Antiproliferative Activity of CCN3: Involvement of the C-Terminal Module and Post-Translational Regulation

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Abstract

Previous work had suggested that recombinant CCN3 was partially inhibiting cell proliferation. Here we show that native CCN3 protein secreted into the conditioned medium of glioma transfected cells indeed induces a reduction in cell proliferation. Large amounts of CCN3 are shown to accumulate both cytoplasmically and extracellularly as cells reach high density, therefore highlighting new aspects on how cell growth may be regulated by CCN proteins. Evidence is presented establishing that the amount of CCN3 secreted into cell culture medium is regulated by post-translational proteolysis. As a consequence, the production of CCN3 varies throughout the cell cycle and CCN3 accumulates at the G2/M transition of the cycle. We also show that CCN3-induced inhibition of cell growth can be partially reversed by specific antibodies raised against a C-terminal peptide of CCN3. The use of several clones expressing various portions of CCN3 established that the CT module of CCN3 is sufficient to induce cell growth inhibition. J. Cell. Biochem. 101: 1475–1491, 2007. © 2007 Wiley-Liss, Inc.

Key words: CCN3; NOV; CCN; cell growth inhibition; cell cycle; proteasome

The CCN proteins constitute a new family of secreted matricellular cell growth and differentiation regulators [Lau and Lam, 1999; Perbal, 2001, 2003, 2004; Brigstock, 2003; Planque and Perbal, 2003; Perbal and Takigawa, 2005; Rachfal and Brigstock, 2005]. Their biological properties are diverse and rely on interactions with multiple targets localized in different sub-cellular compartments. CCN proteins interact with receptors, matrix proteins, ion channels, as well as cytosolic and nuclear proteins. This led to the proposal that CCN proteins act as scaffolding proteins permitting the coordination and functional interconnection of various signaling pathways involved in the regulation of cell proliferation, differentiation, and death [Perbal, 2001, 2003; Lombet et al., 2003; Bleau et al., 2005; Planque, 2006].

The variety of functions attributed to the CCN proteins stem from their conserved tetramodular organization with structural modules showing partial identity with four classes of proteins known to play critical roles in fundamental signaling and/or regulatory pathways. Indeed, the first constitutive module resembles IGFBPs, the second shares identity with Von Willebrand protein, the third with thrombospondin (TSP1), and the fourth module contains a cystine knot that allows dimerization of various growth factors and matricellular proteins [Bork, 1993; Perbal and Planque, 2006].

Very little is known regarding the biochemical activities of the CCN proteins endowed with biological properties that are dependent upon cell origin and developmental state [Planque...
and Perbal, 2003; Perbal and Takigawa, 2005]. Studies performed with CCN3 indicated that inhibition of cell proliferation was a common theme in both normal and tumor cells of various types [Joliot et al., 1992; Gupta et al., 2001; Fu et al., 2004; Geihausen et al., 2004; Benini et al., 2005; Fukunaga-Kalabis et al., 2006; McCallum et al., 2006]. In all systems thus far tested cells that produced CCN3 were themselves responsive to its antiproliferative effects, therefore suggesting an autocrine type of response.

In order to decipher the mechanisms by which cells proliferate in spite of CCN3 anti-proliferative properties, we determined the effects of this protein on cell growth concentrating on the relationships between degree of CCN3-induced inhibition and amounts of CCN3 protein secreted during different stages of cell growth, as well as the various phases of the cell cycle. Our results indicated that the antiproliferative activity of CCN3 is modulated by a post-translational control of the amount of protein produced per cell during growth. We also identified the C-terminal module of CCN3 as sufficient to produce the growth inhibitory effects. This observation reemphasizes the critical role of this module and points to the interactions driven by this module as essential for the biological activity of CCN3.

MATERIALS AND METHODS

Construction of Expression Vectors

The pcDNA-CCN3 vectors were obtained by cloning PCR-amplified CCN3 sequences from the pCMV82 (nucleotide 1–1,943 of human CCN3 inserted at the EcoRI site of pCB6 vector) between the EcoRI sites of pcDNA6/V5-His A vector (Invitrogen) for the pCMV47 construct (nucleotide 1–1,143) [Benini et al., 2005], and between the EcoRI and XbaI sites for the PNH35 and PNH45 constructs [Planque et al., 2006]. For PNH35 and PNH45, the signal peptide of CCN3 was PCR-amplified, inserted in the EcoRI site of a Bluescript derived vector (BBg), sequenced, and then inserted at the EcoRI site of the pcDNA-CCN3 constructs. All constructs were sequenced. Figure 1 outlines the constructs and nomenclature used in this study.

The sequences encoding each exon of human CCN3 were cloned at the EcoRI and SalI sites of pTEF-MF yeast expression vector (DualsystemsBiotech). The recipient pTEF-MF was modified at its EcoRI/HindIII sites by ligating a double-strand oligonucleotide (5'-AATTGGGATCCCATCACCATCACCATCAGAATTCA-3') in order to add a carboxy-6 Histidine (6HIS) tag in the recombinant proteins. Recombinant proteins are secreted in the yeast culture medium.

Cell Lines and Establishment of Stable Clones

The glioblastoma cell line (G59) was originally isolated and provided by Dr. M. Westphal, and has been described previously [Li et al., 1996]. The G540 cells expressing human CCN3 were stably transfected by means of the pCMV-CCN3 plasmid (pCMV82) and G418 selection. G59 cells were also transfected with either pCMV-CCN3 (pCMV47), PNH35, PNH45, or pcDNA DNA. Cell transfections were performed for 48 h in 3.5 cm dishes in 3 ml of completed medium. For each dish, 2 µg of DNA were incubated with 6 µl of EXGEN500 (Euromedex) in a final volume of 100 µl for 15 min at room temperature.
temperature, according to the manufacturer's instructions. Forty-eight hours after transfection, cells were seeded in 10 cm dishes and selected by addition of 5 μg/ml of Blasticidin (Invitrogen) for about 12 days. Individual clones were picked up and tested for expression of the CCN3 protein by Western blot analysis.

Cells were grown in Dulbecco's Modified Eagle's Medium (DMEM) (Sigma) supplemented with 10% fetal calf serum (FCS) (Sigma) and L-Glutamine-Penicillin-Streptomycin solution (Sigma) (complete medium). All cells were maintained at 37°C in 5% CO2.

Production and Purification of Recombinant Proteins

To produce recombinant CCN3 proteins, the full-length CCN3 cDNA and the exon encoding the CT module (NH5) have previously been inserted in the pGEX-4T1 vector (Amersham-GE Healthcare) [Li et al., 2002], producing glutathione-S-transferase fusion proteins (GST-CCN3 and GST-NH5, respectively).

Overnight precultures were diluted 100 times in LB Broth (Lennox) medium containing 100 μg/ml Ampicillin (Sigma) and grown at 37°C until OD600 nm reached 0.7–0.9. Induction of expression of the recombinant proteins was performed by incubation in the presence of 0.1 mM IPTG for 2.5 h. After centrifugation, pellets were resuspended in 50 mM Tris pH 8.0; 1 mM EDTA; 100 mM NaCl and proteinases inhibitors: Complete Mini Tablets (Roche), 100 μM phenylmethylsulphonylfluoride (PMSF), 100 μM N-Tosyl-L-Lysine chloromethyl ketone (TLCK), 1 mM benzamidine, 100 μM N-Tosyl-Phenylalanine chloromethyl ketone (TPCK), and 300 μg/ml lysozyme was added and lysis was performed for 20 min on ice. After the addition of 1% Triton X-100, lysates were sonicated on ice. After centrifugation, supernatants were incubated with Glutathione Sepharose™ 4 Fast Flow beads (Amersham-GE Healthcare) in phosphate-buffered saline (PBS) for 1 h at 4°C on a rotating wheel. For GST-CCN3, PBS was complemented with 5% non-fat milk and 0.5 mM ATP. Beads were then washed several times with PBS-proteinases inhibitors. Recombinant proteins were recovered by three elutions of 1 h on ice with a solution containing 20 mM glutathione, 100 mM Tris pH 8.0, and 120 mM NaCl. Elution supernatants were pooled.

GST-NH5 recombinant proteins were then cleaved by thrombin and subsequently purified by immuno-affinity chromatography. To that purpose, GST-NH5 proteins were incubated in PBS containing 10 U of thrombin (Sigma) overnight at 37°C with weak agitation, then purified by two successive affinity chromatography on anti-GST and anti-NH5 Sepharose columns. NH5 proteins were eluted with 100 mM glycine buffer pH 2.8 and collected into test tubes containing 100 μl 1 M Tris buffer pH 8.0.

For the production of recombinant CCN3 modules in yeast, DSY-S yeasts were treated with polyethylene glycol and lithium acetate (PEG/LiOAc master mix) and transformed with 100 ng of each pTEF-MF recombinant vectors according to the manufacturer's instructions (DualsystemsBiotech). Yeasts were grown in appropriate selective medium (−URA). Non-tagged recombinant NH5 proteins were purified from yeast culture medium by immuno-affinity chromatography (see below). 6HIS-tagged constructs were purified using nickel columns according to the manufacturer's instructions (Invitrogen). Proteins were eluted with 500 mM imidazol.

All recombinant proteins were dialyzed overnight at 4°C against 10 mM NH4HCO3, and lyophilized. Proteins were dissolved in PBS. Quantification was performed after SDS–PAGE followed by Coomassie blue or silver staining of the gels.

Antibodies Production and Purification

The K19M antibody raised against CCN3 [Chevalier et al., 1998] was affinity purified from the immunized rabbit serum by affinity chromatography on Protein A Sepharose beads [Kyurkchiev et al., 2004]. Briefly, 1 ml of nProtein A Sepharose™ 4 Fast Flow (Amersham-GE Healthcare) was equilibrated with 20 mM Tris pH 8.0. An aliquot of 3 ml of K19M anti-CCN3 serum was loaded on the protein A column and then washed with 20 mM Tris pH 8.0 and 20 mM glycine pH 5.0. Antibody was eluted using 100 mM glycine pH 2.5 and neutralized with 1 M Tris pH 8.0. Four fractions were pooled and concentrated on Vivaspin concentrator (Vivascience) to reach a final IgG concentration of 11 mg/ml.

The GST-NH5 protein was also used as antigen to immunize rabbits (Agrobio). IgGs were purified from antisera using protein A
column chromatography. For affinity purification, antisera were passed through a GST column to remove antibodies against GST. Purified rabbit sera were tested for their reactivity against recombinant CCN3 proteins produced in yeast and mammalian cells (Lazar et al., unpublished work).

The anti-NH5 antibody was coupled to CNBr-activated Sepharose™ 4B (Amersham-GE Healthcare) following the protocol recommended by the manufacturer. Briefly, 20 μg of antibodies were dissolved and dialyzed against 100 mM NaHCO₃ pH 8.3 containing 500 mM NaCl (coupling buffer) and added to 3.5 ml CNBr-Sepharose swollen gel and the mixture was rotated end-over-end for 2 h at room temperature. Excess ligand was eluted with 20 ml of coupling buffer and the gel was incubated in 100 mM Tris-HCl buffer pH 8.0 for 2 h at room temperature. The gel was washed five times in cycles consisting of 20 ml of 100 mM acetic acid buffer pH 4.0 followed by 20 ml of 100 mM Tris-HCl buffer pH 8.0, each containing 500 mM NaCl, and then packed into a PD-10 column (Amersham-GE Healthcare).

Growth Curve of Stable Clones

Growth curves were performed by plating 2 × 10⁴ G59 and G540 cells in 6-well plates, attached cells were counted with a hemocytometer at the indicated times (48 h, 72 h, 6, 10, 13, 14, and 16 days). Medium was changed daily to maintain cells for prolonged growth studies. For counting, attached cells were trypsinized and resuspended in DMEM (200 μl to 1.5 ml depending on incubation time). Cell viability was evaluated by staining with 0.1% trypan blue. Cellular extracts were prepared and supernatants collected at each time point of the growth curve for Western blot analysis. Amounts of supernatant were normalized for the number of cells so as to establish the level of secretion per cell.

Antiproliferative Effects of Recombinant CCN3 Proteins

G59 cells (5 × 10⁵) were seeded in 24-well plates in 500 μl of complete medium. GST-CCN3 fusion protein (GST-CCN3) (0.1 μM), GST-CT module (GST-NH5), and control GST was added 2 h after seeding. Alternate sources of recombinant CCN3-CT module were also used: 0.01 μM of NH5 previously released from GST fusion by thrombin-cleavage (Thr-NH5) and NH5 produced in yeast (Yeast-NH5). Cells were counted after 5 and 7 days of incubation.

Conditioned Medium and Neutralization Experiments

To study the effect of conditioned medium on exponentially growing cells, supernatants of cell cultures were collected at two different times (240 and 312 h) and centrifuged at 1,200 rpm for 10 min to remove cell debris and detached cells. After dialysis against DMEM and filtration through 0.22 μm filters (MILLEX-GV, Millipore), the media were supplemented with FCS. G59 and G540 cells were seeded at 2 × 10⁴ cells in 24-well plates and incubated with the purified conditioned medium. Neutralization experiments were performed by plating 5 × 10⁵ G540 cells with addition of 500 μg/ml of the purified K19M antibody or 500 μg/ml of a purified anti-glycated bovine serum albumin (BSA) produced in rabbit as control. Cells were counted after 5 days of incubation. With purified conditioned media, neutralizing experiments were performed by plating 2 × 10⁴ cells in 24-well plates and incubating them for 2 days with 500 μg/ml of purified K19M antibody.

Flow Cytometry

For the analysis of cell cycle distribution, adherent and floating cells were combined and washed in cold PBS. Cells were permeabilized in 1% paraformaldehyde-PBS for 5 min on ice, washed in PBS, fixed in 70% ethanol and kept at −20°C for 30 min or until analysis. Cells were rehydrated in PBS for 10 min, washed with PBS, treated with RNase A (500 μg/ml) (Qiagen) and stained with propidium iodide (50 μg/ml). For analysis of integrin-αvβ3 expression, cells were washed with cold BSA containing 2% FCS and incubated for 45 min on ice with 5 μg/ml of anti-integrin αvβ3 antibody (Mouse monoclonal CD51, clone P2W7, Sigma). Cells were washed with PBS and then incubated with anti-mouse antibody coupled to AlexaFluor 488 (Molecular Probe) (dilution 1:200) for 30 min on ice. In all cases, a total of 10,000 events were acquired for each sample and analyzed on a standard fluorescence-activated cell sorting (FACS) caliber flow cytometer (BD Biosciences). Propidium iodide and AlexaFluor 488 fluorescence were measured by log amplification into the FL3 channel.
Proteasome Inhibition and Synchronization of Cells

For proteasome inhibition, G540 cells were treated with 10 μM MG132 (Z-Leu-Leu-Leu-al) (Sigma), 100 μM LLnL (Calpain Inhibitor I) (Sigma) or dimethylsulfoxide (DMSO) for 6 or 24 h.

For synchronization, G540 cells were arrested at (1) G0 phase by serum starvation for 48 h, (2) in early G1 phase after refeeding for 6 h with complete medium, (3) in late G1 phase after refeeding for 24 h with medium containing 200 μM mimosin (Sigma), (4) in G1/S phase after refeeding for 24 h with medium containing 2 μg/ml of aphidicolin (Sigma), (5) in S phase by washing off aphidicolin for 5 h, and (6) in the G2/M phase 30 h after refeeding with medium containing 0.1 μg/ml of nocodazole (Sigma). Kinetic analysis of cell cycle distribution was performed by plating G59 and G540 cells in the exponential phase of growth. Cells were serum starved for 48 h and arrested in G1/S phase by a 24 h of incubation with 2 μg/ml of aphidicolin. Cells were released to the next phase by washing off aphidicolin. The distribution of cells in the various phases of the cell cycle was determined by FACS analysis at 6, 9, 12, and 24 h.

Quantitative Real-Time RT-PCR (qRT-PCR)

Total RNAs were extracted from cultured human skin fibroblasts, and mRNAs levels were quantified by real-time RT-PCR as previously described [Quan et al., 2005]. Briefly, total RNA (100 ng) was reverse transcribed, with random primers, using Taqman Reverse Transcription Kit (Applied Biosystems). Real-time PCR was performed on a 7700 Sequence Detector (Applied Biosystems) using one-tenth of the reverse transcription reaction and Taqman Universal PCR Master Mix Reagents (Applied Biosystems). PCR primers and probes were from Applied Biosystems custom oligonucleotide synthesis service. Primers and FAM-labeled probe for CCN3 qRT-PCR were as follows: CCN3 sense primer, 5’-CCCGTCTGT-CAGCTGGATGTG-3’; CCN3 anti-sense primer, 5’-AGAAAAATTTGAGGGCCTGGGAG-3’; CCN3 probe, 5’-ACTGCCCTGAGCCTACTGCCAGCTC-3’. Primers and VIC-labeled probe for 36B4 were described previously [Quan et al., 2005]. Multiplex PCR reactions contained primers and probes for CCN3 and 36B4. CCN3 mRNA levels were quantified based on standard curves. For comparison among treatment groups, CCN3 levels were normalized to the housekeeping gene 36B4 (a ribosomal protein used as an internal control for quantification) levels as an internal control.

Expression of MCM3 and MCM6 mRNAs was monitored by qRT-PCR performed on G59 parental and transfected cells. Briefly, 1 μg of total RNA was denatured at 65°C for 10 min and then reverse-transcribed in a 50 μl mixture containing 1.25 U/μl MultiScribe Reverse Transcriptase (Applied Biosystems), 0.4 U/μl of RNase inhibitor (Applied Biosystems), 1× TaqMan reverse transcriptase buffer, 5.5 mM MgCl2, 2.5 μM Oligo(dT) and 500 μM of each dNTP at 48°C for 40 min. Primers were as follows: MCM3 sense primer, 5’-TACCGGCT-GATTGTCAATGTG-3’; MCM3 anti-sense primer, 5’-CAACCAAGCTCTCAGGAG-3’; MCM6 sense primer, 5’-CTCTACCTGTTG-CGGCCCTT-3’; MCM6 anti-sense primer, 5’-GCAAGTCTTGAATGCAAAC-3’. Glyceraldehyde-3-phosphate dehydrogenase (GAPDH) primers and probe as follows: GAPDH sense primer, 5’-GAAGGTGAAGGTGGCCGAGTC-3’; GAPDH anti-sense primer, 5’-GAGATTGATGGGGATTTTC-3’; GAPDH probe: FAM-GAA 5’-GTTCGGTATCTCACGGC-3’. Amplification was performed using 5 ng of cDNA and 200–800 nM primers. GAPDH was quantified using TaqMan Universal PCR Master Mix (Applied Biosystems) while the other genes used a SYBR Green PCR Master Mix (Applied Biosystems). All PCR reactions were performed by using ABI PRISM 7900 Sequence Detection System (Applied Biosystems). The target genes mRNAs were quantified by measuring threshold cycles (CT) to determine the relative expression. Data were normalized to GAPDH. The relative expression of the different mRNAs was also normalized to a calibrator, consisting of the parental cell line mRNA, and was expressed as 2^-ΔΔCT, where ΔΔCT is CT target genes—CT GAPDH, and ΔΔCT is CT sample—ΔCT calibrator.

cDNA Microarray Hybridization and Analysis

Hybridizations were performed on Human 1A (V2) Oligo microarray slides (Agilent Technologies) containing 18,716 oligo probes (60-mer length). Total RNAs were used to obtain labeled cRNAs, according to the manufacturer’s instructions (Low RNA Input Fluorescent
Linear Amplification Kit-Agilent Technologies). The G59 derived cell lines cRNAs were labeled with Cyanine 5-CTP (Cy5) (Perkin Elmer), while G59 was labeled with Cyanine 3-CTP (Cy3). In brief, 5 μg of total RNAs were reverse-transcribed with a T7 promoter primer using Moloney murine leukemia virus reverse transcriptase (Agilent Low RNA Input Fluorescent Linear Amplification Kit-Agilent Technologies). Next, cRNAs were synthesized from the double-stranded cDNAs using T7 RNA polymerase, which simultaneously incorporates Cy3 or Cy5. Labeled cRNAs were purified with RNeasy Mini Protocol for RNA Cleanup (Qiagen). After purification, Cy3 and Cy5 labeled cRNAs were fragmented by 25°C2 fragmentation buffer at 60°C in water bath for 30 min and hybridized to microarray slides at 60°C for 17 h according to the manufacturer’s instructions. The slides were washed with 6× sodium chloride–sodium phosphate-EDTA (SSPE) + 0.005% N-Lauroylsarcosine for 1 min, with 0.06× SSPE + 0.005% N-Lauroylsarcosine for 1 min and with drying solution (Agilent Technologies) for 30 s. Hybridized slides were scanned using GenePix 4000B microarray scanner (Axon Instruments) that acquires data at two wavelengths simultaneously (532 and 635 nm). Image analysis and signal intensity for both channels were calculated by GenePix 3.0 (Axon Instruments). Spots showing evident blemishes were flagged and excluded from analysis. For each array NorTT.exe software (MRC System Toxicology Group, University of Leicester) normalizes gene expression data and performs statistical Student’s t-test on each gene. Ratios between mean net fluorescence from Cy5 channel and mean net fluorescence from Cy3 channel were calculated for each spot (representing ratios between RNA expression values detected in either G59 transfectant cells and those obtained in the parental G59 cell line). Cy5: Cy3 expression ratios were log-transformed (base 2). Functional cluster result analysis was also performed by using the FatigoPlus program (Centro de Investigación Príncipe Felipe, Valencia, Spain).

Gel Electrophoresis and Western Blotting

To prepare proteins for immunoblotting, cells were lysed in NP40 buffer (50 mM Tris-HCl pH 8.0, 150 mM NaCl, 5 mM EDTA, and 2% Noniped P40) with protease inhibitors (Complete Mini Tablets, Roche) and phosphatase inhibitors (50 mM sodium fluoride, 2 mM sodium orthovanadate) for 30 min at 4°C. After centrifugation at 15,000g, extracts were stored at −80°C until use. Conditioned medium and cellular extracts (30 μg) were resuspended in Laemmli buffer containing 2-mercaptoethanol, boiled for 10 min and subjected to electrophoresis under reducing conditions in 12.5% polyacrylamide gels. Separated proteins were subsequently transferred to nitrocellulose (Hybond-C Extra, Amersham-GE Healthcare) by a semi-dry blotter (LKB Biotech) as recommended by the supplier. The nitrocellulose sheet was blocked by incubation for 1 h at room temperature with 5% non-fat milk in PBS-0.2% Tween-20 (PBST) (blocking solution: PBST-milk). The membrane was then incubated in blocking solution with either the purified anti-K19M IgG (dilution 1/2,500) for CCN3 expression, anti-p21 (dilution 1/1,000) (Santa Cruz), anti-cytochrome C (Santa Cruz) (dilution 1/500), anti-β-actin (dilution 1/1,000) (Sigma), and then extensively washed in PBST. The blots were further incubated in goat anti-rabbit or anti-mouse IgG conjugated with horseradish peroxidase (dilution 1/10,000) (Sigma) in blocking solution for 1 h at room temperature. The membranes were developed using the SuperSignal West Pico Chemiluminescent substrate from Pierce.

Statistical Analysis

Differences between two means were analyzed by Student’s t-test and between treatment groups using one-way analysis of variance (ANOVA). Differences during the growth curve were compared by t-test with Bonferroni correction.

RESULTS

Secreted CCN3 is Responsible for Inhibition of Cell Proliferation

Previous results established that human G59 glioma cells expressing CCN3 (G540) grow at a slower rate than the parental G59 cells that do not express detectable levels of CCN3 [Gupta et al., 2001]. By growing these two cell types over extended periods of time, we established that the reduction of proliferation rate associated to CCN3 expression increased with time of culture and that G540 cells reached lower saturation density (P < 0.05) than G59 cells (Fig. 2). Reduction in proliferative rate was
further evidenced by a higher percentage of cells in the G0-G1 phase and decreasing numbers in the S and G2M phase (Table I).

Recombinant CCN3 was previously reported to induce cell growth reduction [Fukunaga-Kalabis et al., 2006]. To determine whether the inhibition of proliferation was dependent upon the concentration of native CCN3, we tested conditioned media. G59 and G540 cells were incubated for 48 h in the presence of either fresh medium, or conditioned medium obtained from G540 cells grown for 240 and 312 h. Both cell lines showed a significant decrease in proliferative capability (Fig. 3A). The rate of inhibition observed with the 312 h conditioned medium was slightly greater than that induced by the 240 h conditioned medium, suggesting that active CCN3 accumulated in the cell culture medium. Conditioned medium from the G59 cells had no effect on proliferation (data not shown).

Through the use of siRNA we demonstrated that the inhibition of CCN3 synthesis resulted in an abrogation of the growth inhibitory effect attributed to CCN3 in chronic myeloid leukemia cell lines and melanocytes [McCallum et al., 2006; Fukunaga-Kalabis et al., 2006]. As a first step to determine whether the secreted CCN3 protein was indeed responsible for the growth inhibition effect on glioma cells, anti-CCN3 antibodies were added to the culture medium in which the cells were reseeded. While the addition of anti-CCN3 antibodies did not affect G59 growth in normal medium, a slight increase in cell number was observed when G540 cells were incubated in normal medium supplemented with anti-CCN3 antibodies (Fig. 3B, left panel). This moderate relief of growth inhibition might result from two opposite effects: (i) the documented partial neutralization of CCN3 activity by the CT-specific antibodies [Kyurkchiev et al., 2004], and (ii) the accumulation of inhibitory CCN3 protein that was expressed by G540 cells during the 48 h culture after replating in fresh medium. A similar increase of cell number (18%) was observed when G59 and G540 cells were grown

### Table I. Cell Cycle Distribution of G59 and G540 Cells Taken at Saturated Density of the Growth Curves

<table>
<thead>
<tr>
<th>Phase</th>
<th>G59</th>
<th>G540</th>
</tr>
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<tbody>
<tr>
<td>Sub-G0</td>
<td>0.25 ± 0.07</td>
<td>1.75 ± 0.07*</td>
</tr>
<tr>
<td>G0-G1</td>
<td>62.15 ± 0.92</td>
<td>71.50 ± 0.14*</td>
</tr>
<tr>
<td>S</td>
<td>17.05 ± 0.64</td>
<td>10.25 ± 0.21*</td>
</tr>
<tr>
<td>G2M</td>
<td>24.45 ± 0.21</td>
<td>18.3 ± 0.08*</td>
</tr>
</tbody>
</table>

*P < 0.05.
in the presence of both conditioned media (240 and 312 h) supplemented with anti-CCN3 antibodies (Fig. 3B, right panel). These results indicated that the inhibitory effect of CCN3 might be partially overcome by the anti CCN3-specific antibodies. Both cell lines responded in the same way, indicating that inhibition of cell growth involved similar mechanisms in both cases. Since we had previously established that anti-CCN3 antibodies did not recognize efficiently the native CCN3 protein [Kyurkchiev et al., 2004], only partial restoration of cell growth is observed. Incubation of cells in the presence of antibodies raised against

Fig. 3. The native secreted CCN3 protein reduces cell proliferation. A: To analyze the effects of secreted CCN3 on cell proliferation, conditioned medium from G540 cells taken at 240 and 312 h was centrifuged, dialedyzed against DMEM, filtered, and supplemented with FCS. G59 and G540 cells were seeded at 2 x 10⁴ cells in 24-well plates and incubated with the conditioned medium (from G540 taken at 240 and 312 h) for 48 h. Fresh medium was also used as control (FM). B: Neutralization of CCN3 activity was performed by plating 2 x 10⁴ G59 and G540 cells in 24-well plates and incubating for 2 days with 500 µg/ml of the purified K19M antibody or 100–500 µg/ml of an anti-glycosylated BSA (irrelevant antibody; IR) produced in rabbit. D: Cytofluorometric analysis of integrin αvβ3 expression. Gray line represents G59 cells and black line G540 cells stained with the anti-integrin αvβ3 antibody. The ordinate represents the number of cells.
glycosylated BSA had no effect on their growth rate (Fig. 3C).

The reduction in cell number induced by incubation in CCN3-containing medium was not the result of a decrease in cell attachment as estimated by cell count numbers of attached cells a few hours after seeding. In addition, G59 cells that expressed CCN3 showed a higher expression of integrin $\alpha_v\beta_3$ (Fig. 3D) that was reported to interact with CCN3 and mediate cell adhesion [Lin et al., 2003, 2005].

The Production of CCN3 is Dependent Upon Cell Density

Previous results obtained with chicken embryo fibroblasts indicated that the expression of CCN3 mRNA was increased when cells reached saturation density [Scholz et al., 1996]. Analysis of CCN3 mRNA expression in cultured G540 cells and normal human skin fibroblasts here confirmed that CCN3 mRNA expression increases with degree of confluence of the cells in a time-dependent manner (Fig. 4A).

Based on the analysis of growth curves for G59 and G540 cells, we could draw that the inhibitory effect of CCN3 increased with the number of cells. Since the culture medium was changed every other day, the greater inhibition observed at later time could not be due to an accumulation of CCN3 in the medium. Therefore, we hypothesized that a larger amount of CCN3 protein was produced per cell at higher densities, hence the greater effect upon cell proliferation. Western blotting performed on cell lysates and conditioned medium at several

![Graph A: CCN3 mRNA expression over time](image)

![Graph B: CCN3 protein expression over time](image)

**Fig. 4.** CCN3 expression increases with time of culture. **A:** Left panel: CCN3 mRNA analysis CCN3 by RT-PCR in G540 cells harvested at 72, 144, 240, and 312 h. Right panel: CCN3 mRNA analysis CCN3 by RT-PCR in normal human fibroblasts. Cells were collected at 48, 72, 120, and 168 h. In all cases, levels of the housekeeping gene 36B4 were used as an internal control. **B:** Western blot analysis of intracellular and secreted CCN3 protein from G540 cells collected at 144, 240, 312, and 336 h (left panel) and in normal human fibroblasts collected at 48, 72, 120, and 168 h (right panel). $\beta$-actin was used as the loading control.
points during cell growth indicated that the cellular production of CCN3 and secretion into the cell culture medium increase with cell number (Fig. 4B). We previously reported that secretion of CCN3 occurs rapidly after its production in the cells [Chevalier et al., 1998]. The present results indicated that the increased intracellular concentration did not result from an alteration of the secretion processes.

To determine whether the production of CCN3 was dependent upon cell density, G540 cells were collected after 192 h of growth, at a stage where they have reached a density of $8 \times 10^4$ cells/cm², trypsinized and seeded in three different-size plates to obtain cell densities of $1 \times 10^4$, $8 \times 10^4$, and $30 \times 10^4$ cells/cm² (Fig. 5A). A measurement of the amount of CCN3 produced per cell was performed by Western blotting. The results showed that the amount of CCN3 expressed per cell was dependent upon cell density, the higher production being observed at high cell density (Fig. 5B).

**CCN3 Interferes With Cell Cycle Progression and Accumulates at the G2/M Transition**

No major difference in the distribution of cells among the different phases of the cell cycle was observed in actively growing of G59 and G540 cells (data not shown and [Gupta et al., 2001]). However, FACS analysis established that a relatively high proportion of G540 cells at saturated density were in the sub-G0 state (Table I), suggesting that the high production of CCN3 was associated to apoptotic events as suggested for CCN1 and CCN3 [Todorovicc et al., 2005; McCallum et al., 2006]. In agreement with G540 cells undergoing apoptosis, cells taken at 312 h were unable to attach and grow as compared to G59 cells ($P < 0.05$) (Fig. 6A). Increased apoptosis and cell cycle arrest in G540 cells reaching confluence was confirmed by increased amounts of cytochrome C and higher levels of the cyclin-dependent kinase inhibitor p21 (Fig. 6B).

We used several inhibitors known to act at various points in the cell cycle to synchronize...
G59 and G540 cells. When cells were synchronized after release from an aphidicolin block, a differential distribution of cells was observed by FACS analysis in CCN3-positive and negative cells (Table II). Whereas 17% and about 60% of G59 cells were detected at the S and G2 phase 9 h after the aphidicolin wash, 43% and about 29% of cells were at the S and G2 phase in the G540 culture. The same type of distribution was observed at 12 h and tends to disappear with time, as cell growth becomes asynchronous. This observation suggested that the decrease in cell proliferation resulted from delayed cell cycle progression mostly at the G2/M transition.

Analysis of CCN3 production during the cell cycle was possible through several means. Serum starvation was used to induce a block at G0. Refeeding for 6 h with complete medium resets the cells at early G1. A late G1 arrest was obtained with the use of mimosin while a G1/S arrest was induced by incubation with aphidicolin. Washing off aphidicolin permitted to reset the cells at the S phase. Block at G2/M was induced 30 h after refeeding the cells with a medium containing nocodazole. At each stage, FACS analysis confirmed that cells were blocked at the expected phase of the cycle, and the quantity of CCN3 expressed by cells was measured by western blotting. While low levels of CCN3 were produced by cells at the G0 phase, the quantity of CCN3 produced gradually increased during G1 and S phase to reach a maximum at the G2/M transition (Fig. 6C).

In agreement with the higher percentage of CCN3-expressing cells at the S phase of the cell cycle, a significantly increased expression of MCM6 and MCM3 genes was detected by gene microarray analysis (not shown) and qRT-PCR (Fig. 7A) in the G59 cells stably expressing the

**TABLE II. Kinetic of Cell Cycle Distribution After Synchronization of the Cells in G0/G1 With Aphidicolin**

<table>
<thead>
<tr>
<th>Aphidicolin</th>
<th>Wash off aphidicolin</th>
</tr>
</thead>
<tbody>
<tr>
<td>Sub-G0</td>
<td>2.4% 0.6% 0.2% 0.4% 1.3%</td>
</tr>
<tr>
<td>G0-G1</td>
<td>74% 26.0% 22.8% 29.3% 61.8%</td>
</tr>
<tr>
<td>G0-S</td>
<td>91% 49.2% 17.0% 9.2% 18.0%</td>
</tr>
<tr>
<td>Sub-G2M</td>
<td>14.5% 23.8% 50.7% 69.0% 19.0%</td>
</tr>
<tr>
<td>G540</td>
<td>3.6% 0.6% 1.1% 1.3% 4.4%</td>
</tr>
<tr>
<td>G0-0.1G1</td>
<td>67.7% 31.4% 27.9% 27.3% 53.0%</td>
</tr>
<tr>
<td>G0-S</td>
<td>12.9% 52.4% 21.7% 17.8%</td>
</tr>
<tr>
<td>G0.1G1-S</td>
<td>16.5% 15.2% 28.6% 50.1% 25.0%</td>
</tr>
</tbody>
</table>

**Fig. 6.** CCN3 accumulates in the G2/M phase of the cell cycle. A: Cells were collected at 312 h, trypsinized, reseeded in large Petri dishes and counts were performed after 48 h. Data are expressed as means ± SE. *P < 0.05. B: Western blot analysis of p21 and cytochrome C in G59 compared to G540 cells. C: Western blot analysis of CCN3 and p21 expression after arrest in each phase of the cell cycle. G540 cells were blocked in (1) G0 phase by serum starvation for 48 h, (2) in early G1 phase after refeeding for 6 h with complete medium, (3) in late G1 phase after refeeding for 24 h with medium containing 200 μM mimosin, (4) in G1/S phase after refeeding for 24 h with medium containing 2 μg/ml of aphidicolin, (5) in S phase by washing off aphidicolin for 5 h, and in the G2/M phase 30 h after refeeding with medium containing 0.1 μg/ml of nocodazole. β-actin was used as the loading control.
Interestingly, the use of two vectors driving the expression of amino-truncated secreted recombinant CCN3 (PNH35 and PNH45) did not result in an increased expression of MCM genes (Fig. 7B), whereas they indeed induced a reduction of cell proliferation similar to the one observed with full-length CCN3 protein (see below). These observations indicated that the induction of MCM genes was not directly related to the antiproliferative activity of CCN3 but reflected instead the higher percentage of cells that remained in the S phase of the cell cycle.

Proteolysis Accounts for Regulated CCN3 Production

Because proteasome-dependent proteolytic degradation of several regulatory proteins occurs during cell cycle progression, we compared the levels of CCN3 produced by G540 cells in the presence and in the absence of proteasome inhibitors.

For this purpose, G540 cells were incubated in the presence of MG132 (Z-Leu-Leu-Leu-al) and LLnL (N-acety-L-leucinyl-L-leucinyl-norleucinal) proteasome inhibitors for 6 and 24 h, and the amount of CCN3 produced during these two periods of time was measured both in cell lysates and in conditioned medium. As shown in Figure 8A, incubation in the presence of proteasome inhibitors resulted in a dramatic increase of CCN3 production.

Similarly, when cells plated at low, medium, and high densities were treated by MG132 and LLnL, a considerable increase of CCN3 was observed (Fig. 8B). Furthermore, the amounts of CCN3 produced in these conditions were identical, whereas, in the absence of inhibitors, CCN3 production had been found to be cell density-dependent.

Fig. 7. Expression of MCM genes is increased in glioma cells following expression of CCN3. MCM mRNA relative expression was evaluated by quantitative real-time RT-PCR and expressed as $2^{-\Delta \Delta CT}$. The parental cell line was used as control and calibrator ($2^{-\Delta \Delta CT} = 1$). Results represent the mean ± SE of duplicate experiments. A: Induction of MCM genes in cells transfected with a plasmid expressing full-length CCN3 (pCMV47). B: CCN3 recombinant proteins lacking one or two amino-terminal modules (PNH35 and PNH45 respectively) do not induce MCM genes expression. G59-pcDNA6: G59 cells stably transfected with pcDNA; G59-pCMV47, G59-PNH35 and G59-PNH45: cells stably transfected with a pcDNA plasmid expressing either the full-length CCN3 or amino-truncated forms of CCN3.
Likewise, the use of proteasome inhibitors increased the amounts of CCN3 produced at the various phases of the cell cycle (Fig. 8C). In that case, the toxicity of the proteasome inhibitors did not permit to conduct a long term experiment with cells synchronized at G1 and G1/S, hence the relative lower amounts detected at these stages in the presence of MG132. In any case, the use of MG132 indicated that the differential amounts of CCN3 detected at S and G2/M could become identical upon inhibition of proteasome-associated proteolysis. These results indicated that in addition to the transcriptional control of CCN3 expression that occurs in the context of cell density, post-translational proteolytic degradation through the proteasome system was responsible for the regulation of CCN3 production by G59 cells and accumulation of CCN3 in confluent cells.

**The CT Module is Sufficient to Mediate an Anti-Proliferative Effect**

As the K19M antibody that can partially counteract the antiproliferative activity of CCN3 is directed against the N-terminal sequence of the CT module, we hypothesized that the carboxy-terminal domain of CCN3 might be responsible for the negative effect on cell proliferation.

As a first step to test this hypothesis, we isolated stable clones, following transfection of G59 cells with expression plasmids that carried truncated versions of the ccn3 gene. The CCN3 mutants lacking either module 1 (PNH35) or both modules 1 and 2 (PNH45) showed the same antiproliferative activity as the full-length CCN3 protein (G540 and pCMV47), as demonstrated by the growth rate of the stable clones PNH45 and PNH35 (Fig. 9A).

To identify what part of CCN3 was responsible for the inhibitory effect on cell growth, we used several sources of purified and recombinant proteins. Addition of GST-NH5 (containing the CT module of CCN3 fused to GST) to the culture medium of G59 cells resulted in a marked decrease in their proliferation (24.5% of growth inhibition compared to GST alone), though less than GST-CCN3 (44% of inhibition) (Fig. 9B). In order to eliminate the possibility that the GST moiety of the fusion proteins could be involved in the inhibitory effect, the CT domain of GST-NH5 was cleaved with thrombin (Thr-NH5) and purified by immuno-affinity chromatography. Incubating G59 cells in the presence of Thr-NH5 resulted in 80.5% of growth inhibition compared to control (addition of PBS only). In parallel, we also used an immuno-affinity-purified NH5 protein isolated from yeast culture medium. As shown in Figure 9B, both sources of CT domain (Thr-NH5 and Yeast-NH5) showed a strong inhibitory effect on G59 glioma cell growth (80.5% and about 55%, respectively), confirming that
sequences contained in this domain conferred the antiproliferative activity on the full-length protein. The other three modules produced either in bacteria or yeast had no obvious effect on G59 cell proliferation (not shown).

**DISCUSSION**

There is strong evidence that CCN3 acts as a negative regulator of cell growth in tumor cells [Gupta et al., 2001; Gellhaus et al., 2004; Benini et al., 2005; Bleau et al., 2005; Planque et al., 2005; Fukunaga-Kalabis et al., 2006; McCallum et al., 2006]. The first indication of an antiproliferative effect of CCN3 was obtained with glioma cells where the ectopic expression of CCN3 was reported to confer on transfected cells both a reduced growth rate and a decreased tumorigenicity [Gupta et al., 2001]. However, these studies could not discriminate whether or not the inhibitory effects of CCN3 had any incidence on cell cycle progression. Additional data obtained with Ewing’s and choriocarcinomas tumor cells confirmed that expression of
CCN3 was associated with inhibition of proliferation [Gellhaus et al., 2004; Benini et al., 2005]. Recent data showed that CCN3 also reduces proliferation of normal cells [Fukunaga-Kalabis et al., 2006]. These observations are well in line with previous data showing increased expression of CCN3 during the developmental differentiation of several tissues [Katsube et al., 2001; Kocialkowski et al., 2001; Perbal, 2001] and heterotypic differentiation of blastemal cells in Wilms’ tumors [Chevalier et al., 1998]. However, they also raise questions regarding the processes that allow CCN3-producing cells to grow quite well in culture and to proliferate within tissues in vivo.

In order to decipher the molecular bases for a controlled production of CCN3 we used our glioma cell model [Gupta et al., 2001] to study the production of CCN3 at the mRNA and protein levels at various stages of cell proliferation. Our results confirmed that CCN3 protein was indeed responsible for cell growth inhibition. Interestingly, cells that produced CCN3 were responsive to the secreted as well as to the purified recombinant protein, therefore indicating that post-translational modification of CCN3, such as glycosylation and phosphorylation that were previously described [Chevalier et al., 1998; McCallum et al., 2006], were not required for growth inhibition. They also suggested that CCN3-induced inhibition of cell growth was an autocrine-type of response.

While CCN3 produced a dose-dependent inhibitory effect on proliferation rate of G540 cells the degree of inhibition did not exceed 50% even when increasing amounts of purified CCN3 proteins were added to the cell culture medium. This suggests that CCN3 would inhibit cell proliferation through a rate-limiting step wherein a component is at low concentration under the assay conditions that we used. At this stage we cannot exclude the possibility that post-translational modifications could play a role in modulating the antiproliferative activity of CCN3.

It was of particular interest to observe that the amount of CCN3 protein produced per cell is dependent upon cell density and phase of the cell cycle. The inhibition of cell growth by CCN3 coupled to the increasing production of CCN3 with cell density offers a comprehensive explanation for the lower number of cells at the plateau phase that was observed with CCN3-producing cells compared to the non-producing parental cells. An opposite relationship was observed for CCN2 (CTGF) mRNA in bovine aorta endothelial (BAE) cells: a positive signal was detected in sparse, actively growing cultures, while expression was low in confluent quiescent cells. Upon removal of half the quiescent cells, CCN2 protein production increased in cells undergoing active proliferation or migration [Shimo et al., 1998]. These observations suggested that CCN proteins might compete for binding sites and that their relative concentration is a critical determinant of the biological outcome.

In this study we also report that the regulation of CCN3 expression takes place both at the transcriptional and post-transcriptional levels. As previously reported for chicken embryo fibroblasts [Scholz et al., 1996], the quantity of ccn3 mRNA expressed in cultured cells increased at confluence. At the same time, large quantities of CCN3 were secreted into the medium, as a result of a decrease in the proteolytic degradation of CCN3 that occurred during the exponential phase of growth through the proteasome pathway. The proteasome system was reported to play an important role in cell cycle regulation, cell growth, and differentiation [Dalton, 2004]. It is responsible for the degradation of the majority of cellular proteins, cell-cycle regulators, signaling molecules, tumor suppressors, transcription factors, and antiapoptotic proteins [Dou et al., 2003]. Control of mRNA decay by the proteasome pathway has also been reported [Laroia et al., 1999, 2002]. Incubation of G540 cells in the presence of proteasome inhibitors for extended periods of time, also results in an accumulation of ccn3 mRNA (data not shown). It will be interesting to determine whether these effects result from a modification of ccn3 mRNA half-life or whether they are a consequence of the cytotoxic effects of the inhibitors over long incubation time. It is tempting to propose that proteolytic degradation could provide an additional means to control the production of CCN3 so as to ensure that its inhibitory effect does not interfere with proliferation at early stages of development, when expansion of specific CCN3-producing cells is needed.

The increased production of CCN3 during the G2 phase of the cell cycle points to a direct role for CCN3 in the reorganization of the cytoskeleton and shed new light on the interactive
networks that are involved in feedback between gap junctions, adherent junctions, and cytoskeleton [Giepmans, 2006].

The CT module of CCN3 clearly stands as an essential and sufficient domain for the antiproliferative activity of this protein. This is consistent with the well-documented participation of this module in the physical interaction of CCN3 with key partners in the regulation of cell proliferation [Perbal et al., 1999; Sakamoto et al., 2002; Lombet et al., 2003; Fu et al., 2004; Gellhaus et al., 2004; Bleau et al., 2005; Planque et al., 2005; Perbal, 2006; Perbal and Planque, 2006; Planque, 2006]. The CT module is represented in five of the six CCN proteins that constitute the CCN family of proteins. This C-terminal module contains six cystein residues that are predicted to be engaged in three interlaced disulphide bonds that form a cystine knot [Perbal and Planque, 2006]. A host of proteins contains CT modules with cystine knots. Crystal structures of transforming growth factor beta (TGFβ), nerve growth factor, platelet-derived growth factor (PDGF), and human chorionic gonadotropin (HCG) all indicated that the active forms of these proteins are either homo- or heterodimers, the formation of which requires the interactions of cystine knot motif. The presence of such a motif in a large number of biologically relevant proteins raises the fascinating possibility that heterodimers may form between several of these proteins and may govern complex biological functions. In light of this model, the involvement of the CT domain in the antiproliferative activity of this protein suggests that physical partners of CCN3 may act in concert with this protein in regulating of cell proliferation. We have already established that the CT module of CCN3 is involved in the physical interaction with fibulin 1C, Notch1 and Connexin 43 [Perbal et al., 1999; Sakamoto et al., 2002; Fu et al., 2004; Gellhaus et al., 2004]. We believe that the CT domain also permits heterodimerization of different CCN proteins and that the formation of heterodimers might play a critical role in regulating the negative biological activity of CCN3. Our present study established that recombinant CCN3 proteins lacking the amino-proximal modules could still inhibit cell growth. Whether the CT module also plays a similar function in other CCN proteins remains an open question.

The ongoing characterization of proteins that physically interact with the CT domain of CCN3 is aimed at this important question. It will also be critical to establish whether the negative activity conferred by the CT module is modulated by other constitutive modules of CCN3. Once the nature of the residues responsible for the antiproliferative activity of the CT module of CCN3 is confirmed, this will hopefully open the way to the use CCN3-derived synthetic peptides as new tools in molecular medicine.

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