to the different protein samples were sliced into 10 bands. Each band was cut into 1-mm cubes, washed with nanopure water, and destained (25 mM ammonium bicarbonate and 50% acetonitrile). After three subsequent destaining steps, the bands were dehydrated for 20 min (100% acetonitrile) and dried in a vacuum centrifuge. The pellets were rehydrated in 30 μl trypsin solution (0.02 μg/μl) (Promega) at 4°C for 45 min, followed by the addition of 400 μl of 50 mM ammonium bicarbonate to
cover the gel pieces. After incubating overnight at room temperature, peptides were extracted two times in 50% acetonitrile in 0.1% trifluoroacetic acid (TFA). The sample was dried in a vacuum centrifuge prior to liquid chromatography coupled to matrix-assisted laser desorption ionization mass spectrometry (LC–MALDI-MS) analysis.

NanomLC and MALDI TOF/TOF analysis. Samples were analyzed as described previously (22) with minor modifications. In brief, peptides were separated on a nanocapillary LC (nanoLC) system (LC Packings/Dionex, Sunnyvale, CA) using an analytical capillary C18 column (150 mm by 100 μm inner diameter [ID]) at 400 nL/min, using a linear increase in the concentration of acetonitrile from 5% to 50% (vol/vol) in 90 min and to 90% in 10 min. The eluted gradient was mixed with matrix solution (7 mg of recrystallized c-α-cyanohydroxyceinamic acid in 1 mL of 50% (vol/vol) acetonitrile, 0.1% (vol/vol) trifluoroacetic acid, 10 mM ammonium dihydrogen phosphate) and spotted off-line to a stainless steel MALDI plate (Applied Biosystems) to form a predefined 16 by 24 array (384 spots) using a ProBiot system (LC Packings/Dionex). The mass spectrometric analysis was carried out using a MALDI tandem time of flight (MALDI-TOF/TOF) instrument (4800 Proteomics Analyzer; Applied Biosystems) with reflector positive ion mode. For MS analysis, an m/z mass range of 800 to 3000 was used, with 1,250 shots per spectrum. A maximum of 20 precursors per spot with minimum signal/noise ratio of 50 were selected for data-dependent tandem MS (MS/MS) analysis. The 2.5 keV collision energy was used for collision-induced dissociation (CID), and 2,500 acquisitions were accumulated for each MS/MS spectrum. All analyses were performed using default calibration, and the mass accuracy was calibrated to within 100 ppm using calibration standards (Applied Biosystems) before each run.

Cell contraction assay. COS-7 and L cells were transfected as described previously (36). In brief, COS-7 or L cells were incubated in the dark, and destained in a solution containing 20% acetonitrile and 50 mM ammonium bicarbonate (Invitrogen) were either Coomassie blue stained or stained with Pro-Q diamond fluorescence (Supersignal; Thermo Scientific) to form a recently discovered and unusual family of evolutionary conserved cytoplasmic proteins composed of an enzymatically active flavoenzyme followed by several protein interaction domains or motifs (18). One MICAL gene has been identified in Drosophila (Mical), while humans and mice have three different MICAL genes (MICAL-1, MICAL-2, and MICAL-3) (26, 37, 39). Genetic inactivation of Mical in fruit flies leads to defects in neural circuit development, myofilament organization, and bristle formation (2, 13, 15, 39). Although Drosophila

RESULTS

Identification of novel MICAL-1-binding partners. MICAlS form a recently discovered and unusual family of evolutionary conserved cytoplasmic proteins composed of an enzymatically active flavoenzyme followed by several protein interaction domains or motifs (18). One MICAL gene has been identified in Drosophila (Mical), while humans and mice have three different MICAL genes (MICAL-1, MICAL-2, and MICAL-3) (26, 37, 39). Genetic inactivation of Mical in fruit flies leads to defects in neural circuit development, myofilament organization, and bristle formation (2, 13, 15, 39). Although Drosophila

centrifugation at 14,000 rpm for 10 min at 4°C, the lysate was transferred to a clean Eppendorf tube and boiled with sample buffer. For immunoprecipitation, the corresponding antibodies were added to each cell lysate and incubated overnight at 4°C. Then prewashed protein A- or G-agarose (Roche) was added to the lysate and incubated for at least 2 h at 4°C. After 4 washes with lysis buffer, proteins were eluted with sample buffer and boiled at 90°C. To analyze proteins from immunoprecipitated (IP) samples or cell lysates, we used SDS-PAGE and transferred proteins to a nitrocellulose (Hybond-C Extra; Amersham) Chemiluminescence (Supersignal; Thermo Scientific) was detected by CL-XPosure film (Thermo Scientific) and quantified using ImageJ.

Recombinant protein purification and binding assay. MBP-MICAL-1-C1 (amino acids 734 to 1048) fusion protein was extracted from E. coli DH5α cells, and the MICAL-1-C1 fragment was cleaved from the MBP moiety following the instructions of the manufacturer (New England BioLabs). To check the binding of GST-NDR2 to MICAL-1, HA-tagged MICAL-1 (HA-MICAL-1) (HEK293 cells expressed full-length protein) or MICAL-1-C1 was immobilized on anti-HA- or anti-MICAL-1-coupled protein G-agarose beads, respectively, and mixed with 4 μg GST-NDR2. Antibody-coupled beads were used as negative control to examine the nonspecific binding. After 2 h of incubation at 4°C, beads were washed 4 times with lysis buffer and boiled in the presence of sample buffer. The proteins were then subjected to SDS-PAGE and Western blot analysis.

Kinase assays. (i) NDR kinase peptide assay. Flag-tagged NDR1 (Flag-NDR1) was immunoprecipitated from cell lysates expressing Flag-NDR1 alone or in combination with HA-MICAL-1 and assayed for NDR kinase activity in the presence of 1 mM NDR substrate peptide (KKRRNLRLSVA), 100 μM [γ-32P]ATP (50 Ci/mmol), and 20 μM ATP in kinase buffer (20 mM Tris [pH 7.5], 10 mM MgCl2, 1 mM benzamidine, 4 μM leupeptin, 3 μM microcystin, 3 mM DTT, 1 μM cyclic AMP-dependent protein kinase inhibitor peptide). The reaction mixture was incubated for 60 min at 30°C and stopped by adding 50 mM EDTA (pH 7.5). The supernatant of the mixture was spotted onto squares of P-81 phosphocellulose paper (Whatman) and washed 4 times with 1% orthophosphoric acid followed by an aceton wash. Radioactivity was measured in a liquid scintillation counter.

(ii) GST-NDR2 kinase assay using HA-MICAL-1 as a substrate. Full-length recombinant NDR2 was cloned into pGEX-T 3’ in frame with GST, expressed in E. coli BL21(DE3) (pLyS3) cells with 0.03 mM isoprropyl-β-d-thiogalactopyranoside (IPTG) induction overnight at 25°C, and purified by glutathione-Sepharose affinity chromatography. For in vitro kinase assays, 3 μg of GST-NDR2 fusion protein was incubated with immunoprecipitated HA-MICAL-1 in the presence of 100 μM [γ-32P]ATP (50 Ci/mmol) and 20 μM ATP in kinase buffer for 30 min at 30°C. The reaction was stopped by adding SDS sample buffer, and protein phosphorylation was analyzed by SDS-PAGE and subsequent autoradiography.

(iii) In vitro phosphorylation with GST-MST1. Recombinant GST-MST1 was obtained from Sigma. To produce immunopurified HA-tagged proteins, COS-7 cells were transfected and processed for immunoprecipitation using anti-HA antibody as described earlier (30). Immunopurified proteins were then washed twice with MST1 kinase buffer (5 mM Tris [pH 7.5], 2.5 mM β-glycerophosphate, 1 mM EGTA, 1 mM Na3VO4, 4 mM MgCl2, 0.1 mM DTT), before incubating at 30°C for 30 min in 20 μl of reaction buffer (5 mM Tris [pH 7.5], 100 μM ATP, 2.5 mM β-glycerophosphate, 1 mM EGTA, 1 mM Na3VO4, 4 mM MgCl2, 0.1 mM DTT, 10 μCi of [γ-32P]ATP [3,000 Ci/mmol; Hartmann Analytic] in the absence or presence of GST-MST1 (100 ng per reaction mixture). The reactions were stopped by the addition of Laemmli buffer, before the proteins were separated by SDS-PAGE. Subsequently, total proteins were visualized by Coomassie blue staining, the gels were then dried, and phosphorylated proteins were visualized by autoradiography.
Mical has been implicated in the control of F-actin assembly (13), the molecular pathways employed by MICAL proteins to exert their diverse cellular effects remain largely uncharacterized, especially in vertebrate species. To obtain more insight into the MICAL signaling complex and related functions, we used a retroviral vector system to generate L cell lines that stably express a Flag-and-HA-tagged (Flag/HA-tagged) version of mouse MICAL-1 (MICAL-1-Flag/HA) (Fig. 1A), the best-characterized vertebrate MICAL thus far (37). Mouse L cell fibroblasts were used because of their murine origin, their endogenous MICAL-1 expression and the ability of a constitutively active MICAL-1 construct to induce morphological changes in these cells (data not shown). This suggests that L cells contain signaling proteins that are crucial for MICAL-1 function. Clonal cell lines were obtained by retroviral transduction followed by puromycin selection, resulting in several clones expressing MICAL-1-Flag/HA at near endogenous levels. One of these lines (D1) was used for large-scale affinity purification of MICAL-1-Flag/HA complexes followed by silver staining, in-gel tryptic digestion, and mass spectrometry analysis. L cells with integrated empty retroviral vector served as a control (C0) (Fig. 1B). The mass spectrometry analysis identified several proteins that were found in protein complexes from pulldown assays using D1 but not C0 cell extracts. The most significant hits included the GTP-binding cytoskeletal protein septin-9 (42), the serine-threonine kinase NDR1 (12), and the actin-binding protein filamin-B (35) (Table 1).

**TABLE 1. Binding partners of MICAL-1 in stable L cells identified by mass spectrometry**

<table>
<thead>
<tr>
<th>Protein</th>
<th>Mascot score</th>
<th>NCBI GI no.</th>
<th>Protein MM* (kDa)</th>
<th>No. of unique peptides</th>
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<td>46396473</td>
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<td>264</td>
</tr>
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<td>Septin-9</td>
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<td>65.5</td>
<td>12</td>
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<td>NDR1</td>
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<td>54.1</td>
<td>8</td>
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<tr>
<td>Filamin-B</td>
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<td>38257404</td>
<td>277.6</td>
<td>8</td>
</tr>
</tbody>
</table>

*The table shows proteins identified with a significant Mascot score in pulldown experiments from D1 cell extracts. No peptides were identified for the indicated proteins in a parallel pulldown from C0 control cells.

**FIG. 1.** Mass spectrometry identifies NDR kinases as novel MICAL-1-binding partners. (A) Schematic depicting the domain organization of mouse MICAL-1 fused to the C-terminal Flag and HA epitopes linked by a PreScission sequence (MC-1-Flag/HA). FBD, FAD-binding domain; CH, calponin homology; PPKP, Casl-binding sequence; CC, coiled-coil. (B) Pulldown assays with anti-Flag monoclonal antibody-conjugated agarose beads were performed on lysates from D1 or C0 cells. Bound proteins were eluted with Flag peptide and analyzed by silver staining. The positions of MICAL-1 and NDR1 on the gel are indicated to the left of the gel. (C) Anti-Flag pulldown assays were performed from D1 and C0 cells as shown in panel B and analyzed by Western blotting (WB) using the indicated antibodies (α-Flag, anti-Flag antibody). IP, immunoprecipitated. (D and E) Lysates from HEK293 cells transfected with epitope-tagged MICAL-1 alone (+) or in combination with Flag-NDR1 (D) or Flag-NDR2 (E) were immunoprecipitated with anti-Myc (α-Myc, anti-Myc antibody). (F and G) Recombinant GST-NDR2 and HA-MICAL-1 purified from HEK293 cells (F) or recombinant MICAL-1-C1 purified from bacteria (G) were mixed and subjected to immunoprecipitation with anti-HA (F) or anti-MICAL-1 (G) antibodies followed by Western blotting with the indicated antibodies. (C) Immunoprecipitation from untransfected L cell extracts followed by Western blotting with the indicated antibodies. Anti-myc and anti-HA antibodies were used as a control for anti-NDR1 and anti-MICAL-1 antibodies, respectively.
the D1 lane following anti-Flag pulldown (Fig. 1C). Immuno-
precipitation experiments from HEK293 cells coexpressing
myc-tagged MICAL-1 (myc-MICAL-1) and Flag-tagged
NDR1 (Flag-NDR1) further confirmed the interaction of
MICAL-1 with NDR1 (Fig. 1D). Using several of the available
antibodies, we were unsuccessful in staining for endogenous
NDR1 or NDR2 in cells (see also references 8 and 10). How-
ever, fluorescence microscopic analysis of COS-7 cells trans-
fected with epitope-tagged MICAL-1 and NDR1 showed that
the distribution of both proteins partially overlapped in the
cytoplasm and in specific membrane regions (data not shown).

Two NDR kinases have been identified, NDR1 and NDR2
(12). NDR1 and NDR2 are 87% identical at the amino acid
level, and one of the peptides identified in our mass spectrom-
etry analysis was shared between NDR1 and NDR2 (data not
shown). It is therefore possible that MICAL-1 also binds to
NDR2. To test this idea, HA-tagged MICAL-1 (HA-MICAL-1)
and Flag-tagged NDR2 (Flag-NDR2) were coexpressed in
HEK293 cells followed by pulldown with anti-Flag antibody.
Indeed, HA-MICAL-1 coprecipitated with Flag-NDR2 (Fig. 1E). A similar result was obtained following commounprecip-
ation of recombinant GST-NDR2 and HA-MICAL-1 protein purified from HEK293 cells (Fig. 1F). To determine
whether the interaction between MICAL-1 and NDR1/2 was
direct, the C-terminal region of MICAL-1 (C1; Fig. 2A) was
purified from bacteria and used in a coimmunoprecipitation
experiment with GST-NDR2. The ability of GST-NDR2 to
coprecipitate MICAL-1 C1 indicates a direct interaction
between NDR2 and the C-terminal region of MICAL-1
(Fig. 1G). Finally, we performed coimmunoprecipitation ex-
periments from naïve L cell lysates using MICAL-1- and
NDR1-specific antibodies. Importantly, endogenous MICAL-1 coprecipitated with endogenous NDR1 and NDR2 and vice
versa from naïve L cell extracts, indicating that in L cells
endogenous MICAL-1/NDR1 complexes can be formed
(Fig. 1H and data not shown).

MICAL-1 and NDR1 interact through domains involved in
the regulation of their enzymatic activity. Both MICAL-1 and
NDR1 are multidomain proteins. MICAL-1 contains an N-termin-
al flavoprotein monoxygenase (FM) domain, a calponin

FIG. 2. MICAL-1 and NDR1 interact through domains involved in the regulation of their enzymatic activity. (A) Schematic representation of
the domain structure of full-length (FL) mouse MICAL-1 and its truncation mutants. The numbers in parentheses and above the schematic
representation are the amino acid positions. (B) Lysates of HEK293 cells transfected with Flag-NDR1 (+) and HA-tagged full-length or truncated
MICAL-1 were immunoprecipitated (IP) with anti-HA antibody (α-HA) and analyzed by Western bloting (WB). (C) Microdensitometry of
Flag-NDR1 coimmunoprecipitated with HA-MICAL-1 (full-length or truncation mutants) from three independent experiments similar to those
shown in panel B. Signal intensity is shown in arbitrary units (a.u.). Values are means plus standard deviations (SD) (error bars). (D) Schematic
depicting NDR1 and its truncation mutants with conserved domains indicated. The positions of the Ser281 and Thr444 regulatory phosphorylation
sites are shown. HM, hydrophobic motif; NTR, N-terminal regulatory region. Numbers indicate amino acid positions. (E) Lysates of HEK293 cells
transfected with HA-MICAL-1 and Flag-tagged full-length or truncated NDR1 were immunoprecipitated with anti-HA antibody and analyzed by
Western blotting. (F) Microdensitometry of Flag-NDR1 coimmunoprecipitated with HA-MICAL-1 (+) from three independent experiments
similar to those shown in panel E. Data are means plus SD. (G) Schematic of the identified interactions between MICAL-1 and NDR1. The LIM
and C-terminal domains of MICAL-1 interact with the C-terminal region of NDR1 harboring the hydrophobic motif and the regulatory Thr444
phosphorylation site.
homology domain, a LIM domain, proline-rich sequences, and coiled-coil motifs (Fig. 2A). NDR kinases contain an N-terminal regulatory (NTR) domain, a catalytic domain, and a C-terminal hydrophobic motif (HM) (Fig. 2D). To determine the domains required for MICAL-1/NDR1 interactions, a series of truncation mutants were generated for MICAL-1 (Fig. 2A) (28) and NDR1 (Fig. 2D) (10) and used in pulldown assays from HEK293 cells coexpressing mutants for MICAL-1 or NDR1 with full-length NDR1 or MICAL-1, respectively. Co-precipitation of endogenous MICAL-1 or NDR1 proteins was not detected in these experimental settings (data not shown). Intriguingly, constructs containing the LIM (N3, C3, and C2) and/or C-terminal domain (C3 to C1) of MICAL-1 showed binding to NDR1, indicating that both domains are required for association with NDR1 (Fig. 2B and C). Since intramolecular interactions between the LIM and C-terminal domains of MICAL-1 have been reported to enforce an autoinhibited protein conformation preventing flavoenzyme activity (28), this suggests that NDR1 might bind MICAL-1 in its autoinhibited state. Conversely, the NDR1 C-terminal region was required for interaction with MICAL-1 (Fig. 2E and F). This is intriguing because the NDR1 C-terminal region harbors the hydrophobic motif surrounding the Thr444 site, whose phosphorylation by MST kinases is required for full NDR1 kinase activity (23, 32, 38). In all, our results reveal an endogenous interaction between MICAL-1 and NDR1. Furthermore, NDR1 associates with two domains of MICAL-1 crucial for regulating its enzymatic activity, while reciprocally, MICAL-1 binds a region of NDR1 contributing to the control of NDR kinase activity (Fig. 2G).

**MICAL-1 negatively regulates NDR1 kinase activation.** To unveil the functional role of MICAL-1/NDR1 interactions, we first explored the idea that NDR kinases regulate MICAL-1. Thus far, two properties of MICAL-1 have been characterized in detail. First, MICAL-1 is an NADPH-dependent flavoprotein monooxygenase with redox activity (25, 30). Second, C-terminally truncated MICAL-1 proteins (N1 to N3 [Fig. 2A]) induce cell contraction in culture (28). This contraction response is most likely dependent on the FM domain and on redox activity, as mutagenesis of essential glycine residues in the MICAL-1 FM domain leads to loss of N1-induced cell contraction (data not shown). Intramolecular interactions between the LIM and C-terminal domains of MICAL-1 can inhibit FM activity (28). Because NDR1 associates with both domains, it is tempting to speculate that NDR kinases regulate MICAL-1 activity. To test this hypothesis, an enzyme-linked assay was used to determine the effect of NDR1/2 on MICAL-1 enzymatic activity based on H2O2 production (Fig. 3A and B) (25, 28). In line with previous observations, full-length (FL) MICAL-1 produced only low levels of H2O2, whereas constitutively active MICAL-1 N1-N3 truncation mutants showed high enzymatic activity (Fig. 3A to D) (25, 28). In a recent study, it was shown that cotransfection of the putative MICAL-1 substrate collapsin response mediator protein-2 (CRMP-2) with MICAL-1 in cells leads to a reduction in MICAL-1 N1-N3 H2O2 enzymatic activity. Similarly, recombinant GST-CRMP-2 induced a reduction in MICAL-1 N3 enzymatic activity (28). It was proposed that rather than producing H2O2, MICAL-1 may perform redox reactions on CRMP-2 in the presence of this substrate (28). In contrast to these findings, MICAL-1 activity was not inhibited by the cotransfection of NDR1 or by the addition of recombinant NDR2. Furthermore, NDR1/2 did not alter the enzymatic activity of FL MICAL-1 (Fig. 3C and D). In addition, shRNA-mediated knockdown of NDR1/2 in L cells did not affect the enzymatic activity of MICAL-1 N3 (Fig. 3E). Previous work has shown that cotransfection of FL MICAL-1 with CRMP-2 or plexin A1 induces COX-7 cell contraction (28). However, coexpression of NDR1/2 or a constitutively active form of NDR1 (NDR1-PIFtide [33]) with FL MICAL-1 did not induce cell contraction, and NDR1/2 overexpression or knockdown did not influence contraction triggered by N3 (Fig. 3F and G; also data not shown). These experimental results argue against a role for NDR1/2 in the control of MICAL-1 enzymatic and contraction activity.

We next determined whether MICAL-1 could serve as a substrate for NDR kinases. Although we were able to establish that MICAL-1 is a phosphoprotein (Fig. 4A), GST-NDR2 failed to phosphorylate MICAL-1 in an in vitro kinase assay, indicating that MICAL-1 is not an NDR2 substrate (Fig. 4B). Unfortunately, production of GST-NDR1 protein is notoriously difficult, and we were unable to obtain sufficient quantities of NDR1 for kinase assays. In summary, our results strongly argue against a direct role for NDR1/2 in regulating MICAL-1 enzymatic activity and the subsequent cell morphological changes.

Therefore, we next examined whether MICAL-1 may regulate NDR kinases. Multisite phosphorylation is a general control mechanism of NDR1 and NDR2, which contain two main regulatory phosphorylation sites: Ser281/282 (NDR1/2, the activation segment) and Thr444/442 (the hydrophobic motif) (Fig. 2D). Upon MOB-1A binding to the N-terminal region, Ser281/282 is autophosphorylated, whereas Thr444/442 is targeted by upstream MST kinases (12). Treatment of cells with okadaic acid (OA), which preferentially inhibits protein phosphatase 2A, results in increased Ser281/282 and Thr444/444 phosphorylation and elevated NDR1/2 kinase activity (3, 32, 38). To study the potential effects of MICAL-1 on NDR1/2, we treated HEK293 cells transfected with Flag-NDR1 alone or in combination with HA-MICAL-1 with OA and assessed Ser281 and Thr444 phosphorylation of precipitated Flag-NDR1 protein using phospho-specific antibodies (Fig. 5A). Sixty minutes of OA treatment strongly induced Ser281 and Thr444 phosphorylation (data not shown). MICAL-1 overexpression did not influence OA-induced Ser281 autophosphorylation (Fig. 5A). Consistent with these data, we did not find an effect of MICAL-1 on GST-NDR2 autophosphorylation in kinase assays (Fig. 4B). It should be noted, however, that the currently available phosphorylated Ser281 (p-Ser281) antibody is not suitable for work with endogenous NDR kinases (unpublished data). In striking contrast, overexpression of HA-MICAL-1 drastically inhibited phosphorylation of Flag-NDR1 on Thr444 following OA treatment (Fig. 5A). This effect was dose dependent, as higher levels of HA-MICAL-1 were more efficient in lowering OA-induced NDR1 phosphorylation (Fig. 5B). Furthermore, MICAL-1 reduced Thr444 phosphorylation of endogenous NDR1 and NDR2 following OA treatment (Fig. 5C). It has been reported that at certain concentrations H2O2 can inhibit Thr444 phosphorylation (8). Since the MICAL-1 FM domain can produce H2O2 under specific conditions or potentially facilitate other redox reactions (Fig. 3) (25, 28), we
next asked whether MICAL-1 enzymatic activity mediates the negative effect of MICAL-1 on Thr444 phosphorylation. Site-directed mutagenesis was used to mutate the first FAD fingerprint in the MICAL-1 FM domain. These mutations disrupt FAD binding and block enzymatic activity without affecting the overall structure of the protein (19, 21, 44). MICAL-1 FL G3/W3 could, however, still reduce OA-induced Thr444 phosphorylation, suggesting that the inhibitory effect of MICAL-1 on Thr444 phosphorylation is independent of its enzymatic activity (Fig. 5C).

Next, to establish the physiological relevance of the MICAL-1 effect, the level of endogenous MICAL-1 in HEK293 cells was decreased by siRNA. In line with our previous results, knockdown of MICAL-1 resulted in an increased level of Thr444 phosphorylation triggered by OA compared to control (Fig. 5D). Thus, MICAL-1 is an endogenous negative regulator of NDR1 activation in HEK293 cells. Since phosphorylation of Thr444 is required for full kinase activity, we next examined the ability of MICAL-1 to reduce NDR1 kinase activity on NDR substrate peptide (Fig. 5E). Flag-NDR1 was expressed in HEK293 cells, alone or in combination with HA-MICAL-1, immunoprecipitated, and used in a peptide kinase assay. In line with its negative effect on Thr444 phosphorylation, MICAL-1 robustly inhibited NDR1 kinase activity, reducing peptide phosphorylation by about 40%. In conclusion, our results show that MICAL-1 is an endogenous negative regulator of NDR1 activation that inhibits T444 phosphorylation, thereby contributing to the control of NDR kinase activity.
The phosphorylation status of MICAL-1 was determined following shRNA-mediated knockdown of MST1 in cells. Interestingly, MICAL-1 phosphorylation was not affected by MST1 knockdown (Fig. 6B). To further test whether the regulation of NDR kinases by MICAL-1 involved the phosphorylation of MICAL-1 by MST1 kinase, we analyzed MICAL-1 as an MST1 substrate. This revealed that while kinase-dead NDR1 [NDR1(kd)] was strongly phosphorylated by recombinant MST1 (Fig. 6C, lane 5), purified MICAL-1 was not (Fig. 6C, lane 6), although MICAL-1 was present at higher abundance than NDR1. As expected, active MST1 also autophosphorylated itself (Fig. 6C, lanes 4 to 6), and all phosphorylation events were dependent on the presence of active MST1 (Fig. 6C, lanes 1 to 3). Notably, no effect of MICAL-1 expression on MST1 autophosphorylation was observed. Taken together, these findings suggest that MICAL-1 is not an MST1 substrate, arguing against the possibility that MICAL-1 interferes with phosphorylation of NDR by MST1 by serving as an alternative MST1 substrate.

A third possibility is that MICAL-1 competes with MST kinases for binding on the hydrophobic motif of NDR1. To test this idea, lysates from HEK293 cells expressing Flag-NDR1 and HA-MST1 were mixed with increasing amounts of lysates from HA-MICAL-1-expressing cells followed by anti-Flag immunoprecipitation. Increasing MICAL-1 levels led to a reduction in the amount of NDR1 that coimmunoprecipitated with NDR1 despite similar MST1 input levels (Fig. 7A). As reported previously (11, 40), overexpression of MST1 leads to an increase in Thr444 phosphorylation (data not shown). To examine whether overexpression of MICAL-1 not only reduces MST1 binding but also causes a consequent decrease in MST1-induced Thr444 phosphorylation, we expressed HA-MST1 in combination with increasing amounts of HA-MICAL-1 and monitored Thr444 phosphorylation of endogenous NDR1/2. Indeed, MICAL-1 decreased MST1-induced Thr444 phosphorylation in a dose-dependent manner (Fig. 7B). In all, these results establish that MICAL-1 interferes with the binding of MST1 to NDR1/2, thereby negatively regulating MST1-induced NDR phosphorylation at the hydrophobic motif.

**MICAL-1 interferes with apoptosis signaling.** Since our findings indicate that MICAL-1 negatively regulates MST1-NDR signaling, we sought to determine whether MICAL-1 could affect a known biological function of the MST1-NDR pathway (40). The activation of NDR1/2 kinases through phosphorylation by MST1 is necessary for apoptosis signaling in mammalian cells (40). In addition, loss of NDR1/2 kinases can result in resistance to apoptosis (4). Therefore, wild-type L cells or stable C0 and D1 cells were treated for 12 h with vehicle or etoposide, a reagent known to induce apoptosis through NDR kinases (4), harvested, and analyzed by immunoblotting (Fig. 8A). Etoposide treatment triggered a robust increase in Thr444/442 phosphorylation in wild-type (not shown) and control (C0) cells concomitant with a strong induction of two apoptotic markers, i.e., cleaved PARP and cleaved caspase-3 (Fig. 8A). Furthermore, many cleaved caspase-3-positive C0 cells were detected following etoposide treatment (Fig. 8B). Intriguingly, etoposide exposure of D1 cells expressing MICAL-1-Flag/HA (Fig. 1) did not result in enhanced Thr444/442 phosphorylation (Fig. 8A). In line with this prominent inhibition of NDR activation, the signals for cleaved PARP and cleaved caspase-3 were dramatically lower in D1 cells than in control C0 cells following etoposide treat-
(Fig. 8A), and only a few cleaved caspase-3-positive D1 cells were detected (Fig. 8B), suggesting that apoptotic signaling is inhibited in D1 cells. Similar results were obtained in a second, independent stable line expressing MICAL-1-Flag/HA (clone D2; not shown). In addition, the activation of NDR kinases (phosphorylated Thr444 [pThr444]) and cleavage of caspase-3 and PARP in response to TNF-α/H9251/CHX, another proapoptotic stimulus known to signal through NDR kinases (40), was reduced in D1 cells than in C0 cells (Fig. 8A). CHX alone did not induce apoptosis or affect pThr444 levels (40; also data not shown). Finally, to assess whether the inhibitory effect of MICAL-1 on apoptosis was dependent on its MO domain and redox signaling, we transiently transfected the constitutively active MICAL-1 N3 mutant or the FL MICAL-1 G3/W3 mutant into L cells and treated these cells with etoposide. Similar to FL MICAL-1, MICAL-1 G3/W3 reduced the etoposide-induced cleavage of caspase-3 (Fig. 8C). No effect of MICAL-1 N3 was observed. These results support the idea that the inhibitory effect of MICAL-1 on apoptosis is independent of its MO domain.

We also addressed the role of endogenous MICAL-1 in apoptosis regulation by RNA interference (RNAi). Wild-type L cells transfected with non-targeting control siRNAs or siRNAs specifically targeting mouse MICAL-1 were treated for 12 h with vehicle or etoposide, harvested, and analyzed by immunoblotting (Fig. 8D). In line with our biochemical and overexpression data, knockdown of MICAL-1 resulted in an increase in the overall amount of Thr444/442 phosphorylated G3/W3.
NDR and in increased signals for cleaved PARP and cleaved caspase-3 (Fig. 8D). Although etoposide and TNF-/H9251 can induce apoptosis through the MST1-NDR pathway, it is formally possible that the observed effects of MICAL-1 on apoptosis do not involve modulation of NDR signaling. To address this point, L cells were transiently transfected with siRNAs targeting MICAL-1, shRNA constructs targeting NDR1/2 or combinations thereof, followed by etoposide treatment. Knockdown of MICAL-1 led to increased signals for cleaved caspase-3, while as shown previously (4), knockdown of NDR1/2 inhibited the cleavage of caspase-3 induced by etoposide (Fig. 8E). Interestingly, following simultaneous knockdown of MICAL-1 and NDR1/2, no induction of caspase-3 cleavage was observed, indicating that MICAL-1 and NDR1/2 function in the same pathway. In all, we conclude that MICAL-1 has inhibitory effects on proapoptotic signaling through the NDR-MST pathway.

DISCUSSION

Phosphorylation of NDR1/2 kinases by MST1 is required for full NDR kinase activity and for the induction of apoptosis in response to proapoptotic stimuli (4, 40). The deregulation of NDR1 and MST kinases predisposes the organism to the development of cancer due to compromised apoptotic responses (4, 31). Despite the critical role of MST-induced NDR phos-

FIG. 6. MICAL-1 is not a substrate for MST1 kinase. (A) Lysates from HEK293 cells transfected with HA-MST1 alone or in combination with MC-1-V5 were immunoprecipitated with anti-V5 antibodies followed by Western blotting (WB). (B) Lysates of U2OS T-Rex cells were transfected with HA-MC-1 and treated with vehicle (−) or tetracycline (+) to induce MST1 shRNA expression. The lysates were subjected to immunoprecipitation with anti-HA antibodies followed by Coomassie blue or Pro-Q staining or WB with the indicated antibodies. The graph shows microdensitometry from a representative experiment. (C) Lysates of COS-7 cells transiently expressing empty vector, HA-NDR1 kinase-dead [HA-DNR1(kd)], or HA-MC-1 were subjected to immunoprecipitation with anti-HA antibody. Immunopurified proteins were then processed for kinase assays without or with GST-MST1. The proteins were then separated by SDS-PAGE and visualized either by autoradiography (top panel) or Coomassie blue staining. In parallel, samples were analyzed by Western blotting using anti-HA and anti-MST1 antibodies. The positions of HA-MC-1, GST-MST1, and HA-NDR1 (in kilodaltons) are indicated by arrowheads to the right of the Western blots. The relative molecular weights (Mr) (in thousands [K]) are indicated to the left of the Western blots.

FIG. 7. MICAL-1 competes with MST1 for NDR1 binding. (A) Lysates from HEK293 cells transfected with Flag-NDR1, HA-MST1, or HA-MICAL-1 were mixed as indicated, immunoprecipitated with anti-Flag antibodies, and analyzed by Western blotting (WB). (B) HEK293 cells were transfected with HA-MST1 in the presence of different amounts of HA-MICAL-1. Lysates were subjected to WB with the indicated antibodies.
phorylation in different cell biological processes and in disease, it has remained unknown if or how this key phosphorylation event is regulated. Our findings identify MICAL-1 as a novel endogenous negative regulator of MST1-NDR signaling and apoptosis in mammalian cells (Fig. 8F). MICAL-1 specifically and endogenously interacts with the hydrophobic motif of NDR1/2 (Fig. 1 and 2) and colocalizes with NDR1 in cells, and ectopic expression of MICAL-1 reduces NDR kinase activation and activity (Fig. 5). Knockdown of endogenous MICAL-1, on the other hand, results in increased NDR kinase phosphorylation (Fig. 5). Furthermore, we found that MICAL-1 does not bind MST1 or serve as a MST1 substrate (Fig. 6) but competes with MST1 for NDR binding (Fig. 7) and thereby reduces MST1-induced NDR activation (Fig. 7). In line with this inhibitory effect, overexpression and knockdown studies show that MICAL-1 negatively affects a known biological function of the MST1-NDR1/2 pathway, namely, proapoptotic signaling (Fig. 8). Together, these results uncover a novel and unique regulatory mechanism of MST-NDR signaling.

Binding of MOB1 to the N-terminal region of NDR1/2 stimulates autophosphorylation and NDR activation (3). Kohler and colleagues recently reported that human MOB2 (hMOB2) can act as an inhibitor of hMOB1-NDR signaling by competing with hMOB1 for binding to the N-terminal domain of NDR1/2 (16). Here we describe an unexpected additional level of NDR regulation. By competing with MST1 for binding to the C-terminal hydrophobic motif domain, MICAL-1 can regulate NDR1/2 activation (Fig. 8F). Together, these studies define a novel level of NDR kinase regulation through competitive inhibitors that directly interfere with the binding of key upstream activating signals, such as MOB1 and MST1. How these competitive inhibitors are controlled is unknown. It is possible that previously identified MICAL-1 interactors such as plexin
proteins contribute to this regulation. Plexins function as evolutionary conserved type I transmembrane receptors for semaphorin proteins in different cell types and organ systems and mediate diverse cellular processes, including apoptosis (7, 27, 46). For example, semaphorin 3A can induce neuron or leukemic cell death through plexin A3 and plexin A1, respectively (1, 24). Intriguingly, MICAL-1 can associate with the cytoplasmic domains of the four class A plexins, and semaphorin ligand stimulation enhances this interaction (28) (Y. Zhou and R. J. Pasterkamp, unpublished observations). Recruitment of MICAL-1 to plexin upon ligand stimulation may reduce the availability of MICAL-1 for inhibiting MST-induced NDR activation, leading to enhanced NDR activity and apoptosis. Alternatively, the regulation of MICAL-1 activity by upstream signaling cues such as protein kinases may be crucial for controlling its inhibitory role in the MST-NDR pathway. This idea gains support from the observation that MICAL-1 is phosphorylated in cells and neural tissues (Fig. 4A), suggesting that it is a substrate for upstream kinases. Future studies will address these and other possible mechanisms.

It is plausible that the MICAL-1-dependent regulatory mechanism delineated in this study also functions in MST- and/or NDR-dependent cellular processes other than apoptosis. For example, the MST1-NDR pathway was recently implicated in the control of centrosome duplication (11). Furthermore, striking similarities exist between the effects of manipulating MST, MICAL, and NDR on neuronal morphology. Knockdown of MST3b or overexpression of constitutively active MICAL-1 in cultured mammalian neurons reduces neurite growth (14, 28), while exogenous NDR2 expression enhances neurite growth (34). The loss of Tricornered (NDR) or Hippo (MST) in Drosophila and the loss of SAX-1 (NDR) in Caenorhabditis elegans leads to altered dendritic arborization and tiling defects (i.e., ectopic overlap between individual dendritic trees) (5, 6, 9). Loss of Drosophila Mical also results in enlarged dendritic fields due to deficits in dendritic pruning (15). These observations together with our own findings support a model in which MST, MICAL, and NDR are components of a common molecular pathway that controls the formation and remodeling of the initial imprecise neuronal network into functional neuronal connections.

In the present study, no effect of NDR1/2 on MICAL-1 enzymatic activity or cell contraction was detected, but it is possible that NDR1/2 influences other proposed MICAL effects such as on Rab-mediated vesicle transport (41) or myofilament organization (2). It is also interesting to note that knockdown of MST3b leads to a reduction in CRMP-1, a putative substrate for the MICAL-1 flavoprotein monooxygenase domain (14, 28). Thus, MICAL-1 may not only negatively regulate MST kinases, but MSTs might also influence MICAL-1 function by controlling its substrate levels.

Taking all the results of this study together, this study reveals a previously unknown biological role for MICAL-1 in apoptosis and defines a novel negative regulatory mechanism of MST-NDR signaling. Future experiments are needed to decipher the precise role of MICAL-1 in the activation of NDR kinases by extrinsic and intrinsic apoptotic stimuli. Further elucidation of the role of MICAL-1 in MST-NDR signaling can be expected to open new avenues for the molecular delineation of MST/NDR functioning in different cell biological processes and for understanding how deregulation of this pathway contributes to disease.

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