Knockdown of microtubule actin crosslinking factor 1 inhibits cell proliferation in MC3T3-E1 osteoblastic cells

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INTRODUCTION

Microtubule actin crosslinking factor 1 (MACF1), also known as actin crosslinking family 7 (ACF7), is a member of the plakin family of large structural proteins that link cytoskeletal elements (1-3). MACF1 is a ~600 kDa cytoskeletal linker that can bind to both F-actin and microtubules (4-6). MACF1 is widely expressed in tissues (7) and is well known for its critical role in controlling the actin and microtubule cytoskeletal dynamics (2, 8, 9). MACF1 performs key roles in numerous cell types, including epidermal cells, neuronal cells, hair cells and others (2, 8, 10, 11).

We have previously found MACF1 to be widely expressed in MC3T3-E1 osteoblastic cells and to partially co-localize with F-actin and microtubules (12). During the osteoblast response to environmental stimuli, the distribution of MACF1 and its co-localization with F-actin and microtubules were altered. Thus, we hypothesized that MACF1 might be crucial in regulating osteoblast function, based on the role of MACF1 in non-osteoblastic cells and our previous findings. However, the function of MACF1 in osteoblastic cells is not well known.

In this study, to advance our understanding of MACF1 in osteoblastic cells, we adopted a stable MACF1-knockdown cell line established with lentivirus-mediated shRNA technology in MC3T3-E1 osteoblastic cells and determined the effects of MACF1 knockdown on cell morphology, cytoskeleton distribution and cell proliferation. We demonstrated that MACF1 knockdown altered cell morphology, increased binuclear/multinuclear cell percentage, disrupted the normal distribution of F-actin and microtubules, and inhibited cell proliferation by inducing G phase cell cycle arrest in MC3T3-E1 cells. Moreover, MACF1 knockdown showed a potential effect on cellular metabolic activity. These results together, for the first time, indicate a direct role of MACF1 in regulating osteoblastic cell morphology and function.
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with the DAPI stained nucleus (Fig. S1) showed a significant increase of the binuclear/multinuclear cell percentage in the MACF1-knockdown cells (Fig. 1E). These findings suggested that MACF1 knockdown altered cell morphology and might cause cytokinesis defect.

MACF1 knockdown induced the redistribution of the cytoskeleton

MACF1 plays key role in regulating the dynamics of F-actin and microtubules (2, 8), which are essential for maintaining cell shape and involved in cytokinesis. We further determined the effects of MACF1 knockdown on the F-actin and microtubules by immunofluorescence staining. As shown in Fig. 2, in both the parental MC3T3-E1 cells and the scrambled control cells, F-actin (red, rhodamine) was distributed at the cell periphery or passed through the cytoplasm as rope-like structures, microtubules (green, FITC) radiated from a microtubule-organizing center (MTOC) to the cell plasma membrane, presenting as tortuous filaments, and the nuclei (blue, DAPI) were regularly shaped. While, in the MACF1-knockdown cells, the F-actin filaments became thinner and were mainly localized at the cell periphery, and the microtubules were reorganized, resulting in longer, straighter filaments with loss of the MTOC in association with compact α-tubulin accumulation around and in the nucleus (Fig. 2).

MACF1 knockdown inhibited cell proliferation and induced S phase cell cycle arrest in MC3T3-E1 cells

Based on the above results, we wondered whether the cell proliferation was affected by MACF1 knockdown. The cell counting detection showed that MACF1 knockdown significantly decreased the cell number after 24 h, 48 h, and 72 h of cell culture (Fig. 3A), without causing cell death (Fig. S2), suggesting an inhibitory effect of MACF1 knockdown on cell proliferation. The cell cycle analysis (Fig. 3B) showed a significant

Fig. 1. Knockdown efficiency of lentivirus-mediated shRNA on MACF1 expression in stable, transfected MC3T3-E1 osteoblastic cells and the effect of MACF1 knockdown on cell morphology. (A) Real time PCR analysis and (B) western blot detection showed significant reduction of MACF1 expression in the MACF1-knockdown (KD) cells at both mRNA level and protein level, compared with the parental MC3T3-E1 cells and the scrambled controls (**P < 0.001). (C, D) Cell morphology of the parental MC3T3-E1 cells, scrambled controls and MACF1-knockdown (KD) cells was observed using either inverted phase contrast microscopy (C) or HE staining (D) after a 24 h culture. MACF1 knockdown induced large binuclear/multinuclear cells. The white dashed line indicates the large cell border and the yellow arrows indicate the binuclear/multinuclear structures. Scale bar = 100 μm. (E) Quantification of the percentage of binuclear/multinuclear cells (*P < 0.05, **P < 0.01). The data represent the mean ± SD.

Fig. 2. MACF1 knockdown induced the redistribution of F-actin and microtubules. Cells were stained with phalloidin, tubulin, and DAPI. MACF1 knockdown disrupted the normal distribution of F-actin (red) and microtubules (green). Yellow arrows indicate the localization of the F-actin, the distribution and MTOC for the microtubules. Scale bar = 50 μm.

Fig. 3. MACF1 knockdown inhibited cell proliferation and induced S phase cell cycle arrest in MC3T3-E1 cells. (A) After 24 h, 48 h, and 72 h of cell culture, the scrambled control cells and MACF1-KD cells were harvested and automatically counted using Vi-Cell XR cell viability analyzer. The cell proliferation was significantly inhibited by MACF1 knockdown (**P < 0.05, ***P < 0.001). (B, C) Cell cycle analysis by flow cytometry (B) showed that MACF1 knockdown dramatically increased the S phase cell percentage (C), MACF1-KD versus scrambled control. *P < 0.01, **P < 0.001. (D) Real time PCR analysis showed significant down-regulation of S phase-related genes expression, MACF1-KD versus scrambled control, **P < 0.01. The data represent the mean ± SD.
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increase of the S phase cell percentage in MACF1-knockdown cells (139.4% as scrambled control, Fig. 3C). Further examination also showed remarkable reduction of the expression of critical S phase-related genes, including cyclin A2, cyclin E1, and cyclin-dependent kinase 2 (CDK2), in the MACF1-knockdown cells (Fig. 3D). These results suggested that the inhibition of cell proliferation by MACF1 knockdown might occur through the induction of an S phase cell cycle arrest.

The effect of MACF1 knockdown on cellular MTT reduction activity and mitochondrial content
As cell proliferation and cell cycling process are related with cell metabolic status, the effects of MACF1 knockdown on cellular metabolic activity was further examined using the MTT assay, which reflects cellular metabolic activity by measuring the conversion of MTT into purple-colored MTT formazan crystals by mitochondrial dehydrogenases (13). The results showed that MACF1 knockdown did not cause significant changes in the absorbance after 24 h, 48 h and 72 h of cell culture (Fig. 4A), indicating that the cellular MTT reduction activity was not affected. This finding is interesting for its disagreement with the cell proliferation results (Fig. 3A). Because the MTT assay is frequently used to detect cell proliferation based on the theory that the MTT reduction activity correlates with the number of living cells (13, 14). However, as shown in Fig. 3A, the cell proliferation was significantly inhibited by MACF1 knockdown.

To determine a possible explanation for this result, the absorbance of the same number of cells was measured. As expected, the data showed that the absorbance of MACF1-knockdown cells was substantially higher than that of the same number of the scrambled controls (P < 0.05, Fig. 4B). Previous reports have demonstrated the relationship between cell size, cell cycle and mitochondrial activity (15-17). Therefore, based on our findings, we hypothesized that the MACF1 knockdown-induced increase of absorbance may be due to the increased large binuclear/multinuclear cells. To assess this, we examined the active mitochondrial content using MitoTracker Green staining. Compared with the scrambled control, more mitochondrial content were observed in the large binuclear/multinuclear cells (Fig. 4C). Thus, our findings indicated that MACF1 knockdown increased the population of large cells, and consequently mitochondrial content and metabolic activity.

DISCUSSION
MACF1 plays key role in regulating the dynamics of F-actin and microtubule cytoskeleton, and has been demonstrated roles in numerous cell types (2, 8-10, 18). Here, we demonstrate for the first time the role of MACF1 in osteoblastic cells using MACF1-knockdown osteoblastic cells, based on our previous findings (12). The results showed that MACF1 knockdown altered cell morphology, disrupted the distribution of F-actin and microtubules, inhibited cell proliferation, and induced S phase cell cycle arrest.

Cell morphology, the basis of cell function, was first examined. The parental MC3T3-E1 cells and the scrambled control cells exhibited fusiform and polygonal morphology, while there was significant increased population of large, binuclear/multinuclear cells (Fig. 1C-E), suggesting that there might be alterations in the cytoskeleton and cytokinesis.

Fig. 4. The effect of MACF1 knockdown on cellular MTT reduction activity and mitochondrial content. (A) After 24 h, 48 h, and 72 h of cell culture, the MTT assay was conducted, showing no significant difference in the absorbance between the scrambled control and MACF1-KD cells. (B) Detection of the absorbance of the same cell number in the scrambled control or the MACF1-KD group showed increased cellular MTT reduction activity by MACF1 knockdown. MACF1-KD versus scrambled control, *P < 0.05. (C) Cells were stained with MitoTracker green and DAPI, more mitochondrial content (green) was observed in the large binuclear/multinuclear cells. Scale bar = 50 μm.
Further investigations confirmed our hypothesis. Striking F-actin and microtubule changes occurred in the MACF1-knockdown cells (Fig. 2). F-actin became thinner and was mainly localized at the cell periphery, and microtubule filaments became straight and longer associated with the disappearance of MTOC. This finding was consistent with the results of Kodama et al. (2). They also found that ACF7 (MACF1) deficiency resulted in less stable, longer microtubules associated with defective polarization of the MTOC in endodermal cells. These results indicate the importance of MACF1 in regulating cytoskeletal dynamics.

Disruption of the normal dynamics of F-actin or microtubules causes cytokinesis defects that result in binuclear/multinuclear cells and cell cycle arrest, which affects cell proliferation (19-21). Thus we wondered whether the cell proliferation would be changed by the MACF1 knockdown. As expected, MACF1 knockdown dramatically inhibited the cell proliferation (Fig. 3A). In addition, further cell cycle analysis showed that MACF1 knockdown induced an S phase cell cycle arrest in association with the down-regulation of S phase-related genes (Fig. 3B-D). These findings suggest that the inhibitory effect of MACF1 knockdown on cell proliferation may be induced through the increase of S phase arrest. Although we found that MACF1 knockdown dramatically inhibited cell proliferation and caused cytokinesis defective characteristic in MC3T3-E1 osteoblastic cells, Wu et al. reported that ACF7 absence did not cause significant decrease of cell proliferation or mitosis defects in either epidermal or endodermal cells (2, 8, 9). This difference may be due to the different cell types. Menon et al. have reported that differential knockdown of core septicin SEPT7, a cytoskeletal protein, for cytokinesis (22). Besides, as cytokinesis is a complex process that involves many proteins (23), we wonder that there may be additional as-yet unidentified osteoblast-specific proteins that interact with MACF1 in regulating cytokinesis. Further studies need to be carried out.

Another interesting finding was that MACF1 knockdown increased the cellular MTT reduction activity (Fig. 4B) as this was in contrast with the cell number result. Previous studies have reported the discrepancies between MTT assay and cell counting, and revealed that the cellular MTT reduction activity was related with mitochondrial content and activity rather than cell number (16). In addition, a strong coupling between cell size and mitochondrial content has been demonstrated (17). Moreover, there is correlation between cell cycle and mitochondrial activity, showing that cell size increases when cell entering S phase, together with increased mitochondrial activity (15). We also found a greater mitochondrial content in the large binuclear/multinuclear cells in MACF1-knockdown group (Fig. 4C). Thus, our findings suggest that the MACF1 knockdown-induced increase of cellular MTT reduction activity may be due to the increased number of large binuclear/multinuclear cells, which show more active mitochondrial content.

In conclusion, present study demonstrates for the first time the role of MACF1 in osteoblastic cells. Our results suggest an essential and positive role of MACF1 in maintaining cell morphology, cytoskeleton organization and cell proliferation. Furthermore, this work demonstrates that the inhibitory effect of MACF1 knockdown on cell proliferation may be due to a cytokinesis defect and an S phase cell cycle arrest. Moreover, present studies indicates a potential effect of MACF1 knockdown on cellular metabolic ability by increasing large binuclear/multinuclear cells, and consequently the mitochondrial content. Further studies including the in vivo experiments will be carried out in future.

MATERIALS AND METHODS

Cell culture and construction of stable MACF1-knockdown cell line

The murine MC3T3-E1 osteoblastic cells were provided by Dr. Hong Zhou of the University of Sydney. MC3T3-E1 cells were cultured in α-MEM medium (Life Technologies, USA) supplemented with 10% fetal bovine serum (FBS) (Life Technologies, USA), 100 μg/ml streptomycin and 100 units/ml penicillin, in a humidified, 37°C, 5% CO2 incubator. For the construction of the stable MACF1-knockdown osteoblastic cell line, shRNA specifically targeting murine MACF1 (NM_001199136.1) and one scrambled shRNA were designed and synthesized by Genepharma Co. Ltd (Shanghai, China). MC3T3-E1 cells were transfected with either MACF1-shRNA lentivirus vector or scrambled shRNA vector. Finally, the stably transfected cell lines were selected under the same selection condition with puromycin, and the knockdown efficiency was determined using both real time RT-PCR and western blot.

Real time RT-PCR

Real time RT-PCR was performed as previously described (12). Briefly, total RNA was extracted from cells using TRIzol reagent (Invitrogen, USA) and reverse transcribed into complementary DNA (cDNA). Then, real time PCR detection of gene expression was performed with specific primers and SYBR Green using β-actin or GAPDH as an internal control. The thermal cycling conditions included initial denaturation step at 95°C for 30 s, 40 cycles at 95°C for 10 s, 60°C for 20 s, 72°C for 5 s. The relative expression was calculated via 2^ΔΔCt method (24). The gene specific primers are: MACF1, sense: (5'-GAAACATTACCAAGTGTTCAAC-3') and antisense (5'-TGTCCATCGCCAAGGTCTTCATAG-3'); cyclin A2, sense (5'-AGTCCGCTCTTACCATGCTG-3') and antisense (5'-TCTGGTAGAAGGCTCAAGACAAG-3'); cyclin E1, sense: (5'-GCTGCGGTTCTAGTCCAA-3') and antisense (5'-GAGATGAAAGAGCAGGGTGCC-3'); CDK2, sense: (5'-TGTGGGCTTCCTCCCTGAGTAGAAG-3') and antisense (5'-CATCCTGGAAAGAAGGT-GA-3'); β-actin, sense: (5'-AGTGGAGCAGTGACATCGTGA-3') and antisense (5'-GCCAGAGCAATATCTCCTTCT-3').
Western blot

Protein extraction was performed using cell lysis buffer (50 mM Tris-base, 1 mM EDTA, 150 mM NaCl, 0.1% SDS, 1% Triton X-100, and 1 mM PMSF) on ice. Equal amounts of proteins were subjected to 6% SDS-PAGE and transferred to PVDF membrane. After incubation with the blocking buffer (5% nonfat milk), the membrane was subjected for overnight incubation at 4°C with primary antibody against MACF1 (Abcam, USA), or GAPDH (Sigma-Aldrich, USA). The horseradish peroxidase (HRP) conjugated secondary antibody was further used. Protein bands were visualized by chemiluminescence using an ECL kit (Pierce, USA) and exposed to X-ray film.

Cell morphology observation

Cell morphology was observed using an inverted phase contrast microscope (Olympus, Japan) and images were captured. In addition, HE staining was applied to determine the changes in cell morphology using the Hematoxylin and Eosin Staining Kit (Beyotime Institute of Biotechnology, China) as previously described (25). Cells (1 × 10^4/cm²) were seeded on coverslips and cultured for 24 h. Then, cells were fixed with 95% ethanol, stained with HE, and observed under a microscope (Nikon 80i, Japan) with images being captured.

Immunofluorescence staining

Cells (1 × 10^4/cm²) were seeded on coverslips. After a 24 h culture, immunofluorescence staining of F-actin, α-tubulin and the nucleus was conducted. After 15-min fixation with 4% paraformaldehyde and permeation with 0.5% Triton X-100 (TBS), the cells were stained with mouse rhodamine phalloidin (1:40, Invitrogen, USA) and anti-α-tubulin (1:20, Calbiochem, USA) overnight at 4°C. Then, the cells were incubated with goat anti-mouse FITC-IgG secondary antibody (1:100, CoWin Biotech Co, China) for 1 h. 4,6-diamidino-2-phenylindole (DAPI) (1 μg/ml) was used to counterstain the nucleus. The cells were enveloped with Fluoromount-G (SouthernBiotech, USA) and visualized using a laser confocal scanning microscope (Leica TCS SP5, Germany) or a fluorescence microscope (Nikon 80i, Japan). Images were captured, and the percentage of binuclear/multinuclear cells was determined from more than 200 cells and statistically analyzed using the GraphPad Prism software.

Cell proliferation assay

The cell proliferation and viability was detected using Vi-Cell XR cell viability analyzer (Beckman Coulter, USA) as previously described (26, 27). This technique detects the cell number and cell viability by applying trypsin blue dye-exclusion staining combined with image-based data analysis. Cells were plated into 6-well plates in triplicate at a density of 1 × 10^5/cm² and cultured for 24 h, 48 h and 72 h. Then, the cells were harvested by digestion with trypsin/EDTA and re-suspended in 1 ml medium for automatic cell counting with Vi-Cell analyzer. The cell number and percentage of living cells were determined. The percentage of living cells was presented as number of viable cells/total number of cells × 100%.

Cell cycle analysis

Cells (1 × 10^4/cm²) were incubated for 48 h and harvested. The cells were fixed in 70% ice-cold ethanol overnight at 4°C. After washing with PBS, cells were stained with PI solution (1 mg/ml sodium citrate, 0.3% Triton X-100, 0.1 mg/ml PI, 0.02 mg/ml RNase A) in the dark at 4°C for 30 min. The cell cycle was determined by flow cytometer (FACSCalibur, BD Biosciences, USA) and the percentage of cells in the G1, S and G2 phases was calculated using the ModFit software.

MTT assay

MTT assay measures the conversion of MTT into purple-colored MTT formazan by mitochondrial enzymes (13) and was performed as previously described (28). Cells were seeded into 96-well plates at a density of 1 × 10^4/cm² in triplicate. After 24 h, 48 h and 72 h of culture, 20 μl of MTT (5 mg/ml) solution was added to 96-well plates and the cells were incubated for 4 h at 37°C, followed by removal of the culture medium and the addition of 150 μl of dimethyl sulfoxide (DMSO). After a 10 min vibration, the absorbance was measured at 570 nm on a multi-functional plate reader (Bio-Tek, USA) with the medium culture subtracted as the baseline. For further investigation, the MTT formazan absorbance of the same number of scrambled control and MACF1-knockdown cells (3.2 × 10^4/well, 6.4 × 10^4/well, or 12.8 × 10^4/well) was determined. Briefly, cells at three different densities including 1 × 10^4/cm², 2 × 10^4/cm², and 4 × 10^4/cm², were seeded in 96-well plates in triplicate and the absorbance was measured after a 12-h culture.

Mitochondria detection

Live cells were incubated with 150 nM MitoTracker Green FM (Life Technologies, USA) for 45 min and the nucleus were counterstained with DAPI for 30 min at 37°C in 5% CO₂. Then, cells were imaged with a laser confocal scanning microscope using 488 nm excitation for MitoTracker Green and 405 nm excitation for DAPI.

Statistical analysis

All experiments were independently repeated at least three times with cells established in triplicate for each single assay, and the data were reported as the mean value ± standard deviation (SD). Statistical analyses of the data were performed using the GraphPad Prism software, and a student t-test was used. P values < 0.05 were considered significant and indicated in the figures.

ACKNOWLEDGEMENTS

The authors would like to thank Dr. Hong Zhou (The University of Sydney, Australia) for her generous gift of the MC3T3-E1 cell line. This work was supported by grants from the National Natural Science Foundation of China (grant number 31400725, 80i, Japan) with images being captured.

ACKNOWLEDGEMENTS

The authors would like to thank Dr. Hong Zhou (The University of Sydney, Australia) for her generous gift of the MC3T3-E1 cell line. This work was supported by grants from the National Natural Science Foundation of China (grant number 31400725, 80i, Japan) with images being captured.

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