Novel Transcription Factor-Like Function of Human Matrix Metalloproteinase 3 Regulating the CTGF/CCN2 Gene\textsuperscript{v}

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Matrix metalloproteinase 3 (MMP3) is well known as a secretory endopeptidase that degrades extracellular matrices. Recent reports indicated the presence of MMPs in the nuclei (A. J. Kwon et al., FASEB J. 18:690–692, 2004); however, its function has not been well investigated. Here, we report a novel function of human nuclear MMP3 as a trans regulator of connective tissue growth factor (CCN2/CTGF). Initially, we cloned MMP3 cDNA as a DNA-binding factor for the CCN2/CTGF gene. An interaction between MMP3 and transcription enhancer dominant in chondrocytes (TRENDIC) in the CCN2/CTGF promoter was confirmed by a gel shift assay and chromatin immunoprecipitation. The CCN2/CTGF promoter was activated by overexpressed MMP3, whereas a TRENDIC mutant promoter lost the response. Also, the knocking down of MMP3 suppressed CCN2/CTGF expression. By cytochemical and histochemical analyses, MMP3 was detected in the nuclei of chondrocytic cells in culture and also in the nuclei of normal and osteoarthritic chondrocytes in vivo. The nuclear translocation of externally added recombinant MMP3 and six putative nuclear localization signals in MMP3 also were shown. Furthermore, we determined that heterochromatin protein gamma coordinately regulates CCN2/CTGF by interacting with MMP3. The involvement of this novel role of MMP3 in the development, tissue remodeling, and pathology of arthritic diseases through CCN2/CTGF regulation thus is suggested.

Connective tissue growth factor (CTGF/CCN2) is a member of the CCN family of matricellular proteins and also has been designated Hcs24, FISP12, IGFBP8, IGFBP-rP2, β1IG-M2, and ecogenin. The other CCN proteins include Cyr61/CCN1, NOV/CCN3, WISP1/CCN4, WISP2/CCN5, and WISP3/CCN6 (5, 26, 38, 39) as well, and they are structurally and functionally related glycoproteins involved in cell differentiation, proliferation, adhesion, migration, and the formation of the extracellular matrix. These matricellular functions of CCNs are involved in physiological processes such as wound healing, angiogenesis, morphogenesis, and embryogenesis as well as in pathological states including cancers, myocardial infarction, fibrotic disorders, rheumatism, and osteoarthritis (33, 49). Cartilage is a connective tissue that is constructed by chondrocytes embedded within an ECM predominantly composed of collagen and proteoglycans. ECM remodeling is achieved by degrading extracellular matrix proteins. MMPs, which comprise a large family of enzymes that cleave growth factors and their binding proteins, thereby activating or inhibiting specific signaling events (15). Of note, the expression and role of MMP3 have been investigated in the pathological status of articular cartilage, such as in osteoarthritis and rheumatism (1, 52).

Recent study has demonstrated the existence and functions of intracellular MMPs and tissue inhibitors of metalloproteinases (TIMPs). TIMP-1 accumulates in the cellular nuclei in association with the cell cycle (54). Alternative splicing and promoter usage generate an intracellular MMP11 isoform directly translated as an active MMP (31). MMP2 is found in the nuclei of cardiac myocytes and is capable of cleaving poly-(ADP-ribose) polymerase (PARP) in vitro (28). MMP3 also is detected in the nuclei of hepatocytes and is involved in apop-
tosis (47). MT1-MMP exhibits an intracellular cleavage function and causes chromosome instability, and it cleaves centrosomal pericentrin in human cells but not in murine cells (12).

The mechanisms of CCN2/CTGF induction/production have been well investigated (4, 26, 29); however, there have been few approaches to directly bind regulatory proteins of the CCN2/CTGF gene. Recently, we investigated cell type-specific mechanisms of CCN2/CTGF gene regulation and found a cis-acting element, transcription enhancer dominant in chondrocytes (TRENDIC), between positions −202 and −180 from the transcription start site of CCN2/CTGF (7), a region that previously had been predicted to contain an NF-1-like site (7, 13). In our previous study, the strong production of CCN2 from chondrocytic cells was estimated to be mediated by TRENDIC rather than by a juxtaposing Smad-binding element (SBE) (7, 23). In this present study, we subsequently cloned the cDNAs encoding TRENDIC-binding factors and unexpectedly found MMP3/stromelysin-1 to be one of them. We then investigated whether or not MMP3 is localized in the nuclei of chondrocytes in vitro and in vivo. Having found such localization, we also examined if the nuclear MMP3 could bind with enhancer sequences in the CCN2/CTGF promoter and activate CCN2/CTGF transcription, and we showed that it did so. Finally, we evaluated the properties of MMP3 as a transcription factor by analyzing nuclear MMP3-associated proteins (NuMAPs).

MATERIALS AND METHODS

HCS-2/8 cDNA phage library. The cDNA phage library was constructed by using a ZAP cDNA synthesis kit (Toyobo, Osaka, Japan) and a ZAP express cDNA phage library. HCS-2/8 cDNA phage library. The cDNA phage library was constructed by using a ZAP cDNA synthesis kit (Toyobo, Osaka, Japan) and a ZAP express cDNA phage library.

Cell culture. The following human-derived cells were used: human chondrosarcoma-derived chondrocytic HCS-2/8 cells (40, 51), MDA-MB-231 breast carcinoma cells, HeLa cells derived from human cervical cancer, and SaOS-2 osteosarcoma-derived cells. A COS7 monkey kidney-derived cell line also was used. Cells were cultured in Dulbecco’s modified Eagle’s minimum essential medium (DMEM) supplemented with 10% fetal bovine serum (FBS) in humidified air containing 5% CO2 at 37°C. For immunohistochemistry and internalization studies, glass chamber slides (4 or 8 well) were coated with 50 μg/ml of collagen (Cellmatrix type I-C; Nitta-Gelatin, Osaka, Japan) for 30 min before cells were seeded.

Antibodies. We used the following anti-MMP3 antibodies: anti-MMP3 N-terminal region (5025; Sigma, St. Louis, MO), anti-MMP3 hinge region (4802; Sigma), anti-MMP3 C-terminal domain (4927; Sigma), and anti-MMP3 catalytic (CAT) domain antibody (4190; Sigma). We also used anti-CCN2/CTGF (AF666; R&D), anti-histone H3 (06-599; Upstate), anti-Smad 2 (ST919; Sigma), anti-α-tubulin immunoglobulin G (IgG) (T9026; Sigma), anti-cathepsin D (C-20; Santa Cruz Biotech, Santa Cruz, CA), anti-lamin A/C monoclonal IgM (sc-7293; Santa Cruz), and anti-β-actin (AC-74; Sigma). We used the following antibodies to detect tags: anti-Flag M2 (Sigma), anti-Myc tag (Abcam, Cambridge, United Kingdom), and anti-glutathione S-transferase (anti-GST) (GE Healthcare) antibodies. An antidiogestin alkaline phosphatase-conjugated Fab (Roche, Basel, Switzerland) was used in electrophoresis mobility shift assays (EMSA). For Western blotting, we used horseradish peroxidase-conjugated secondary antibodies against anti-mouse IgG (Amersham) and anti-rabbit IgG (Dako, Copenhagen, Denmark). An anti-rabbit IgG rhodamine conjugate (Sigma) was used for immunostaining. We used these antibodies at the concentrations instructed by the manufacturer. The MMP3 enzyme-linked immunosorbent assay (ELISA) kit was purchased from Daiichi Pure Chemicals (Tokyo, Japan).

Preparation of subcellular fraction proteins. The nuclear and cytoplasmic proteins were prepared by using a Celllytic NuCLEAR extraction kit (Calbiochem, San Diego, CA) according to the manufacturer’s protocol. The subcellular fractions were prepared by using a Proteolipid extraction subcellular proteome extraction kit (Calbiochem, San Diego, CA) according to the manufacturer’s protocol. Total cell lysate was prepared by using a Celllytic M reagent (Sigma) according to the manufacturer’s protocols. The protease inhibitor cocktail (Sigma) was added at the appropriate stage.

SDS-PAGE and Western blot analysis. Extracted proteins were heated at 95°C for 5 min in sodium dodecyl sulfate (SDS) sample buffer in the presence of 5% 2-mercaptoethanol and separated by SDS-polyacrylamide gel electrophoresis (PAGE) in 12% polyacrylamide gel. Alternatively, proteins were heated at 70°C for 10 min in lithium dodecyl sulfate sample buffer containing 50 mM DTT and
were separated in a 10% Bis-Tris NuPAGE gel (Invitrogen) in morpholinopropanesulfonic acid running buffer containing an antioxidant. Semidy electroblotting was carried out using a polyvinylidene difluoride membrane (Hyb P; GE Healthcare). The membrane was blocked in Tris-buffered saline containing 0.05% Tween 20 (TBST) and 5% skim milk for 30 min at room temperature (RT). After being blocked, the membrane was incubated with the primary antibody overnight at 4°C and subsequently was incubated with the secondary antibody for 1 h at RT in the blocking solution. The blot was visualized by using an enhanced chemiluminescence (ECL) Western blotting analysis system (GE Healthcare) with chemiluminescence detection. The photogram was obtained by autoradiography or by using an ECL minicamera (GE Healthcare). The band signals obtained by Western blotting were quantified by using ImageJ, version 1.37 (Wayne Rasband, NIH).

Recombinant proteins, MMP3 activator, and inhibitors. A recombinant human proenzyme MMP3 (rhMMP3; R&D) was purchased and used for the molecular weight control and the internalization assay. For other experiments, we used our purified rhMMP3, prepared as described below. Organomercurial (4-aminophenyl)mercuric acetate (APMA) (Sigma) was dissolved in 50 mM NaOH. Active MMP3 was prepared by incubating proenzyme MMP3 (final concentration, 83.3 nM) with 1 mM of APMA in Tris buffer (50 mM Tris-HCl, pH 7.5, containing 5 mM CaCl2 and 0.05% Triton X-100) at 37°C for 4 h. All MMP inhibitors were purchased from Calbiochem and were dissolved and stored in dimethyl sulfoxide (DMSO). Before use, they were diluted 100-fold in a cell culture medium (1% DMSO) and then added to cultured cells (final concentration, 0.01% DMSO). Transforming growth factor β (TGF-β) (10 ng/ml) was purchased from Peprotech (Rocky Hill, NJ).

Stromelysin endopeptidase activity assay. The stromelysin endopeptidase activity assay was performed using the fluorescence resonance energy transfer peptide substrate Mca-RPKPV-Nval-WR-KDp-NH2 fluorogenic substrate II (R&D, Minneapolis, MN) (32). The substrate (10 μM) was mixed with enzyme or nuclear extract in a Tris buffer (50 mM Tris-HCl, pH 7.5, containing 5 mM CaCl2 and 0.05% Triton X-100) and incubated at 37°C. The relative fluorescence units (RFUs) of the reactant in a 96-well black plate were measured by using a Fluoroskan AscentFL (Labsystem, Helsinki, Finland).

Immunochemical immunocytochemistry. Cells were fixed in 4% (wt/vol) paraformaldehyde in phosphate-buffered saline (PBS) for 15 min, washed in TBS, and permeabilized with 0.2% Triton X-100 for 15 min. Incubation with primary and secondary antibody was performed in PBS containing 1.5% normal goat serum and 0.05% Triton X-100 for 1 h, respectively. After three washes with TBS, the mounting and DNA staining were performed by using a ProLong gold antifade reagents with 4',6-diamino-2-phenylindole (DAPI) (Molecular Probes, Invitrogen). Cells were observed by confocal laser microscopy as described below, TE2000-U microscope (Nikon) with a laser (excitation wavelengths of 488 and 568 nm) (Bio-Rad) through an Eclipse Nikon filter set (NH1). The cells were observed by confocal laser microscopy (Nikon) with a laser (excitation wavelengths of 488 and 568 nm) (Bio-Rad) through an Eclipse Nikon filter set (NH1).

Probe CCN2p160, which contains a CCN2 promoter DNA fragment between positions −292 and −137, was prepared from a plasmid template by PCR and labeled with digoxigenin. For EMSA performed with rhMMP3, MMP3 inhibitor (0.5 mM) was added to each lane. For supershift assays, an antibody (2 μg) was preincubated at 4°C for 40 min before the addition of the probes. For competitive EMSA, the recombinant protein and competitor were preincubated at 4°C for 5 min before the addition of the probe. Poly-L-lysine (0.5 or 1 μg/reaction) was used.

ChIP assay. The chromatin immunoprecipitation (ChIP) assay was carried out according to the manufacturer’s protocol (Upstate) with a slight modification. A hammer was used to disrupt HCS-28 cells that were cultured for 48 h with a medium change at 24 h. Formaldehyde (final concentration, 1%) was added to the medium, and the cells were incubated for 10 min at 37°C. The cells were washed twice and scraped in ice-cold PBS containing a protease inhibitor cocktail (Sigma) and then centrifuged for 4 min at 200 × g at 4°C. Subsequently, the cells were lysed in 200 μl of SDS lysis buffer (50 mM Tris-HCl, pH 8.1, containing 10 mM EDTA and 1% SDS) for 10 min on ice. DNA was sheared by three 10-s rounds of sonication on ice by using a Handy Sonic model UR-20P (Tomy Seiko, Tokyo, Japan) at 30% of maximum power. After centrifugation at 10,000 × g at 4°C for 10 min, the supernatant was diluted 10-fold in ChIP dilution buffer (16.7 mM Tris-HCl, pH 8, containing 167 mM NaCl, 1.2 mM EDTA, 0.01% SDS, 1% Triton X-100, and proteinase inhibitor cocktail), and 1% of it was retained as the input. The sample was precleared by mixing salmon sperm DNA (50 μg) and anti-rhMMP3 (50 μg) antibodies with 5% Protein A–agarose (Upstate) for 30 min at 4°C. After a brief centrifugation, the supernatant was rotated with or without the anti-MMP3 CAT domain antibody (1:500; Sigma) overnight at 4°C and was further incubated after adding 60 μl (3%) of the 50% salmon sperm DNA–50% protein A-agarose slurry at 4°C for 1 h. The protein A-agarose–antibody–antigen complex was centrifuged at 300 × g at 4°C for 1 min, and the pellet was washed once with low-salt wash buffer (20 mM Tris-HCl, pH 8.1, containing 150 mM NaCl, 2 mM EDTA, 0.1% SDS, and 1% Triton X-100), once with high-salt wash buffer (20 mM Tris-HCl, pH 8.1, containing 500 mM NaCl, 2 mM EDTA, 0.1% SDS, and 1% Triton X-100), once with LiCl wash buffer (10 mM Tris-HCl, pH 8.1, containing 0.25 M LiCl, 1% NP-40, 1% deoxycholate, and 1 mM EDTA), and twice with TE buffer (10 mM Tris-HCl, pH 8.1, 0.1 mM EDTA). The antigen complex was eluted by being mixed and rotated in 250 μl of freshly prepared elution buffer (1% SDS, 0.1 M NaHCO3). The elution step was repeated, and the eluates were combined. A 4% volume of 5 M NaCl was added to the eluate and to the input sample. The cross-linked chromatin complex was reversed by being heated at 65°C for 4 h, and DNA was purified using QIAquick spin columns (Qiagen, Hilden, Germany). PCR was carried out using Prime STAR HS DNA polymerase (Takara) according to the manufacturer’s protocol. Ten percent of the total purified DNA was used for the PCR in 50 μl of reaction mixture. The 204 bp of CCN2 enhancer fragment was amplified with primers between –292 and –88 with an annealing temperature of 63°C. The PCR products were separated on a 2% agarose gel in 0.5× TBE buffer, run at 100 V for 2 h, transferred to a nitrocellulose membrane, and visualized by using the developed film.
GTC ATC ATC TC-3’ and GAPDH NEO LCR (5'-GTC TTC TGG GTG GCA GTG AT-3') in 30 cycles of PCR under the following conditions: 98°C for 10 s, 65°C for 20 s, and 72°C for 20 s. The PCR products were analyzed by 2% agarose gel electrophoresis.

**Plasmid constructs.** The cDNA of MMP3 cloned in pBCK-CMV by Southwestern screening was subcloned into the pFlag-myc vector (Sigma) via a PCR-based method, and it was designated pFlag-MMP3-myc. A series of CGFCCN2 promoter-luciferase reporter constructs were described previously (7–9). The cDNA of MMP3 was subcloned from the pFlag-MMP3-myc vector to the pCold-TFI vector for expression in *E. coli*. Expression plasmids for MMP3 domains (full length [FL] and active forms, the prodomain, the combination of the CAT and Hinge domains [CAT + Hinge], and hemopexin-like repeat [PEX]) and MMP3 point mutants (H218R, H220R, and H216/228R) were constructed via a PCR-mediated mutagenesis method. The cDNA of GST was recombined from the pGEX-6p-3 vector (Amersham) to the pCold-TFI vector (Takara) using HindIII and SalI sites by a PCR-mediated method to constitute pCold-GST. The cDNAs of heterochromatin protein 1x (HP1x), HP1y, NF45, nuclear receptor corepressor 1 (NCOR1), and chromatin assembly factor 4 (CAF4) were predicted and synthesized by IGENE (Sapporo, Japan). The specific short interfering RNAs (siRNA) and gene knockdown studies.

For reporter gene assays, 500 ng of the reporter construct and 50 ng of the pRL-TK(int) vector (Promega, Madison, WI) were co-transfected with a total of 1 μg of plasmid DNA in the reporter/effector control ratio of 2:1 or 10:1. Medium was changed 24 h after the transfection. After being cultured for a further 24 h, cells were lysed in 200 μl of 1× passive lysis buffer (Promega, Madison, WI) with gentle rocking for 20 min and were collected. Luciferase assays were carried out by using a Dual Glo luciferase assay system (Promega), as described previously, on a smaller scale (17). Relative luciferase activities were calculated as the ratios of firefly luciferase activity to renilla luciferase activity.

DNA transcription and luciferase assay. Cationic liposome-mediated DNA transfection was carried out with a Fugene 6 transfection reagent according to the manufacturer’s optimized methodology (Roche). For reporter gene assays, cells were seeded in 12-well plates and cultured for 12 to 24 h. Subsequently, cells were transfected with 1 μg of plasmid DNA in the reporter/effector control ratio of 2:1 or 10:1. Medium was changed 24 h after the transfection. After being cultured for a further 24 h, cells were lysed in 200 μl of 1× passive lysis buffer (Promega, Madison, WI) with gentle rocking for 20 min and were collected. Luciferase assays were carried out by using a Dual Glo luciferase assay system (Promega), as described previously, on a smaller scale (17). Relative luciferase activities were calculated as the ratios of firefly luciferase activity to renilla luciferase activity.

siRNA and gene knockdown studies. The specific short interfering RNAs (siRNAs) were predicted and synthesized by iGENE (Sapporo, Japan). The siRNA targeting MMP3 (siMMP3-1 (5'-GAG AGC UUG UUA UAU UCA CCA CUA A-AU-3') and siMMP3-2 (5'-CUU UCU UUG CAU -3')) was designed and synthesized by Invitrogen (Invitrogen, Carlsbad, CA). The cDNA of MMP3 was cloned in pBCK-CMV by Southwest screening. For transient gene knockdown studies, which improves protein solubility. 5°C. The induction of the protein synthesis was carried out by cooling the E. coli at 15°C for 30 min and further shaking the sample for 24 h at 15°C after the addition of 0.5 mM IPTG. Cellular pellets were obtained by centrifugation, frozen at −80°C, melted, suspended in appropriate volumes of lysis buffer (50 mM Tris, pH 8.0, containing 500 mM NaCl, 1% Triton X-100, 1 μM pepstatin A, and 1 mM phenylmethylsulfonyl fluoride [PMSF]), disrupted by four 30-s sets of sonication, and centrifuged at 17,000 × g for 15 min at 4°C. To confirm the recombinant protein production in the supernatants, we carried out SDS–10% PAGE and Coomassie brilliant blue (CBB) staining using CBB R-250 (Sigma). Bovine serum albumin (BSA) was used as a concentration standard.

**GST pull-down assay.** The GST pull-down assay was carried out by using the soluble fractions including recombinant proMMP3, GST, and GST-fused proteins as described above. Twenty microliters of glutathione-Sepharose 4B beads (GE Healthcare) was washed once with 500 μl of the Tris buffer (50 mM Tris-HCl, pH 7.5, containing 150 mM NaCl, 1 mM EDTA, and 1% Triton X-100) with protease inhibitors (0.5 mM PMSF and 1 mM pepstatin A). The beads were mixed with the soluble fractions containing 3, 10, or 30 μg of GST-fused recombinant proteins in 500 μl of the Tris buffer, and the soluble fraction containing 10 μg of recombinant proMMP3 tagged with Flag, was added. The mixture was rotated for 2 h at 4°C, and the precipitant was washed four times with 500 μl of the Tris buffer. The precipitant was dissolved in 20 μl of 2× SDS sample buffer containing 8 M urea, boiled at 95°C, and separated by SDS–10% PAGE. Western blot analysis was performed using an anti-Flag M2 antibody (Sigma) or anti-GST antibody (GE Healthcare).

**Animals, tissue preparation, and immunohistochemistry.** The hind limbs of 8-week-old female BALB/c nu/nu mice (Clea, Tokyo, Japan) were fixed with...
10% neutral phosphate-buffered formalin. The experimental osteoarthritis was induced by injecting monoinoiodoacetic acid (MIA; Sigma) into the intraarticular spaces of 6-month-old Wistar rats, as described previously (36). These procedures were approved by the Animal Committee of the Okayama University Dental School. The animals were processed for histological analysis at 6 weeks after the injections. The specimens were decalcified in a 14% EDTA solution for 2 to 3 weeks and then embedded in paraffin, and the sections were prepared. After deparaffinization, the antigen in the section was activated by immersion in 10 mM target retrieval solution (Dako) in a microwave oven operating for 2 min. The sections were blocked in TBS with 10% FBS for 10 min and incubated with anti-rabbit IgG tetra-
methyl rhodamine isothiocyanate conjugate (1:100) in TBS containing 3% BSA at 4°C overnight. Afterwards, the sections were incubated with anti-rabbit IgG tetra-
the anti-MMP3 C-terminal antibody (1:100) in TBS containing 3% BSA at 4°C overnight. The sections were blocked in TBS with 10% FBS for 10 min and incubated with

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RESULTS

MMP3 in the cell nucleus. To identify TRENDIC-binding nuclear factors, we performed Southwestern screening by using a cDNA library of HCS-2/8 cells, since TRENDIC was first identified as an enhancer element dominant in these cells (7). Among the three genes cloned (Table 1), we further analyzed MMP3, because MMP3/stromelysin-1 is known to have an important role in cartilaginous tissues, although it had been recognized to act in the extracellular environment. In order to investigate if MMP3 was present in the cell nucleus, we prepared subcellular fractions of HCS-2/8 cells and analyzed the distribution of MMP3 by Western blotting. Surprisingly, MMP3 was detected in the nuclear extract of HCS-2/8 cells (Fig. 1A). The molecular weight of the major nuclear MMP3 was confirmed to be identical to that of rhMMP3 by Western blotting with antibody against the MMP3 CAT domain (Fig. 1A). Next, we further analyzed MMP3 in a variety of subcellu-
lar fractions by using an anti-MMP3 C-terminal antibody. Initially, the quality of subcellular fractions was verified. We detected histone H3 as well as the transcription factor Sox6 specifically in the nuclear fraction, α-tubulin in the cytosolic fraction, cathepsin D in cytosolic and membrane/organelle fractions, and lamin A/C in the cytoskeletal matrix fraction (Fig. 1B, left). The distinct localization of these marker pro-
tains verified the purity of the fractions. Among these fractions, FL and putative fragments of MMP3 were distinctly detected in the nuclear fraction (Fig. 1B, center). MMP3 also was detected immunocytochemically in the nuclei of HCS-2/8 cells. The signal of MMP3 overlapped with that of the nuclear stain DAPI (Fig. 1C). Finally, the existence of MMP3 inside the circle of lamin A/C, which is a component of the nuclear lamina, was confirmed by confocal laser scanning microscopy (Fig. 1D). We also detected a significant amount of secreted MMP3 in the cell culture supernatant of HCS-2/8 cells as evaluated by an ELISA (480 ng/ml) (Fig. 1B, right). Taking these findings together, we confirmed that MMP3 was actively synthesized in HCS-2/8 cells and was secreted or/and translocated into the nuclei.

DNA-binding ability of MMP3. We originally cloned MMP3 as a DNA-binding factor (Table 1). To examine whether MMP3 could bind to TRENDIC or not, we first performed EMSAs. The TRENDIC probe (pTRENDIC) was shifted in the gel with nuclear proteins of HCS-2/8 cells (Fig. 2A, lanes 1 and 2), as observed previously (7). The band shift was attenuated by the anti-MMP3 PEX repeat antibody (Fig. 2A, lane 5) but not by the addition of other anti-MMP3 antibodies (Fig. 2A, lanes 3 and 4). The control anti-rabbit IgG also did not interrupt the protein-TRENDIC interaction (data not shown). We also examined the disruptive effect of the PEX antibody on the interaction of nuclear proteins with BCE1/TbRE and Seq2 probes, which are other enhancers in the CCN2/CTGF promoter (7, 13). The band shift of pBCE1/TbRE or pSeq2 also was attenuated by the PEX antibody (Fig. 2B, lanes 3 and 6), indicating that MMP3 also is involved in the gene regulation mediated by these elements.

To confirm the specificity of the TRENDIC-MMP3 binding, a competitive EMSA was performed. The probe-protein inter-
action was inhibited by the addition of cold TRENDIC in a dose-dependent manner (10-, 100-, or 1,000-fold excess of cold TRENDIC) (Fig. 2C).

To further examine the interaction between the genomic CCN2 enhancer and the nuclear MMP3, we carried out a ChIP assay. In the chromatin immunoprecipitate obtained from HCS-2/8 cells by using anti-MMP3 antibody, the genomic CCN2 enhancer region between positions −292 and −88, which contains enhancers such as TRENDIC, BCE1/TbRE, and Seq2, was detected by the ChIP assay (Fig. 2E). No such signal was detected without the anti-MMP3 antibody (Fig. 2E). The open reading frame (ORF) fragment of GAPDH was not detected by the ChIP, either with or without the anti-MMP3 antibody (Fig. 2E). Both the genomic CCN2/CTGF enhancer region and the ORF fragment of GAPDH were detected in the cross-linked chromatin sample before immunoprecipitation (Fig. 2E). These results indicate that MMP3 exists in the protein complex, binding to the CCN2 enhancer region. Taken together, these results indicate that MMP3 not only interacted in vitro with dsDNA having the enhancer sequences but also interacted in situ with the genomic CCN2/CTGF enhancer region in the HCS-2/8 cells.

Effects of intracellular MMP3 on CCN2 promoter activity. To clarify if MMP3 could modulate the CCN2 promoter activity, we cotransfected several cell lines with an MMP3 expres-
sion vector (p3xFlagMMP3-myc; schemed in Fig. 3A) and CCN2 promoter reporter constructs and then quantified the promoter activity by conducting luciferase assays. MMP3 expression by transient transfection was confirmed by immuno-
 blotting (Fig. 3B). The intracellular overexpression of MMP3 activated the CCN2/CTGF promoter (Fig. 3C, D). TGF-β and intracellular MMP3 synergistically activated the CCN2/CTGF promoter (Fig. 3C). The CCN2 promoter activity with MMP3 overexpression was 2.2-fold higher in HCS-2/8 cells, 1.8-fold higher in MDA231 cells, and 2.2-fold higher in COS7 cells than in their controls (Fig. 3D). In contrast, the effect of overexpressed MMP3 was not significant in HeLa or SaOS-2 cells (Fig. 3D). It should be noted that the relative CCN2 promoter activity in HCS-2/8 and MDA231 cells was over 30-fold higher than in SaOS-2 and HeLa cells (data not shown). To further investigate this mechanism, we utilized mutants of the CCN2 promoter (Fig. 3E). Mutants pDS4 and pDS5, lacking the element between −202 and −88, lost the response to overexpressed MMP3 in HCS-2/8 cells (Fig. 3F). The response
Effects of MMP3 knockdown on CCN2 gene expression. To investigate if the endogenous MMP3 was involved in the abundant production of CCN2 from HCS-2/8 cells (7, 9), we knocked down MMP3 mRNA by RNA interference technology and evaluated the effect of the siRNA on CCN2 mRNA and protein expression and promoter activity. Transient transfection of siMMP3-1369 (20 nM) decreased MMP3 mRNA expression to 14% of the control dsRNA (siNC) level (Fig. 4A, left), indicating that siMMP3-1369 successfully degraded MMP3 mRNA. Under the condition of MMP3 being knocked down, the CCN2 mRNA level also was decreased to 70% of the control level (Fig. 4A, right).

Next, to evaluate the effect of the MMP3 siRNA (siMMP3) on MMP3 and CCN2 protein production, 0, 20, 50, or 100 nM
of siMMP3-1302 or control dsRNA (siNC) was transfected into HCS-2/8 cells, and MMP3, CCN2, and GAPDH protein production in the lysate was evaluated by Western blotting. As a result, the levels of FL MMP3 (Fig. 4B, upper left) and MMP3 fragments (Fig. 4B, upper right) decreased in a dose-dependent manner, while the levels of these MMP3 molecules were not decreased by control dsRNA transfection. Under the MMP3 knockdown conditions, the CCN2 protein level also was decreased (Fig. 4B, lower left) compared to that with control dsRNA. The GAPDH protein production level was not affected by the knockdown of MMP3 (Fig. 4B, lower right). These results indicate that MMP3 regulates CCN2 gene expression in HCS-2/8 cells. In agreement with this finding, the CCN2 promoter activity was attenuated down to 40% of the control level in HCS-2/8 cells (Fig. 4C, left) and 60% of the control level in MDA231 cells (Fig. 4C, right) by the siMMP3-1369 transfection.

To further confirm the regulation of CCN2 gene expression by MMP3 in HCS-2/8 cells, a retroviral vector that expressed shMMP3 or scrambled shRNA was constructed and infected into HCS-2/8 cells, and MMP3 and CCN2 expression was evaluated 2 weeks after the infection. The shMMP3 retroviral vector successfully knocked down the level of expression of MMP3 mRNA to 35% of that of the control (Fig. 4D, left). In the retrovirus-mediated long-term MMP3 knockdown condition, CCN2 mRNA expression was as low as 28% of the control level (Fig. 4D, right).

If MMP3 is a critical and general regulator of CCN2, the expression of CCN2 is anticipated to be significantly lower in other cells without MMP3 expression. To confirm this, we comparatively analyzed the expression of CCN2 and MMP3 in HCS-2/8 and HeLa cells. In contrast to its expression in HCS-2/8 cells, MMP3 mRNA expression was deficient in HeLa cells (Fig. 4E). As we expected, the CCN2 expression level in HeLa cells was strikingly low (12%) compared to that in HCS-2/8 cells. These results indicate that abundant MMP3 regulates CCN2 gene expression in HCS-2/8 cells, while CCN2 expression in HeLa was low because of the deficiency of MMP3, a
crucial factor in CCN2 induction. Additionally, both MMP3 and CCN2 expression levels in SaOS2 osteoblastic cells were quite low (data not shown). Taken together, these data demonstrated that cellular MMP3 regulates CCN2 gene expression at the transcription level.

MMP3 domains required for CCN2/CTGF promoter activation. In order to determine the involvement of individual domains of MMP3 in the CCN2/CTGF trans-activation function, several assemblages of MMP3 domains were expressed in COS7 and HCS-2/8 cells, and the CCN2/CTGF promoter reporter assay was carried out. The expression of the MMP3 domain deletion mutants in COS7 cells was confirmed by Western blotting (Fig. 5C). The FL and active forms of MMP3 trans-activated the CCN2/CTGF promoter, while the prodo-
main alone did not (Fig. 5A). Interestingly, the PEX domain of MMP3 lacking a CAT domain trans activated the CCN2/CTGF promoter in the cells, and the trans activation ability of the MMP3 PEX domain was comparable to that of the FL and active form (Fig. 5A). Moreover, CAT+Hinge also trans activated the CCN2/CTGF promoter, while Hinge alone had no effect for trans activation (Fig. 5A). Similar results of MMP3 domain functions were obtained in HCS-2/8 cells (Fig. 5A and B). These results clarified that not only the FL and active forms of MMP3 but also a few MMP3 fragments, such as PEX and CAT+Hinge domains, could trans activate the CCN2/CTGF promoter.

A proteolytic activity of MMPs has been known to be zinc dependent. In the part of the CAT domain forming the active site, a Zn$^{2+}$ ion is coordinated by a chelate bond with three histidine residues found in the conserved sequence HEXXHXXGXXH and is called a zinc-binding motif. The change of any one of the histidines to arginine was reported to cause the loss of the proteolytic activity of MMPs (27). To clarify if the trans-activation ability of MMP3 for CCN2 expression is dependent on proteolytic activity or not, we constructed plasmids expressing catalytically dead mutants of MMP3. As a result, the alteration of a histidine residue at position 218 (H218) to arginine (H218R mutant) resulted in a decrease of the trans-activation ability of MMP3 for the CCN2 promoter (Fig. 5D). The H228R mutant and the double mutant (H218R/H228R) yielded similar results (Fig. 5D). Both the CAT domain and the PEX domain lacking the CAT domain hold the trans-activation ability (Fig. 5A, B). Taking these findings together, the trans-activation ability of MMP3 for the CCN2 promoter is partly dependent on the catalytic activity, and the PEX domain greatly contributes to the activation of the CCN2 promoter.

Uptake and nuclear translocation of extracellular MMP3.
Next, to clarify the subcellular dynamics of MMP3, we prepared Cy3-labeled rhMMP3 (Cy3-MMP3), added it to the medium of HCS-2/8 cells in culture, and observed its behavior with a confocal laser scanning microscope. The Cy3-MMP3 signals emanated from the cellular membrane or cytoplasm between 5 and 60 min after the addition of Cy3-MMP3 to the cell culture (Fig. 6A, images b to f, i, and j). Of note, Cy3-
FIG. 4. Effects of MMP3 siRNA on CCN2 gene expression. (A) Knockdown of MMP3 also downregulated CCN2 gene expression. mRNA expression of MMP3 or CCN2 was quantified after the transfection of HCS-2/8 cells with siMMP3 or negative control dsRNA (siNC). siMMP3 knocked down the MMP3 mRNA to 15% of the level of the control while decreasing the amount of CCN2 mRNA to 70% of the control. (B) The knockdown of MMP3 also decreased the amount of cell-associated CCN2/CTGF protein. Negative control RNA (siNC) or siMMP3 was transfected at a final concentration of 20, 50, or 100 nM into HCS-2/8 cells. MMP3, CCN2, or GAPDH protein in the cell lysate at 48 h after the transfection was analyzed by Western blotting. R indicates the transfection-reagent-only control. Levels of FL (left upper panel, short exposure) or fragments (right upper panel, long exposure) of MMP3 were decreased by the knockdown. Under the MMP3 knocked-down condition, CCN2/CTGF protein also was decreased (left lower panel), indicating that MMP3 regulates CCN2. (C) CCN2 promoter activities under the condition of the MMP3 knockdown, as quantified by luciferase assays. siMMP3 also downregulated the CCN2 promoter activities in HCS-2/8 cells (to 40% of the control level) and also in MDA231 cells (to 60% of the control). (D) Retrovirus-mediated MMP3 knockdown decreased CCN2 mRNA expression. HCS-2/8 cells were infected by a retroviral vector that expressed shMMP3 or scrambled shRNA (Scr.), and then the mRNA expression levels of MMP3 and CCN2 at 2 weeks after the infection were analyzed by real-time PCR. MMP3 mRNA expression was successfully knocked down to 35% of the control level (Scr.). The CCN2 mRNA level under the MMP3 knockdown condition was 28% of the control level (Scr.). (E) Relative MMP3 and CCN2 expression levels in HCS-2/8 and HeLa cells. MMP3 mRNA expression was deficient in HeLa cells (left), while CCN2 mRNA expression in HeLa cells was 13% of that in the HCS-2/8 cells. These results indicate that MMP3 is a major regulator for CCN2 gene expression. *, P < 0.05 (n = 3 or 4); #, below the detection limit.
MMP3 signals also were observed in the cell nuclei between 15 and 30 min after the addition (Fig. 6, images f to i). The Cy3-MMP3 signal was barely detected at 60 min after the addition, indicating the possibility of degradation or resecretion. The Cy3-MMP3 localization in the Cy3-MMP3-positive cell population was quantitatively analyzed (Fig. 6B). The Cy3-MMP3 was taken up into the cells within 5 min after the addition, showing dominant localization in the cytoplasm, which was followed by the translocation into the nucleus between 15 and 60 min after the addition. Additionally, to demonstrate that MMP3-Cy3 was certainly in the nucleus, a sequential horizontal view of the MMP3-Cy3 signal and DNA in
FIG. 6. Extracellular MMP3 was internalized into the cells and translocated into the nucleus. (A) Cy3-MMP3 was added to the culture medium of HCS-2/8 cells. The cells were fixed at 0 (a), 5 (b to d), 15 (e to g), 30 (h, i), or 60 min (j) after the addition of MMP3 and then were observed under a confocal laser scanning microscope. Red signals indicate Cy3-MMP3, and green signals indicate DNA. Cy3-MMP3 was observed on the cell surface or in the cytoplasm between 5 and 60 min after the addition (b to f, i, and j). Cy3-MMP3 was observed in the nuclei at 15 and 30 min (f to i). Bars, 5 μm.

(B) Quantitative analysis of the Cy3-MMP3-positive cells in relation to subcellular localization. N, nucleus; C, cytoplasm; N/C, both. n = 4 to 7.

(C) Sequential view of the cells by using confocal laser scanning microscopy revealed that MMP3-Cy3 (red) is inside the nucleus filled with DNA (green).

(D) Effect of LMB on the activation of the CCN2/CTGF promoter by intracellular MMP3 expression. The addition of LMB (20 ng/ml) enhanced both the basal level and the trans-activation effect of MMP3 for the CCN2/CTGF promoter. * P < 0.05 as indicated by brackets (n = 4). vec., vector.
A cell was presented (Fig. 6C). These observations clarified that extracellular MMP3 is taken up into the HCS-2/8 cells and subsequently translocated into the nucleus.

Effect of LMB on the induction of CCN2/CTGF by MMP3. In order to demonstrate that nuclear translocated MMP3 transactivates the CCN2/CTGF promoter, leptomycin B (LMB) was employed. LMB is an antibiotic with membrane permeability, and it has been known to inhibit the nuclear export function of CRM1/exportin 1 by directly binding to its cysteine residue. We hypothesized that the nuclear export of MMP3 could be CRM1 dependent, and thus LMB could cause the nuclear accumulation of MMP3 and the enhancement of the activation of the CCN2/CTGF promoter by nuclear MMP3. Exactly as we predicted, LMB enhanced the trans-activation effect of MMP3 on the CCN2/CTGF promoter (Fig. 6D). The increase in the CCN2 promoter activity without MMP3 overexpression may represent the effect of LMB on endogenous MMP3. These results represent that nuclear-cytoplasmic trafficking is crucial in the trans activation of the CCN2/CTGF promoter by MMP3.

Domain structure and nuclear localization signals of MMP3. In addition to the classical domain structure of MMP3, we found and predicted several novel domains and signal sequences in MMP3 (Fig. 7A). As an anti-MMP3 PEX antibody blocked protein-DNA interactions, a DNA-binding domain can be included in the PEX domain. The hinge region has been known to be just a linker; however, it contains proline-rich sequences found in some transcription factors. The PEX domain was revealed to be a DNA-binding domain that forms a propeller-like structure. Putative NLSs in MMP3 are mapped beneath the structure. The putative NES containing four leucines also was found. (B) List of NLSs in MMP3 and schematically depicted EGFP-NLS fusion proteins. The arginine (R)- and lysine (K)-rich sequences were picked up from the FL amino acid sequence of MMP3 and designated NLS0 to NLS5. The cDNAs encoding these NLSs were cloned into the pEGFP-C1 vector. Arginine and lysine residues are indicated by boldface. The positions of the NLSs in proMMP3 and the domain in which the NLSs were found are shown. PEX, hemopexin-like repeat. NLS1, NLS2, and NLS3 in MMP3 also were predicted by using PSORTII pattern 4 (pat4) or pat7. (C) Subcellular localization of the GFP-NLSs. The subcellular localization of EGFP-NLSs and the control (Cont.) was examined in COS7 cells by confocal laser microscopy. EGFP-NLS1, EGFP-NLS4, and EGFP-NLS5 were observed in the nuclei as well as in the cytoplasm. Similar results were obtained for EGFP-NLS0, EGFP-NLS2, and EGFP-NLS3 (data not shown). EGFP-MCS (control) was observed only in the cytoplasm. Scale bars, 5 μm. The photogram is representative of 10 independent cells with similar results.
terminus of MMP3 contains proline at the fifth position. It can work as a helix breaker, causing the partition sorting of MMP3 to the endoplasmic reticulum (ER)-Golgi secretion pathway as well as to the cytoplasm (44). A leucine-rich sequence in the PEX domain can work as a nuclear export signal (NES).

Classically, the nuclear import of proteins is mediated by importins \( \alpha \) and \( \beta \), which recognize basic amino acid clusters, designated NLSs, on the protein to be imported (3, 34). In order to assess if MMP3 entered the nucleus via this classical pathway, we sought putative NLSs in MMP3. Six lysine- and arginine-rich sequences were found in the amino acid sequence of MMP3, and we designated them NLS0 to NLS5 (Fig. 7A, B). The putative NLSs were scattered among all of the domains of MMP3. Since MMP3 can be (auto)cleaved, the
scattered NLSs can function individually in the resultant fragments. In order to investigate the function of individual NLSs, we constructed the EGFP-NLS expression plasmids for all six putative NLSs in MMP3, overexpressed them in COS7 cells, and observed their subcellular localizations. The native EGFP (control) was observed only in the cytoplasm and not in the nucleus (Fig. 7C). Surprisingly, all EGFP-NLSs were observed in the nuclei as well as in the cytoplasm, as shown in Fig. 7C (data not shown for NLS0, NLS2, and NLS3). These results indicate the possible involvement of the NLS-mediated classical pathway in the nuclear translocation of MMP3 even for the FL and the fragments of MMP3.

**Identification of NuMAPs.** If MMP3 plays a certain role in the nuclei, it should be associated with other proteins to exhibit the downstream phenomenon, whether it cleaves them or not. To clarify this point, we applied the nuclear extract of HCS-2/8 cells to anti-MMP3 antibody columns and then eluted the bound proteins and analyzed them by using an LC-MS/MS system (Fig. 8A). Identified proteins included heterochromatin proteins, transcription activators/repressors, RNA polymerase II, nucleosome/chromatin assembly protein, and others (Table 2 and data not shown). Among them, we confirmed the mRNA expression of HP1α, HP1γ, NF45/interleukin enhancer binding factor 2 (ILF2), NCoR1, and CAFlp48/RBBP4/RbAp48 in the cells; thereafter, we prepared their GST-fused recombinant proteins and recombinant MMP3 (Fig. 8B, C). The bindings of HP1γ, NF45, NCoR1, and CAFlp48 to MMP3 were confirmed from the results of a GST pull-down assay (Fig. 8D); thus, these proteins were designated NuMAPs. These data suggest that MMP3 associated with these NuMAPs in the nucleus in exerting its nuclear function.

To clarify the function of these NuMAPs and the collaborative function with intracellular MMP3 for CCN2/CTGF transcription, mammalian NuMAP expression plasmids were constructed (Fig. 8E) and coexpressed in HCS-2/8 cells for a reporter gene assay. HP1γ recently was found to exhibit multiple functions beyond heterochromatin formation (19). In our study, the overexpression of HP1γ activated the CCN2/CTGF promoter (to a level 2.5-fold above that of the control), while the expression of other NuMAPs (NF45, RBBP4, and DDOST) caused no substantial change compared to that of the controls [pcDNA3.1(−) and pcDNA3.1(−)3HA] (Fig. 8F). In addition, the coexpression of HP1γ and MMP3 synergistically activated the CCN2/CTGF promoter (Fig. 8G). These results suggested that HP1γ, a member of the NuMAPs, regulates CCN2/CTGF transcription in cooperation with nuclear MMP3.

![FIG. 8. Identification of NuMAPs. (A) The methodology to identify the NuMAPs. The nuclear proteins extracted from HCS-2/8 cells were immunoprecipitated by using the anti-MMP3 affinity beads for the MMP3 CAT domain (Cat.), Hinge region, or PEX antibody or IgG (control). The eluted proteins were digested with trypsin at the C-terminal peptide bonds of lysine and arginine. Samples were applied to the nanoflow high-performance liquid chromatography (HPLC) chip MS/MS, and the data were analyzed to identify the proteins. The NuMAP candidates are shown in Table 2. (B) Structures of the recombinant MMP3 and several GST-fused NuMAP candidates. Trigger factor (TF) was added to all of the proteins to increase their solubility. (C) CBB staining of the recombinant proteins. The soluble fractions (1 or 5 µl) or the BSA control was electrophoresed in 10% polyacrylamide gels and stained. Arrows indicate the recombinant proteins. (D) MMP3-NuMAP interactions. Flag-tagged MMP3 (10 µg) was mixed with the GST-fused NuMAP candidates (3, 10, or 30 µg) and then pulled down with glutathione-Sepharose beads. The pulled down proteins were analyzed by immunoblotting using an anti-Flag or anti-GST antibody. The direct interaction of MMP3 with HP1γ, NF45, NCoR1, and RBBP4 was confirmed. (E) Expression of HA-tagged HP1γ, NF45, RBBP4, and DDOST in COS7 cells. Schemes of the expression constructs are illustrated on the top. Their expressions were confirmed by examining the lysates by Western blotting (lower panels). (F) Trans activation of the CCN2/CTGF promoter by HP1γ among NuMAPs, revealed by using reporter assays. *, P < 0.05 as indicated by brackets (n = 4). (G) Cooperative trans activation of the CCN2/CTGF promoter by MMP3 and HP1γ. HCS-2/8 cells were cotransfected with pB2CCN2promoter-luc (400 ng), pHRL-TK(int−) (100 ng), p3Flagmyc or pMMP3 (1 µg), and p3xHA or pHPlγ (1 µg) in each well of a 12-well plate for the luciferase assay. *, P < 0.05 (n = 4); #, P < 0.05 as indicated by brackets (n = 4). (H to K) Colocalization of HA3 tag was faintly observed to be diffused. Scale bars, 5 µm. IP, immunoprecipitation; vec., vector.](http://mcb.asm.org/FIG8.jpg)
MMP3 can be (auto)cleaved into the CAT domain and the PEX domain in or out of the cells, and the PEX domain and CAT+Hinge domain possess a trans-activation ability for the CCN2/CTGF promoter at a level comparable to that of FL MMP3. In addition, both contain putative NLSs. To clarify the localization of NuMAPs (HP1γ, NF45, and RBBP4) with MMP3s (FL, PEX, and CAT domains), these proteins or domains were coexpressed in COS7 cells, and then the colocalization was investigated by using confocal microscopy. Firstly, HP1γ, NF45, and RBBP4 were observed mainly to be localized only in the nucleus (Fig. 8H to J), while the HA3 tag was observed to be faintly diffused (Fig. 8K). The FL, PEX, or CAT domain of MMP3 was observed to be localized in the nucleus as well as in the cytoplasm. Neither NuMAPs (HP1γ, NF45, and RBBP4) nor MMP3s (FL, PEX, and CAT domains) were observed in the nucleolus (Fig. 8H to K). These findings indicated that HP1γ, NF45, and RBBP4 can interact with MMP3 in the nucleus in vivo, as demonstrated by in vitro GST pull-down assays (Fig. 8D). Taking these results together, among all NuMAPs tested, HP1γ can trans activate the CCN2/CTGF promoter by interacting with MMP3 in the nucleus.

Subcellular localization of MMP3 in normal and osteoarthritic cartilaginous tissues. To investigate the relationship between subcellular MMP3 localization and the pathophysiology of articular cartilage in animals, we immunohistochemically examined MMP3 in normal and osteoarthritic articular cartilages with a confocal laser microscope. MMP3 was immunopositive in the normal articular cartilage of 2-month-old mice (Fig. 9A, row a), while control IgG brought no significant signal. The colocalization of the MMP3 signals with DNA was very evident at high-power magnification (Fig. 9A, row b). In a rat osteoarthritic cartilage, the articular and semilunar chondrocytes were positively stained by the MMP3 antibody (Fig. 9B). We also observed the colocalization of the MMP3 staining with DNA in articular chondrocytes at high-power magnification (Fig. 9B, rows e and f). MMP3 in the cytoplasm also was observed in some osteoarthritic articular chondrocytes (Fig. 9B, rows e and f; yellow in merged view). It should be noted that the fibrochondrogenic cells, in which CCN2 would be expressed and involved in the tissue remodeling, showed MMP3 existed alongside DNA (Fig. 9B, row g; yellow in merged view). The data that MMP3 is localized in the nuclei of normal developing and osteoarthritic articular chondrocytes in animals suggest a role of the nuclear MMP3 in the development and regeneration of articular cartilage.

Effects of MMP inhibitors on CCN2/CTGF promoter activity and on nuclear stromelysin-like endopeptidase activity. To investigate if the proteinase activity of MMP3 was associated with the observed transcriptional activation of CCN2, we added MMP inhibitors to the culture medium of HCS-2/8 cells and subsequently quantified the CCN2 promoter activities. A specific inhibitor of human MMP3 (MMP3 inhibitor II) suppressed the CCN2 promoter activity in a dose-dependent manner (Fig. 10A). In contrast, a specific inhibitor of MMP2 and MMP9 did not suppress the CCN2 promoter activity (Fig. 10A), nor did GM6001, a well-known broad-spectrum hydroxamic acid inhibitor of MMPs (Fig. 10A). These results indicate that the structural hindrance caused by the specific inhibitor blocked the interaction between MMP3 and its binding target, thus strongly suggesting the involvement of the CAT domain in the transcriptional activation events.

To determine further if the nuclei of HCS-2/8 cells exerted stromelysin endopeptidase activity, we tested the ability of a nuclear extract of HCS-2/8 cells to cleave a fluorogenic sub-
strate of stromelysin (32). Initially, the stromelysin activity of MMP3 (8.3 nM) activated by 1 mM APMA was successfully detected by measuring the number of RFUs emitted by the cleavage of the peptide substrate (Fig. 10B). This stromelysin endopeptidase activity of active MMP3 was blocked by MMP3 inhibitor II or GM6001 but not by the MMP2/MMP9 inhibitor (Fig. 10B). The endopeptidase activity of proenzyme MMP3 also was tested. Proenzyme MMP3 (1.7 nM) cleaved the fluoro-

FIG. 10. Characterization of stromelysin-like activity in HCS-2/8 nuclei. (A) Effects of MMP inhibitors on CCN2 promoter activity in HCS-2/8 cells. MMP2/MMP9 inhibitor I (M2/9i), MMP3 inhibitor II (M3i), or GM6001 was added to the culture medium of HCS-2/8 cells at a final concentration of 0, 0.02, 0.2, 2, or 20 μM. The cells were transfected with the CCN2/CTGF promoter reporter 8 h after the addition, and then reporter gene expression was evaluated 48 h after the transfection. Only MMP3 inhibitor II at a concentration of 20 μM suppressed the CCN2/CTGF promoter activity (to 66% of the control level); however, neither GM6001 nor the gelatinase inhibitor affected the promoter activity. These results indicate that endogenous MMP3 regulated CCN2/CTGF gene expression via the CCN2/CTGF promoter. *, P < 0.05 (n = 4). (B) Establishment of the stromelysin activity assay in vitro. A stromelysin-specific peptide substrate, Mca-RPKPVE-Nval-WRK(Dnp)-NH₂ fluorogenic substrate, was tested for cleavage by rhMMP3 as well as by a nuclear extract of HCS-2/8 cells, and the endopeptidase activity was quantified by measuring the RFUs. Active MMP3 (8.3 nM) fully cleaved the peptide in 2 h, and the endopeptidase activity was perfectly blocked by MMP3 inhibitor II (M3i) or GM6001 but not by the MMP2/MMP9 inhibitor (M2/9i). *, P < 0.05 (n = 4). (C) Proenzyme MMP3 (1.7 nM; 100 ng) cleaved the substrate as well, suggesting autoactivation along the assay. (D) Nuclear extract (NE) of HCS-2/8 cells (100 ng) also cleaved the substrate. (E) The endopeptidase activity of the nuclear extract was perfectly blocked by 1 mM APMA. w/o, without.
orogenic substrate relatively slowly (Fig. 10C), suggesting that proenzyme MMP3 was becoming activated in the reaction buffer at 37°C. Instead of the recombinant MMP3, we next tested the stromelysin endopeptidase activity of the nuclear extract prepared from HCS-2/8 cells. This extract also cleaved the peptide substrate in a dose-dependent manner; 6 μg of the nuclear extract exerted enough activity to cleave the peptide to the same extent as 100 ng (1.7 nM) of proenzyme MMP3 (Fig. 10D). These findings support our idea that active forms of FL or/and smaller forms of MMP3 exist in the nuclei of chondrocytes. Unexpectedly, this stromelysin activity of HCS-2/8 nuclear extracts was perfectly blocked by 1 mM APMA (Fig. 10E) but not by any inhibitor tested, i.e., GM6001, MMP3 inhibitor II, and the MMP2/MMP9 inhibitor (data not shown). Previously, PEX domain-deleted mini-stromelysin mutants were revealed to show an altered response to APMA, in that precursor processing was inhibited rather than accelerated (11). In our data, FL and several shorter MMP3s were detected by using anti-MMP3 C-terminal domain antibody (Fig. 1B). Thus, we speculate that the FL or shorter MMP3s in the nucleus possess stromelysin activity, which was blocked by APMA. In contrast to the results of the endopeptidase activity test using nuclear extracts in vitro, in a reporter gene assay, MMP3 inhibitor II inhibited the CCN2 promoter activity, whereas GM6001 did not (Fig. 10E). Therefore, GM6001 may not be able to efficiently enter the cells, and MMP3 inhibitor II may have blocked the MMP3 action in any of the ensuing steps, e.g., the release from ECM, endocytosis, or nuclear import, rather than in the nucleus. These findings suggest that the possible modification of MMP3 or some interaction partner of MMP3 in the nucleus protects MMP3 from MMP inhibitors to sustain stromelysin activity.

Specificity of MMP3-TRENDIC interaction and its consensus sequence. The DNA-binding ability of MMP3 was already demonstrated in Fig. 2, in which we showed that TRENDIC and the other cis elements were able to bind MMP3 in the nuclear extract. To further investigate the sequence specificity of MMP3-targeted DNA, we initially performed in silico analyses and, subsequently, a competitive EMSA (Fig. 11). TRENDIC sequences from different species were aligned and compared (GenBank accession numbers were shown in a previous study) (Fig. 11A) (7). Obviously, the TRENDIC sequence is highly conserved among vertebrates (Fig. 11A). The TRENDICs were 92% conserved between human and bovine species. The TRENDICs also were highly conserved in rodents, particularly mice and rats (96%). The amphibian TRENDIC sequence was relatively different from those of other vertebrates (77% conserved between humans and frogs). Nevertheless, a sequence

![FIG. 11. Sequence specificity of MMP3-TRENDIC binding. (A) Sequence conservation of TRENDIC among species. Human, bovine, murine, rat, and frog TRENDIC sequences are aligned. The nucleotides conserved between human TRENDIC and those of other species are shown in boldface. The TRENDIC core sequence predicted from the TRENDIC sequences of the species is shown at the top. (B) Alignment of the cis elements in the human CCN2 promoter. The nucleotides identical to those in TRENDIC are shown in boldface. The TRENDIC core sequence predicted from the TRENDIC sequences of the species is shown at the top. (C) Specific binding of MMP3 to the CCN2 enhancer sequence in the presence of an excess of nonspecific competitor. Probe CCN2p160, which contains the three enhancers shown in panel B, was shifted with recombinant MMP3 in a dose-dependent manner in the presence of 0.5 μg of poly(dA-dT). (D) The MMP3-enhancer complex was supershifted with anti-PEX antibody but not with anti-Hinge or control IgG. Poly(dA-dT) (1 μg/reaction) was used. f., free probe; s., shifted probe; s.s., supershifted probe. (E) Sense sequences of modified TRENDIC competitors. The corresponding dsDNAs were used for the competitive EMSA shown in panel F. (F) The MMP3-p160 interaction was interfered with by TRENDIC but not by its modified sequences. The competitors (TRENDIC, SeqA, and SeqB; 1-, 10-, 100-fold excesses of probe) were preincubated with rhMMP3 before the addition of the probe p160 with poly(dI-dC) (0.1 μg/reaction). TRENDIC competed with p160 for binding with rhMMP3 (lanes 1 to 3), whereas the SeqA dsDNA failed to compete with the binding reaction (lanes 4 to 6). SeqB was modestly competitive with the probe (lanes 7 to 9).]
promoter activity (Fig. 10). All of these data together support our contention that MMP3 is translocated to the cell nucleus and, once there, trans activates the CCN2 gene.

In this study, most of the data were obtained by using HCS-2/8 chondrocytic cells, a cell line that has been a useful model in chondrocyte biology. The cells have been used as a model of normal and osteoarthritic chondrocytes in a number of studies (30, 40, 51). Of note, CCN2 originally was cloned as a hypertrrophic chondrocyte-specific gene product 24 (Hcs24) from HCS-2/8 cells and was demonstrated to function as an endochondral ossification-promoting factor (50). Indeed, HCS-2/8 cells produced abundant amounts of CCN2 compared to the amount produced by other cell types (7, 9). The HCS-2/8 cell line was established from a chondrosarcoma, but the expression of CCN2 in cartilaginous tumors is negatively related with the level of their malignancy (45), indicating that HCS-2/8 cells can be a model of a normal phenotype of chondrocytes. This cell line also has been used for arthritis studies, because HCS-2/8 produces MMP3 and matrix degradation-related molecules in response to tumor necrosis factor alpha (16, 17, 41). In addition, MMP3 immunopositive cells were specifically expressed in normal articular cartilage of a 14-year-old human, while MMP9, MMP13, and ADAMTS4 were not detected (H. I. Roach, personal communication). We observed nuclear MMP3 in both normal and osteoarthritic cartilages (Fig. 8). In addition, the strong expression of CCN2/CTGF in chondrocytes in osteoarthritic tissues already has been reported (36). Thus, nuclear MMP3 may function when chondrocytes are in a physiological or pathological status.

The trans activation of the CCN2/CTGF promoter by the nuclearly translocated MMP3 is a major discovery in this study. One piece of evidence for this finding is that the knockdown of MMP3 resulted in a decrease in CCN2/CTGF expression (Fig. 4). Moreover, a major CCN2/CTGF-producing cell line, HCS-2/8, was found to produce MMP3, while MMP3 expression was undetectable in HeLa cells that produce small amounts of CCN2/CTGF (Fig. 4) (7–9). MMP3 is known as a secretory proteinase that degrades ECM proteins such as proteoglycans and collagens, while MMP3 also cleaves and activates other MMPs, such as gelatinases, and growth factors, such as TGF-β. In an analysis of the signal sequence at the amino terminus of MMP3, proline was found at the fifth position of the sequence. The proline in the signal sequence can work as a helix breaker and may result in bisorting between the ER-Golgi secretion pathway and cytoplasm after the signal peptide emerges from the ribosome (44). If this partition trafficking occurred, some MMP3 would be located in the cytosol for further transport to the nucleus, while other MMP3 would be secreted through the ER-Golgi complex. This may result in both direct nuclear transport and the internalization-nuclear transportation of MMP3.

The TGF-β/Smad signal regulates CCN2/CTGF gene expression at the level of transcription, and the activation of TGF-β by MMP3 in the extracellular microenvironment can indirectly activate CCN2/CTGF through TGF-β/Smad signaling (7); however, the intracellular overexpression of MMP3 from the pMMP vector trans activated the CCN2/CTGF promoter (Fig. 3). Therefore, we consider that nuclearly translocated MMP3 directly trans activates the CCN2/CTGF gene. Interestingly, the overexpression of intracellular MMP3 and TGF-β synergistically activated the CCN2/CTGF promoter.
(Fig. 3C). If the Smads and intracellular MMP3 signals are activated simultaneously, some signal cross talk may occur, because the MMP3-binding site as well as TRENDIC is located immediately upstream of the Smad binding site (SBE) in the CCN2/CTGF promoter (7). Thus, MMP3 can regulate CCN2/CTGF transcription directly by itself and through activation of TGF-β/Smad signaling. Further investigation about this signaling cross talk is under way.

MMP3-null mice have been reported to show delayed wound healing, joint inflammation/osteoarthritis, abnormal CD4+ physiologic, increased susceptibility to bacterial infection, abnormal neuromuscular synapse morphology, and abnormal miniature end plate potential (48). CCN2/CTGF has been known to be a key player in the wound-healing process. Together with our discovery that nuclear MMP3 regulates CCN2/CTGF, the insufficient supply of CCN/CTGF in MMP3-null mice should result in a delayed wound-healing phenotype. Further investigation by using MMP3-null mice is under way.

We should be careful to explain the mechanism of trans activation by MMP3 and the novel domain structure of MMP3, because MMP3 can cleave and activate itself. The overexpression of the MMP3 deletion mutants revealed that the PEX domain and CAT+Hinge domain can trans activate the CCN2/CTGF promoter per se as well as the FL or active form of MMP3 (Fig. 5A). Catalytically dead mutants of MMP3 decreased the trans activation ability for the CCN2 promoter compared to that of the wild-type MMP3 (Fig. 5D). These results indicate that both the CAT domain and PEX domain of MMP3 can activate the CCN2 promoter independently. The catalytic activity partially contributed to the trans activation of the CCN2 promoter, however, it is not essential for the trans activation. For the mechanisms of activation, the involvement of NuMAPs was strongly suggested (Fig. 8D to G). Indeed, HP1γ synergistically trans activated the CCN2/CTGF promoter with FL MMP3 (Fig. 8E-G). The cell type-specific trans-activation mode of MMP3 for CCN2/CTGF may be ascribed to the cell type-specific involvement of cofactors such as HP1γ. Locating the HP1γ-binding domain in MMP3 also is important and is under investigation (Fig. 7A).

The DNA-binding domain of MMP3 was revealed to be in the PEX domain, because the TRENDIC-nuclear protein interaction was blocked by the anti-PEX antibody (Fig. 2A, B). In sharp contrast, the anti-N-terminal, anti-Hinge antibodies caused no significant change in the protein-DNA interactions (Fig. 2A). Moreover, no upper-shift band was observed with the other anti-MMP3 antibodies by the supershift assay (Fig. 2A). Of note, in another experiment with purified rhMMP3, the anti-PEX antibody made an upper shift with the MMP3-p160 complex (Fig. 11D). Several mechanisms can be considered in accounting for the differential effects of the anti-PEX antibody in the EMSA. For example, (i) the ZnMe/CAT domain can also bind to p160, and such a secondary DNA-binding domain may be hidden by other factors, such as NuMAPs, in the nuclear extracts; (ii) the DNA-binding ZnMe/CAT domain of nuclear MMP3 may have been cleaved away from the PEX domain; or (iii) enhanced MMP3 dimerization under the purified condition may yield a new MMP3 complex that binds to both DNA and the anti-PEX antibody.

In our study, recombinant MMP3 was translocated from outside of the cells into the nuclei (Fig. 6). Also, endogenous MMP3 was in the cells and on the cell surface (Fig. 1C and D and 9). Recent studies indicated that proMMP2 made a complex with TIMP1 and low-density lipoprotein-related protein 1 (LRP1) and then was internalized into the cells (6, 14). The LRP1/proMMP2/TIMP complex was also reported to be internalized (10). LRP1 is abundantly expressed in HCS-2/8 cells (25). We also observed that LRP1 was colocalized with MMP3 in HCS-2/8 cells by using laser confocal microscopy (data not shown). Thus, MMP3 or the MMP3/TIMP complex may bind to the trans-membrane type of receptor molecules, such as LRP1s, for internalization into cells. The delivery of certain molecules into the nucleus from the plasma membrane by endocytosis already has been reported (3). Therefore, MMP3 may be transported to the nucleus through such a pathway. The regulation of MMP3 internalization is under investigation by using LRP1 and its inhibitor-receptor-associated protein.

We also found six putative NLSs in MMP3 for nuclear entry through the nucleopores (Fig. 7). In addition, we found one putative NES in MMP3 (Fig. 7). Indeed, externally added rhMMP3 translocated to the nucleus in 30 min, whereas the amount was decreased at 60 min after the addition (Fig. 6). This loss of nuclear MMP3 may have been caused by degradation or nuclear export. LMB, which suppresses CRM1-dependent nuclear export, caused the activation of the CCN2/CTGF promoter (Fig. 6E). This result may be due to the nuclear accumulation of MMP3 caused by the blockade of CRM1-mediated nuclear export. Therefore, MMP3 can shuttle back to the cytoplasm after being translocated to the nucleus.

Of note, Si-Tayeb et al. (47) also reported an NLS in MMP3 that is the same as one of the NLSs identified in our study (Fig. 7B). It is possible that multiple NLSs are recognized synergistically for efficient nuclear translocation. Also, an NLS must be exposed on the surface of the molecule to be transported for recognition by importins for the nuclear translocation from the cytoplasm. The posttranslational modification of MMP3 can remove or hide the primary NLS and expose another NLS on the molecular surface, suggesting the differential use of multiple NLSs. We also identified a RAN-binding protein as an MMP3-associated protein. This protein is involved in the nuclear import event. As such, two independent pathways to the nucleus are indicated at present. Further investigation will specify the precise mechanism of nuclear translocation of MMP3.

We detected FL MMP3 dominantly in the nuclear fraction of HCS-2/8 cells (Fig. 1) with minor signals for its fragments, which can exert stromelysin proteolytic activity (Fig. 5). In addition to stromelysin activity in the nuclear extract of HCS-2/8 cells, there have been two reports of stromelysin activity in the nucleus (47, 54). Si-Tayeb et al. (47) reported a 35-kDa nuclear MMP3 in HepG2 cells that displayed stromelysin activity. Of note, a specific MMP3 inhibitor repressed the promoter activity of CCN2 (Fig. 4). Thus, the proteinase activity may be involved in the trans activation of CCN2 by MMP3. We speculate that an appropriate amount of nuclear MMP3 is under the control of TIMPs, so that it can work properly as a transcription factor, since we also detected TIMP-1 in the nuclear fraction of HCS-2/8 cells (data not shown). It was previously reported that TIMP-1 accumulates in the cell nuclei in association with the cell cycle (54), and multiple roles of
TIMPs other than their roles as proteinase inhibitors have been indicated, such as stimulating cell growth and acting as an antiapoptotic molecule (18). Thus, it is anticipated that MMP3 interacts with TIMPs in the cells as well as in the extracellular environment. It was reported that the intracellular overexpression of MMP3 induced apoptosis (47). Here, we have speculated about the mechanism of the apoptosis elicited by cellular MMP3. One possibility is that a vast amount of cellular/nuclear MMP3 induces apoptosis by digesting cytoplasmic and intranuclear protein components. A death-associated protein that was identified as one of the NuMAPs may be involved in the apoptosis process (data not shown). Another possibility is that the nuclear MMP3 induces the overexpression of CCN2/CTGF, which subsequently induces cellular apoptosis (22).

We identified several NuMAPs by performing the coimmunoprecipitation study (Fig. 8; Table 2). These putative cofactors included heterochromatin proteins, transcription coactivators, corepressors, RNA polymerase II, chromatin assembly factors, and others. One may expect to find MMP3 in the proteins pulled down by the beads with anti-MMP3, but we found no MMP3. We believe that the elution condition we employed dissociated MMP3-NuMAP complexes but did not interfere with stronger antigen-antibody interactions. It also is possible that TIMPs have not been detected for a similar reason. One of the NuMAPs, HP1γ, was demonstrated to interact with MMP3 and to coactivate the CCN2/CTGF promoter with MMP3 (Fig. 8). Although multiple functions of heterochromatin proteins as transcription factors and as binding targets of RNA interference machinery (19, 21) have been reported, heterochromatin proteins are known to recognize the trimethylated ninth lysine of histone H3 (H3K9), resulting in chromatin condensation (2). Moreover, abundant trimethylated H3K9, representing the active form of chromatin, was detected in the promoter regions but not in the enhancer regions (20, 43). Hence, it is possible that nuclear MMP3 binds to HP1/ Swi6, one of the NuMAPs, on the histone of the CCN2 enhancer region to trans activate the CCN2 gene. Indeed, we detected an interaction between MMP3 and the genomic CCN2 proximal promoter region by ChIP assay (Fig. 4E). Therefore, native HP1γ may collaborate with MMP3 on the genomic DNA locus in the cells.

As a NuMAP, NCoR1 also was identified, which is a transcription repressor that acts by promoting chromatin condensation and by preventing access to the gene by the transcription machinery. Another possible mechanism of CCN2 gene trans activation is that MMP3 degrades NCoR1 to prevent transcription repression. Recently, MMP2 was found in the nuclei of cardiac myocytes and is capable of cleaving PARP in vitro (28). Thus, PARP can be inactivated or activated by MMP3 itself, or MMP2 can be activated by MMP3. PARP has been known to perform poly(ADP-ribosyl)ation of the chromatin DNA for chromatin DNA maintenance. The cleavage and inactivation of PARP result in the inhibition of poly(ADP-ribosyl)ation and expose the DNase hypersensitive region. The exposure of such naked DNA can supply an MMP3-accessible region on DNA. Further genetic and epigenetic approaches will clarify the precise mechanisms of CCN2 induction by MMP3 acting with such cofactors.

In this report, we showed that MMP3 from outside of the cells and in the cytoplasm was translocated into the nucleus, where MMP3 bound to DNA/chromatin and trans activated the CCN2 gene (Fig. 12). During this process, some of the NuMAPs may have contributed to this transcription factor-like function of MMP3. Based on our present findings and numerous past reports, we propose a dual role for MMP3, i.e., its functioning both extra- and intracellularly. In this model, MMP3 acts to degrade ECM outside, whereas it induces CCN2 and other matrix-related proteins in the nucleus. This extra- and intracellular, dual-functioning MMP3 may play an important role in the development and matrix remodeling of cartilage and bone. Abnormalities in the MMP3 dynamics may be involved in the MMP/CCN2-related matrix diseases, e.g., ostearthritis and rheumatism, and in fibrotic diseases such as systemic sclerosis and atherosclerosis. Clarifying and controlling the behavior of extracellular/intracellular MMP3 may assist in the establishment of a new therapeutic approach to combat such matrix-related diseases.

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