Binding of carbon nanotube to BMP receptor 2 enhances cell differentiation and inhibits apoptosis via regulating bHLH transcription factors

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Biomaterials that can drive stem cells to an appropriate differentiation level and decrease apoptosis of transplanted cells are needed in regenerative medicine. Nanomaterials are promising novel materials for such applications. Here we reported that carboxylated multiwalled carbon nanotube (MWCNT 1) promotes myogenic differentiation of mouse myoblast cells and inhibits cell apoptosis under the differentiation conditions by regulating basic helix-loop-helix transcription factors. MWCNT 1 attenuates bone morphogenetic protein receptor (BMPR) signaling activity by binding to BMPR2 and attenuating the phosphorylation of BMPR1. This molecular understanding allowed us to tune stem cell differentiation to various levels by chemical modifications, demonstrating human control of biological activities of nanoparticles and opening an avenue for potential applications of nanomaterials in regenerative medicine.

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Nanotechnology has evolved rapidly and has an impact on nearly every aspect in modern society including medicine. For example, carbon nanotube (CNT) materials have shown promising potentials in drug1 and gene delivery,2 cancer cell imaging3 and regenerative medicine. CNT has been able to directly differentiate mouse embryonal carcinoma stem cells4 and promote neural stem cell differentiation from human embryonic stem cells.5

Adult stem cells, the undifferentiated cells residing in mature organs or tissues, can renew themselves and differentiate to cells of the specialized tissues or organs. Because of the less controversial aspect of their use than embryonic stem cells, the easy accessibility, and their capability to divide in cell cultures, adult stem cells provide valuable cell sources to treat and cure many regenerative diseases. Induction of myogenic differentiation from different adult stem cells is promising to cure diseases like muscular dystrophies,6 cardiac myocyte death7 and muscle damage. For example, mesenchymal stem cells identified from bone marrow have been successfully used for repairing heart muscle in cardiac ischemia.7,8 However, this application is limited by the poor viability of the transplanted cells caused by the pathological milieu and loss of survival signals from matrix attachments and cell–cell interactions.9 In order to increase the survival of stem cell grafts, some transgenic strategies (for instance, by engineering stem cells to express AKT, a serine threonine kinase and a powerful survival signal) have been pursued.10 In spite of these endeavors, there is still a need to develop strategies to enhance differentiation in a controllable manner and improve the viability of transplanted cells. C2C12 is a mouse myoblast cell line, which is capable of myogenic or osteogenic differentiations.11 In this work, we investigated the myogenic differentiation of C2C12 under the effects of carboxylated multiwalled CNT (MWCNT 1) and studied the involved molecular mechanisms. The results demonstrated that MWCNT 1 promoted myogenic differentiation of C2C12 without using any protein or chemical factor. MWCNT 1 also improved cell survival under the differentiation conditions. These effects were originated from the binding of MWCNT 1 to bone morphogenetic protein receptor 2 (BMPR2), suppressing the BMP signaling pathway and regulating basic helix-loop-helix (bHLH) transcription factors involved in differentiation and apoptosis. Based on this mechanistic understanding, we successfully modulated myogenic differentiation of C2C12 to various levels by surface-chemistry modifications on MWCNTs.

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Abbreviations: bHLH, basic helix-loop-helix; BMP, bone morphogenetic protein; BMPR, bone morphogenetic protein receptor; BRE, BMP-responsive element; BSA, bovine serum albumin; ChIP, chromatin immunoprecipitation; CNT, carbon nanotube; DAPI, 4′,6-diamidino-2-phenylindole; DM, differentiation medium; DMEM, Dulbecco’s minimum essential medium; FITC, fluorescein isothiocyanate; FRET, fluorescence resonance energy transfer; GAPDH, glyceraldehyde-3-phosphate dehydrogenase; GM, growth medium; Id, inhibitor of DNA-binding/differentiation; PLA, proximity ligation assay; MWCNT, multiwalled carbon nanotube; MYH, myosin heavy chain; p-BMPR1A, phosphorylated BMPR1A; RT-PCR, reverse transcription PCR; qRT-PCR, quantitative reverse transcription PCR; TEM, transmission electron microscope; TRITC, tetramethyl rhodamine isothiocyanate; TUNEL, terminal deoxynucleotidyl transferase dUTP nick end labeling

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Results

MWCNT 1 promotes myogenic differentiation and inhibits apoptosis. The physicochemical characterization of MWCNT 1 was summarized in Table 1. MWCNT 1 has an inner diameter of 5–12 nm and an outer diameter of 30–50 nm. On the basis of the functional group loading (0.4 mmol/g) as determined by the elemental analysis and measured surface area (500 m²/g), we estimated that the average density of modification is approximately five functional groups per 10 nm². At pH 7.0, MWCNT 1 carries negative charges. Its solubility in water and plasma is much better compared with the pristine MWCNT. Inductively coupled plasma mass spectrometry analysis indicated a negligible amount of metal impurities excluding the possibility that they may affect cellular function.

C2C12 can differentiate into adipocytes, muscle cells or osteoblasts under different conditions. It has been reported that C2C12 cells underwent myogenic differentiation after transventricular injection into the arterial circulation of the murine heart. MWCNT 1 showed little cytotoxicity at concentrations up to 100 µg/ml (Supplementary Figure S1). To mimic the differentiation microenvironment of transplanted cells, C2C12 cells were incubated in the differentiation medium (DM) to evaluate their myogenic differentiation under the influence of MWCNT 1. C2C12 cells when treated with MWCNT 1 expressed myosin heavy chain (MYH) protein, formed myotubes and showed a dose-dependent increase in fusion index compared with DM controls (Figures 1a–c). These data demonstrated that MWCNT 1 accelerated the myogenic differentiation under the differentiation environment.

Differentiation conditions (low serum) deprived mitogens of cells and caused an increase in apoptotic cells (20–30%). Microscopic observation indicated that the cell death was significantly decreased by MWCNT 1 treatment (Figures 1d and e). Terminal deoxynucleotidyl transferase dUTP nick end labeling (TUNEL) assay showed that MWCNT 1 treatment decreased the percentage of TUNEL-positive cells from 23 to about 8% in 48 h (Supplementary Figure S2 and Figure 1f).

Table 1 Characterization of MWCNT-COOH

| Parameter                  | Value                      |
|----------------------------|----------------------------|
| Diameter (nm)              | 30–50 (outer); 5–12 (inner) |
| Length (µm)                | 0.5–2.0                    |
| Zeta potential (mV in H²O) | −42.5                      |
| Zeta potential (mV in plasma) | 9.3                      |
| Functional group loading (mmol/g) | 0.4                      |
| Fe, Al, Cr and Mn (%) by ICP-MS | 0.3, 0.02, 0.002 and 0.00001 |
| Other 19 metals (%) by ICP-MS | <0.00008                  |

The activity of apoptotic effector protein caspase 3 dropped to nearly control level (1.2-fold compared with that in growth medium) (Figure 1g). These data showed that MWCNT 1 promotes myogenic differentiation as well as inhibits apoptosis of C2C12 under the differentiation conditions.

The myogenic differentiation of C2C12 was coordinated by a family of proteins containing bHLH structures including MyoD, myogenin, myf5, mrf4 and E2A proteins such as HEB, E12 and E47. The inhibitor of DNA-binding differentiation (Id) proteins are negative regulators of myogenic differentiation. The Id proteins form a complex with myogenic regulatory factors and inhibit their binding to DNA to activate the expression of muscle-specific genes. We first evaluated whether MWCNT 1 promoted myogenic differentiation by altering functions of these two families of proteins or their interactions.

MWCNT 1 positively regulates functions of bHLH proteins. We first analyzed the effects of MWCNT 1 on myogenic regulatory factors and Id proteins. Because of the functional redundancy of MyoD with mrf4 and myogenin with myf5, we analyzed the protein levels of two myogenic regulatory factors MyoD and myogenin and one E2A protein HEB. The expression level of MyoD and HEB did not change, whereas the expression of myogenin increased (Figure 2a). In the normal medium, MWCNT 1 treatment decreased the expression of all the three Id genes on both mRNA and protein levels (Supplementary Figure S3a and Figure 2b). Under the differentiation conditions, MWCNT 1 treatment also downregulated Id1 and Id3 at the mRNA level (Supplementary Figure S3b). MyoD is known as a marker for myogenic commitment and its major function is to induce the expression of myogenin and cell-cycle inhibitors such as p21. MyoD forms a complex with E2A protein and activates the expression of target genes by binding to E-box motifs in the promoter region of these genes. However, because of their higher affinity to E2A proteins, Id proteins compromise the MyoD–E2A complex formation and block the gene expression by binding to E2A proteins. The nanotube-induced increase in myogenin expression and decrease in Id proteins expression suggested that MWCNT 1 might abrogate the formation of HEB–Id complex, thus favoring the formation of HEB–MyoD complex. Co-immunoprecipitation with HEB followed by probing the co-precipitated proteins with anti-Id1 and anti-MyoD antibodies suggested that MWCNT 1 might abrogate the formation of HEB–Id complex. MWCNT 1 treatment recruited more HEB–MyoD complex to the cell nucleus compared with the normal medium (Figure 2c). In order to further validate that the enhanced myogenin-expression level after MWCNT 1 treatment is indeed caused by the elevated HEB–MyoD complex formation, chromatin immunoprecipitation (ChIP) assay with anti-HEB and anti-MyoD antibodies was carried out. The results indicated that MWCNT 1 treatment recruited more HEB–MyoD complex to the E-box region of the myogenin promoter (Figure 2d). As a consequence of the increased DNA binding by HEB–MyoD complex, the activity of the myogenin promoter was increased by six-folds (Figure 2e).

Myogenin triggers the expression of muscle-specific genes by replacing MyoD and forming a dimer with HEB after the initiation of differentiation (24–48 h under the
A significant increase in HEB–myogenin complex formation after MWCNT 1 treatment was observed (Figure 2c). A subsequent event is the nuclear translocation of myogenin to activate the expression of muscle-specific genes. Immunocytochemistry showed that MWCNT 1 accelerated both the expression and nuclear translocation of myogenin (Figures 2f and g). Together, these data showed that by negative regulation of Id proteins, MWCNT 1-enhanced positive regulation of cell-differentiation machinery.

p21Cip1/waf-mediated apoptosis inhibition. We were also interested in investigating how MWCNT 1 suppressed cell apoptosis. Cdk inhibitors, such as p21Cip1/waf (hereinafter p21), can protect cells from apoptosis, and p21 expression was controlled by the bHLH proteins. We found that MWCNT 1 treatment upregulated the p21 promoter activity (Supplementary Figure S4a) and recruited more HEB–MyoD complex to bind to p21 promoter (Supplementary Figure S4b). Western blot analysis showed an increased expression of p21 from 24 to 72 h after MWCNT 1 treatment (Figure 3a). On the basis of these results and the previous reports, we hypothesized that the MWCNT 1-induced elevation of p21 expression accounted for the apoptosis suppression. p21 inhibits apoptosis only when it is translocated from nucleus to mitochondrial membrane. To verify the mitochondrial localization of p21, we labeled mitochondria with a mouse-originated anti-Core II antibody and a Cy3-labeled secondary antibody, and p21 was labeled with rabbit-originated anti-p21 antibody and a fluorescein isothiocyanate (FITC)-conjugated secondary antibody. Our results showed that p21 proteins were mainly localized in mitochondria in MWCNT 1-treated cells. In the untreated group, p21 proteins resided in the nucleus (Figure 3b). At the mitochondrial membranes, p21 forms a complex with procaspase 3 and inhibits its activation. The enhanced p21–procaspase 3

Figure 1 MWCNT 1 promotes C2C12 differentiation and inhibits apoptosis. (a) Time-dependent MYH protein expression assayed by western blot with anti-MYH antibody. (b) Myotube formation in C2C12 cells cultured in DM or MWCNT 1 (25 μg/ml) for various time points. Immunocytochemistry was performed with anti-MYH antibody and FITC-conjugated secondary antibody (scale bar: 200 μm). (c) Quantitative analysis of time-dependent cell differentiation under MWCNT 1 concentrations of 25 and 100 μg/ml assayed by nuclear fusion index (*P < 0.05). (d) and (e) Phase-contrast photomicroscopy revealed cell apoptosis after 48 h in DM and MWCNT 1 concentration of 25 μg/ml. (f) Quantitative analysis of BrdU-positive nuclei. The percentage of BrdU-positive nuclei in 0 h was also shown. (g) Quantitative analysis of caspase 3 activity. Caspase 3 activity in GM group was normalized to 1.0 (*P < 0.05). GM, growth medium.
complex formation was observed in a co-immunoprecipitation assay (Figure 3c). MWCNT 1 also inhibited caspase 3 activity (Figure 2f). Therefore, our findings suggested that upregulation of the expression of p21 proteins was at least one mechanism to protect cells from apoptosis under the differentiation conditions.

The above data demonstrated that MWCNT 1’s downregulation of Id protein levels has a critical role in promoting myogenic differentiation and inhibiting apoptosis of C2C12. The Id proteins are the targets of the BMP signaling pathway. We have reported that CNTs inhibit the BMP signaling pathway and this was confirmed by...
an independent group.\textsuperscript{29} Here we found that MWCNT 1 also inhibited Id protein expression by suppressing the BMP signaling pathway. In order to gain further understanding of interplays between MWCNT 1 and the BMP signaling pathway, we investigated the further details in this context.

**MWCNT 1 perturbs the phosphorylation of BMPR1.** BMP signaling is suppressed when the initial membrane-receptor activation or any of the following signal transduction steps is partially blocked.\textsuperscript{30} In order to reveal how MWCNT 1 blocked BMP signaling, we examined the effects of MWCNT 1 on individual signaling steps.\textsuperscript{31} We first analyzed the phosphorylation of Smad1/5/8. Western blot indicated that while the level of Smad1/5/8 was unchanged, MWCNT 1 treatment decreased their phosphorylation level (Figure 4a), suggesting that Smad1/5/8 phosphorylation or an upstream step was affected. In cells, Smad1/5/8 was phosphorylated by activated BMPR1. In order to clarify whether MWCNT 1 affected any step downstream of BMPR1 phosphorylation, a transgenic cell line C33A-2D2 containing firefly luciferase reporter driven by Id1 promoter was constitutively phosphorylated Smad1/5/8 and transduced intracellular signals.\textsuperscript{30} Incubation of both the transfected cell lines with MWCNT 1 at various concentrations for different times showed no dose-dependent inhibition on Id1 promoter activity, whereas the specific Smad1/5/8 inhibitor dorsomorphin did (Figures 4c–f). These results suggested that MWCNT 1 did not interfere with any step downstream of BMPR1.

Phosphorylation of BMPR1 by BMPR2 after ligand binding is the first step to trigger the intracellular BMP signaling cascade.\textsuperscript{31} In order to investigate whether MWCNT 1 interferes with the phosphorylation process of BMPR1, the phosphorylated BMPR1A in whole-cell extract was determined after co-immunoprecipitation with its antibody. The data showed that MWCNT 1 strongly suppressed BMP4-induced phosphorylation of BMPR1A (Figure 4b), suggesting that MWCNT 1 inhibited BMPR1 phosphorylation by BMPR2.

**MWCNT 1 binds to BMPR2 and suppresses BMP signaling process.** The suppression of BMPR1 phosphorylation by MWCNT 1 suggested that its target might be membrane-bound BMP receptors. Under the transmission electron microscope (TEM), MWCNT 1 bound to cell surface or was endocytosed into cells with continued membrane interactions (Figures 5a and b). We first investigated the spatial proximity of MWCNT 1 with BMPRs. To do this, we conjugated mouse IgG to MWCNT 1. By an \textit{in situ} proximity ligation assay (PLA),\textsuperscript{32} using rabbit antibodies to BMPR1A...
and BMPR2, we detected a close proximity between MWCNT 1 and BMPR2 (10.6 PLA signals per nuclear, compared with 20.5 PLA signals per nuclear in positive control of BMPR1A/BMPR2 and 1.8 PLA signals per nuclear in negative control of bovine serum albumin (BSA)-conjugated MWCNT 1 (Supplementary Figures S5a and b) other than with BMPR1A (4.0 PLA signals per nuclear) (Figures 5c and d). This was likely due to a direct binding between MWCNT 1 and BMPR2. In order to substantiate MWCNT 1/BMPR2 binding, we also determined fluorescence resonance energy transfer (FRET) between fluorescent-labeled MWCNT 1 and BMPR2. MWCNT 1 was labeled with FITC-conjugated BSA (FITC-BSA) and BMPR2 was labeled with tetramethyl rhodamine isothiocyanate (TRITC)-conjugated secondary antibody. FRET acceptor bleaching method measures the donor ‘de-quenching’ in the presence of acceptor by comparing donor fluorescence intensity before and after destroying the acceptor by photobleaching. If FRET is initially present, a resultant increase in donor fluorescence will occur on photobleaching of the acceptor. We found that photobleaching TRITC resulted in an increase in the FITC intensity with an efficiency of 15–20% (Figures 5i–n). This increased efficiency is comparable to that reported in interaction between two fluorescent proteins; hence this result confirmed the binding between MWCNT 1 and BMPR2. This binding event abrogated the BMPR1A–BMPR2 heterodimer formation as shown also by PLA using antibodies to BMPR1A and BMPR2 after BMP4.
The interruption of the formation of BMPR1A–BMPR2 heterodimer markedly suppressed the BMP4-induced phosphorylation of BMPR1A as analyzed by immunoprecipitation (Figure 4b). We also found that MWCNT 1 in culture system neither adsorbed BMP4 ligand (Figure 5e) nor affected their binding to BMPRs (Supplementary Figures S5c–f). These data demonstrated that MWCNT 1 affected the BMP signaling pathway by binding to BMPR2 and compromising its signaling function.

Our finding can be summarized into a working model (Figure 6). By binding to BMPR2 and inhibiting the phosphorylation of BMPR1, MWCNT 1 downregulates the phosphorylation of Smad1/5/8 and reduces the expression of Id proteins. The decrease in Id proteins expression enhances the formation of HEB–MyoD complex, which in turn promotes the expression of myogenin and p21. Myogenin forms complex with HEB to activate the differentiation-specific genes (e.g., MYH) and leads to myogenesis. p21 proteins translocate to mitochondria to bind and inhibit the activation of proapoptotic 3, thereby suppressing apoptosis.

**Fine tuning of cell differentiation by surface modifications on MWCNT 1.** Biological perturbations by nanoparticles can be attributed to their size, shape, structure and surface properties. When the size, shape and structure of nanoparticles are kept the same, their surface chemistry will dictate the surface properties such as hydrophobicity, stereochemistry and charges. The altered surface properties of CNTs determine their interactions with proteins and other biological molecules. We hypothesized that the interactions of CNTs with BMPRs might be modulated by their surface chemistry. We therefore examined a diverse array of surface-modified MWCNTs, which contains 84 members (MWCNT 1–84) for their effects on the activation of BMPRs as indicated by Id1 promoter activity in C33A-2D2 cells. MWCNTs with diverse surface-chemistry modifications generated various Id1 promoter activations (Supplementary Figure S6 and Figure 7b). To explore whether we could tune cell differentiation by modulating BMP signaling activation, we further investigated MWCNTs 6, 7, 11, 12, 20 and 43 (Figure 7a) for their effects on cell signaling and differentiation. First, we found that they enter cells in a similar manner.
as MWCNT 1 as shown by TEM (Supplementary Figure S7a). Similar to MWCNT 1, they did not inhibit caALK-induced Id1 activation in C33A-2D2-ALK2 or C33A-2D2-ALK3 cells (Supplementary Figure S7b). These MWCNTs also showed various inhibitions on BMPR1A phosphorylation (Figure 7e). Furthermore, they downregulated phosphorylation of Smad1/5/8 and the expression of Id proteins in the same order (Figure 7e). These data strongly showed that surface-chemistry modifications conferred different modulations on BMPR and caused different levels of suppression on the BMP signaling pathway. Therefore, this approach is promising for fine tuning of cell differentiation through modulating BMP signaling to a desired level. To test this possibility, we analyzed the differentiation profile of C2C12 cells. Our data showed that the differentiation of C2C12 cells was regulated to various levels as shown by the inversely correlated myogenin promoter activity (Figure 7c) and different expression levels of both myogenin and the differentiation-marker MYH proteins (Figure 7e). Expression of the apoptosis-inhibitor p21 was also regulated (Figure 7e). As a quantitative measure for terminal differentiation, myotube formation and the nuclear fusion index were also modulated to various controllable levels (Figures 7f and d).

Discussion

Rapid and controllable induction of differentiation and decreasing cell apoptosis are both desired attributes of medicinal agents in regenerative medicine. These two aims have been sought out by various strategies without much success. Nanomaterials hold enormous promises in this aspect because they exhibit suitable physical characteristics and they are able to mimic surface properties of natural tissues. Therefore, the molecular understanding of the effects of nanoparticle on stem cell differentiation is critical to fully develop their application potential.

In a previous study, incubation of CNT with cell lysate enhanced protein binding and MS analysis revealed that CNT bound to murine scavenger receptor macrophage receptor with collagenous structure in addition to other proteins.
Our study provides the evidence that nanoparticles bound to cell-membrane receptors in live cells. Discoveries from this work in combination with previous reports demonstrate how CNTs alter cell functions such as differentiation and apoptosis (Figure 6). The binding of CNTs to BMPR2 weakens the interactions between BMPR2 and BMPR1 and inhibited BMPR1 activation. Such weakening is propagated through the BMP signaling pathway as shown by a downregulation of Smad phosphorylation, Smad nucleus translocation and Id protein expression. Reduced Id expression in cell nucleus changes the regulation of HLH transcription factors and upregulated cell-differentiation genes and p21, which suppresses cell apoptosis. The combined effect is the enhanced cell myogenic differentiation.

The driving forces for CNTs/protein binding are likely electrostatic, hydrophobic and π–π stacking interactions. Such interactions are probably not specific. In fact, CNTs perturbed cell genome by up- and downregulation of more than 400 genes and the BMP pathway is only one of the top ten most-affected signaling pathways. However, one significant finding from this work is the identification that perturbations on the BMP signaling pathway are resulted from interactions of CNT with BMPR2. On the basis of this finding, chemical modifications on CNTs to alternate its interactions with BMPR2 become possible. By carrying out chemical modifications on CNT surface, we successfully modulated myogenic differentiation of C2C12 to various levels by controlling CNT–BMPR2 interactions. Chemical basis for

![Figure 7](image-url)
such regulation was explored by analyzing the relationships between various structural parameters of CNTs’ surface molecules and the Id1 promoter activities. Preliminary studies showed that molecular surface areas and polar surface areas exhibited some correlations with the effects of CNTs on BMP signaling. However, the correlation was not very tight, suggesting that more complex factors such as other molecular and CNT interactions were also involved in the regulation of cell differentiation.

Although the biological impacts of nanomaterials are mostly unknown, and some effects are found to be harmful, this work points out that beneficial effects can be initiated by CNTs. This investigation characterized in molecular terms how CNTs promote cell myogenic differentiation by binding to BMPR2 and suppress BMP signaling. Furthermore, we successfully modulated effects of CNTs on differentiation, modifying surface chemistry on CNTs. By this result, we demonstrate the human control of the biological impacts of nanomaterials and open an avenue for the applications of nanomaterials in regenerative medicine.

Materials and Methods

Materials, cell lines, reagents and antibodies. MWCNT 1 and the combinatorial array of surface-modified MWCNTs were prepared as reported.37 Stock solutions of MWCNTs (1 mg/ml) in ddH2O were prepared and stabilized at 121 °C for 30 min and sonicated before use.

C2C12 cells <20 generations were grown in growth medium (Dulbecco’s minimum essential medium (DMEM) medium supplemented with 20% fetal bovine serum) and were placed in DMEM medium supplemented with 2% horse serum) when culture reached 80% of confluency (t = 0 h). C33A-ZD2 was constructed using human cervical carcinoma cell line C33A. Single-cell clones from C33A that responded to human recombinant BMP4 were selected after they were transfected with a multimerized BMP-responsive element (BRE) from the mouse Id1 promoter linked to the firefly luciferase gene (pGL4-BRE-Luc). The EC50 value for BMP4-induced specific BRE-mediated activity was 1 ng/ml. The control C33A line expressing the pGL4-luciferase vector only (C33A-empty vector) was also constructed as a control for all the assays.

Constitutive active ALK2 and ALK3 constructs were gifts from Paul B Yu, Harvard Medical School. Myogenin-luciferase and p21-luciferase constructs were gifts from Michael A Rudnicki, Ottawa Health Research Institute, and Carme Gallego, Universitat de Lleida.

Lipofectamine 2000 Transfection Reagent, Opti-MEM I Reduced-Serum Medium, and RIPA Buffer (10 ×, no. 9806) were purchased, respectively, from Invitrogen (Grand Island, NY, USA) and Cell Signaling (Danvers, MA, USA). The luciferase assay system (E1500) and the ChIP assay kit (no. 17–295) were purchased, respectively, from Promega (Madison, WI, USA) and Millipore (Billerica, MA, USA). Caspase3/GPP32 Colorimetric Assay Kit (k106–100) was purchased from BioVision (Milpitas, CA, USA).

Goat anti-mouse and goat anti-rabbit secondary antibodies (no. 926–32 220 and 926–32 221) were used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for immunocytochemistry were commercial, obtained from Jackson ImmunoResearch (West Grove, PA, USA). Primary antibodies used in immunocytochemistry were obtained from BioVision (Milpitas, CA, USA) and Cell Signaling (Danvers, MA, USA). The fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA). All the fluorescent-labeled secondary antibodies used for western blot analysis are products of LI-COR Biosciences (Lincoln, NE, USA).

Calculation of nuclear fusion index. The nuclear fusion index was calculated by the number of nuclei in cells containing two or more nuclei in total nuclei × 100%. For each experiment, 300–500 nuclei were counted.

Primers for PCR. Id1, forward: 5’-GCTGAGGCTGCTAGTGTT-3’, reverse: 5’-TGGAACATCAGCCTGTCGGG-3’ (391 bp); Id2, forward: 5’-GCAATCTGCTGCTGACCTGGC-3’ (376 bp); Id3, forward: 5’-CTCAAGGCCTGGTCTGCG-3’, reverse: 5’-TGCTGCTGGAGGGAGTGACG-3’ (329 bp); and β-actin, forward: 5’-AAATCTGGTGATGATCAAAA-3’, reverse: 5’-AGAAAGAAGCTGCGAAAAAG-3’ (180 bp). Primers for ChIP assay, p21, forward: 5’-ACAGTGGTGAGCAAGACACCA-3’, reverse: 5’-TCTGGCTGCTGAGATGTTGG-3’ (305 bp); myogenin, forward: 5’-GAATCACTGATTAACCCAGTGA-3’, reverse: 5’-ACCACCACTGGTTGGCACA-3’ (180 bp) (E-box motif sequences highlighted by red); and β-actin, forward: 5’-AAATCTGGTGATGATCAAAA-3’, reverse: 5’-AGAAAGAAGCTGCGAAAAAG-3’ (180 bp).

Semi-quantitative reverse transcription PCR (RT-PCR) and quantitative RT-PCR (qRT-PCR). RNA was extracted with an RNeasy Mini Kit and CDNA was synthesized with SuperScript III first-strand synthesis system for RT-PCR (both kits are commercial, obtained from Invitrogen). For semi qRT-PCR, 1.5 µl of cDNA and 5 µl of primers (10 µM) were used for one reaction. A total of 20 cycles were run with a C1000 thermal cycler (Bio-Rad, Hercules, CA, USA) and each cycle consisted of denaturation at 94 °C for 30 s, annealing at 62 °C for 30 s and extension at 72 °C for 1 min. qRT-PCR was run in a 7900HT fast real-time PCR system with the following cycling conditions: enzyme activation at 95 °C for 20 s; 40 cycles for denaturing at 95 °C for 1 s; and anneal/extend at 60 °C for 20 s. Finally, 0.25 µl of CDNA and 0.2 µl of primers (10 µM) in one 10 µl reaction were used with Fast SYBR Green Master Mix (Applied Biosystems, Carlsbad, CA, USA). Every reaction was run in triplicate in a 384-well clear optical reaction plate. All the instruments and reagents used for real-time PCR were from Applied Biosystems.

TUNEL assay. TUNEL assays were conducted with an ApopTag Plus Fluorescein In Situ Apoptosis Detection Kit (S7111; EMD Millpore, Billerica, MA, USA) following the indicated protocol. Briefly, cells were treated as indicated, fixed in 1% paraformaldehyde, washed in PBS three times, incubated in equilibration buffer for 10 min and incubated with terminal deoxynucleotidyl transferase for 60 min at 37 °C. Cells were washed with stop buffer for 10 min and then incubated with anti-digoxigenin conjugate for 30 min at 25 °C. After washing in PBS, cells were mounted with DAPI and viewed and photographed on a microscope with a fluorescent attachment.

Caspase 3 activity assay. After being exposed to DM with or without CNTs for 24 h, cell lysates were prepared and 200 µg of proteins were used for caspase 3 activity assays. Samples were read at 400 nm in a microtiter plate reader and fold increase in caspase 3 activity was determined by comparing with the level in cells cultured in growth medium.

Co-immunoprecipitation. Cell lysates were extracted with 1 × RIPA Buffer, and 200 µg of lystate was used for each immunoprecipitation. Before precipitation (ab27504) was from Abcam (Cambridge, MA, USA). Glyceraldehyde-3-phosphate dehydrogenase (GAPDH; G9545) antibody was purchased from Sigma-Aldrich (St. Louis, MO, USA). The Id1 (BCH-1/372-50), Id2 (BCH-3/2-4-8-50) and Id3 (BCH-4/6-1-50) antibodies were from BioCheck (Foster City, CA, USA). Complex III subunit Core 2 monoclonal antibody (MS304-SP) was commercial, obtained from MitoSciences (Eugene, OR, USA).

Immunocytochemistry. For immunocytochemistry assays, cells were seeded on 22 × 22-mm cover slips (Corning Life Sciences, Tewksbury, MA, USA) and treated with poly-c-lysine. After fixing with cold 4% paraformaldehyde in PBS for 15 min and permeabilization with 0.25% Triton X-100 for 10 min, cells were blocked with 1% BSA in PBS containing 0.5% Tween-20 for 30 min followed by 1 h incubation with primary antibodies and 1 h incubation with FITC- or Cy3-conjugated secondary antibodies at room temperature. Cells were rinsed and mounted on cover slips with mounting medium containing 4′,6-diamidino-2-phenylindole (DAPI; Vector Lab, Burlingame, CA, USA). Fluorescence images were taken using a Zeiss (Peabody, MA, USA) LSM 510 confocal microscope with a × 40 objective lens.
with primary antibodies, lysate was preclarified by adding 1.0 μg of the appropriate normal mouse or rabbit IgG (corresponding to the host species of the primary antibody) together with 20 μl of suspended Protein A/G-Plus-Agarose. Each precipitation used 1 μg of primary antibodies and 20 μl of Protein A/G-plus-agarose. After centrifugation, samples were boiled in 1% SDS solution and analyzed by western blot. All western blots were imaged with an Odyssey Infrared Imaging System (LI-COR Biosciences).

ChIP assay. ChIP is a method to determine the identity and quantity of DNAs that bind particular proteins of interest. Following the breakage of DNA molecules into pieces by sonication, the protein–DNA complex can be immunoprecipitated out from the cellular lysate by using an antibody that is specific to a putative DNA–binding protein. Then the identity and quantity of DNA bound by the protein can be analyzed using PCR. After optimization of DNA-sheeting conditions, the manufacturer’s standard protocol for ChIP was followed. About 1 × 10^7 C2C12 cells in a 10-cm dish were treated with 1% formaldehyde for 10 min at 37 °C. Cells were washed twice with cold PBS containing protease inhibitors and suspended in 200 μl of SDS lysis buffer in a 1.5 ml eppendorf tube, which was then held in ice water. Sonication was applied to the cell pellets for a total of 18 min using a 130-W ultrasonic processor (model: VCX 130) accessorized with a 6-mm probe (model: CV18) (VWR International Ltd, Hunter Boulevard, England) with the following setting: amplitude setting 15% using 15 s pulses with 30 s cool-down intervals between pulses. About 20 μl of sonicated cell supernatant was kept as input samples and the rest was diluted by 10-fold. After immunoprecipitation with anti-HEK and anti-MyoD antibodies, 60 μl of protein A-agarose/salmon sperm DNA was added for 1 h of culture, followed by centrifugation. The complex was eluted and histone–DNA crosslinks were reversed by heating at 65 °C for 4 h. DNA was recovered by phenol/chloroform extraction. Finally, 5 μl of DNA suspension was used for PCR cycling.

In situ PLA. Cells were seeded on 35-mm glass-bottom dishes (MatTek Corp., Ashland, MA, USA). After treatment, cells were fixed with cold 4% paraformaldehyde for 15 min at room temperature followed by cell-membrane permeabilization with 0.25% Triton-X100 in PBS for 10 min. The cells were washed three times with PBS, blocked for 1 h at room temperature with 1% BSA in PBS containing 0.1% Tween-20 and incubated with the indicated antibody pairs in a cell incubator at 37 °C. Duolink in situ PLA (Olink Bioscience, Uppsala, Sweden) was performed according to the manufacturer’s instructions. Images were taken using an Olympus Research Inverted System Microscope (Olympus, Center Valley, PA, USA) with X81 LUCPlan FLN objectives and cellSen Dimension software (1.4.1; Olympus). The paired primary antibodies used included rabbit and mouse BMP4 antibody, rabbit BMPRA1 antibody and rabbit and mouse BMPR2 antibody. For preparation of IgG-conjugated MWCNT 1, mouse IgG (100 μg, Santa Cruz Biotechnology, Inc.) or BSA (100 μg) was mixed with MWCNT 1 (50 μl of the supernatant of aqueous MWCNT 1 suspension after being centrifuged at 10 000 × g for 1 min) in water (final volume, 400 μl) and shaken overnight at 4 °C.

The mixture was centrifuged at 16 000 × g for 30 min and the pellet was resuspended in 100 μl of water. Finally, 50 μg/ml-conjugated MWCNT 1 was used for cell treatment (2 h).

FRET. FRET was performed using a Zeiss LSM 510 Meta microscope with a × 63 objective lens, all controlled by Zen 2009 software (Peabody, MA, USA). For labeling MWCNT 1, 200 μl of the supernatant of aqueous MWCNT 1 suspension (0.5 mg/ml, centrifuged at 10 000 × g for 1 min) was mixed with 4.2 mg of FITC-BSA (Sigma, St. Louis, MO, USA) in 0.6 ml of ddH2O and shaken overnight at 4 °C. The mixture was centrifuged at 16 000 × g for 30 min to pellet MWCNT 1, and the pellet was washed once with PBS. Finally, the pellet was suspended in 0.2 ml of ddH2O. C2C12 cells were seeded on 35-mm glass-bottom dishes (MatTek Corp.). After incubation with FITC-conjugated MWCNT 1 (50 μg/ml) for 2 h, cells were washed with PBS twice to remove free MWCNT 1. Samples were prepared as for immunocytochemistry. FRET was detected by an acceptor photobleaching method. This method involves photoquenching in the presence of an acceptor. This can be done by comparing donor fluorescence intensity in the same sample before and after destroying the acceptor by photobleaching. If FRET was initially present, a resultant increase in donor fluorescence will occur after photobleaching of the acceptor. A selected region of interest was irradiated with the 543-nm laser line (70% bleach power, 50 iterations) for 12 s to photobleach TRITC, FITC and TRITC images were collected immediately after photobleaching, and FRET was measured as an increase in FITC fluorescence intensity. The FRET efficiency was calculated as \( FRET_{\text{eff}} = 1 - \frac{D_{\text{post}}}{D_{\text{pre}}} \) using the Zen 2008 software, taking into account background noise in each channel, where \( D_{\text{pre}} \) and \( D_{\text{post}} \) are, respectively, the even FITC fluorescence intensity in the selected region before and after photobleaching.

Cell transfection and luciferase activity assay. C2C12 or C33A-2D2 cells were seeded in 24-well plates. After overnight culture, the cell confluence reached about 80–90%. Next, 0.6 μg of plasmids mixed with Lipofectamine 2000 at a ratio 1:3 (w/w) in 100 μl of Opti-MEM I Reduced-Serum Medium was used for transfecting cells in one well. A pcDNA3.1 empty vector was used as control for caALK2 and caALK3 constructs and a pGL3 empty vector was used as control for myogenin–luciferase and p2l-luciferase constructs. A pGL3 control vector containing SV40 promoter and enhancer sequences was used to monitor the transfection efficiency simultaneously. Six hours after transfection, cells were changed to fresh medium, and 24 h after transfection, cells were exposed to CNTs. After the indicated time, cell lysates were prepared by adding 80 μl of 1 × RIPA buffer to each well. Protein concentration was detected with a BCA Protein Assay Kit (Thermo Scientific, Barrington, IL, USA). A luciferase activity assay was conducted in a white 384-well plate and detected with an Envision 2012 Multilabel Reader (PerkinElmer, Waltham, MA, USA). Finally, 4 μl of the lysates and 25 μl of SteadyLite Plus substrate were used for a one-well assay. The luciferase activity was normalized as luminescence intensity divided by the protein concentration of the same lysate sample.

Id1 promoter activity assay. Approximately 20 000 C33A-2D2 cells per well were seeded in white 96-well plates and cultured overnight. Each CNT (100 μg/ml) from the combinatorial array was incubated with cells for 24 h before BMP4 (25 ng/ml) was added. After incubation for another 24 h, 100 μl of SteadyLite Plus substrate was added and luminescence intensity was determined. To normalize luciferase activity, a cell viability assay with the same CNT exposure was performed with the XTT method. The luminescence intensity divided by the number of viable cells in each well was used and the value in medium was set as 1.

Statistical analysis. The two-sided student’s t-test was used to analyze differences between experiments. Data are reported as mean values ± S.D. from multiple determinations. P-values < 0.05 were considered statistically significant and all the statistical calculations were carried out using SigmaPlot 10.0 (Systat Software, Chicago, IL, USA).

Conflict of Interest

The authors declare no conflict of interest.

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