Methyltransferase-like protein 7A (METTL7A) promotes cell survival and osteogenic differentiation under metabolic stress

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INTRODUCTION

Bone can heal itself, but it is exceedingly difficult to reconstitute large bone defects induced by heavy trauma or resection of malignant tumor. In addition, if blood supply to bone defect area is deficient, bone regeneration is seriously disturbed, exemplified in osteonecrosis of the femoral head. Regenerative medicine, including the implantation of stem cells, has been studied as a novel solution to treat these conditions. However, when the local vascularity is impaired, even the transplanted cells undergo rapid necrosis before differentiating into osteoblasts and regenerating bone. Thus, to increase the effectiveness of stem cell transplantation, it is quintessential to improve the viability of the implanted stem cells. In this study, given that the regulation of glucose may hold the key to stem cell survival and osteogenic differentiation, we investigated the molecules that can replace the effect of glucose under ischemic microenvironment of stem cell transplantation in large bone defects. By analyzing differentially expressed genes under glucose-supplemented and glucose-free conditions, we explored markers such as methyltransferase-like protein 7A (METTL7A) that are potentially related to cell survival and osteogenic differentiation. Overexpression of METTL7A gene enhanced the osteogenic differentiation and viability of human bone marrow stem cells (hBMSCs) in glucose-free conditions. When the in vivo effectiveness of METTL7A-transfected cells in bone regeneration was explored in a rat model of critical-size segmental long-bone defect, METTL7A-transfected hBMSCs showed significantly better regenerative potential than the control vector-transfected hBMSCs. DNA methylation profiles showed a large difference in methylation status of genes related to osteogenesis and cell survival between hBMSCs cultured in glucose-supplemented condition and those cultured in glucose-free condition. Interestingly, METTL7A overexpression altered the methylation status of related genes to favor osteogenic differentiation and cell survival. In conclusion, it is suggested that a novel factor METTL7A enhances osteogenic differentiation and viability of hBMSCs by regulating the methylation status of genes related to osteogenesis or survival.

RESULTS

Glucose is essential for osteogenic differentiation and survival of hBMSCs

In this study, assuming that the regulation of glucose may be the key to stem cell survival and osteogenic differentiation, we aimed to find the molecules that potentially replace the effect of glucose in ischemic microenvironment of stem cell transplantation for large bone defects [7, 10]. By analyzing differentially expressed genes under glucose-free conditions, which generate microenvironmental stress, and under glucose-supplemented conditions, the genetic markers that are possibly related to cell survival and osteogenic differentiation were explored. We mined methyltransferase such as 7A (METTL7A) gene, which codes for a putative enzyme that induces methylation [11–13], resulting in enhanced viability and osteogenesis of hBMSCs. The in vitro and in vivo effects of METTL7A in both glucose-free and glucose-supplemented environments as well as the underlying mechanism of action were investigated.
osteo-induced hBMSCs were cultured under osteogenic medium supplemented with 5.5 mM glucose [G(+)OM] or under osteogenic medium without glucose [G(−)OM] for 21 days. Normally, normal blood glucose levels are defined as 4–5 mM and low glucose concentrations below 2.2 mM. Several studies also set 0–1 mM glucose as the starting level of low glucose condition [14–17]. The glucose supplementation conditions were set to the normal range of 5.5 mM and the glucose-free condition was set to 0 mM. Osteogenic differentiation was significantly enhanced in G(+)OM than in G(−)OM while almost no staining was detected under G(+)CM (Fig. 1A). When the intensity of Alizarin Red staining was analyzed graphically, the intensity was >20-fold greater in G(+)OM than in G(−)OM on day 21 of culture (Fig. 1B). The gene expression of osteogenic marker type I collagen (COL1) also significantly greater in G(+)OM than in G(−)OM while that of osteocalcin (OCN) was not on day 7 of culture (Fig. 1C). Cell viability also significantly decreased by more than 70% from day 7 with G(−)OM compared with G(+), while no significant difference in cell viability was found between G(+)-CM and G(+)OM (Fig. 1D). Western blot showed an increase in apoptosis-related protein, BAX, and a decrease in the survival-related gene, BCL-XL, under glucose-free condition. p-AKT (s473), which is activated by metabolic process, stem cell proliferation, endodermal cell differentiation, and osteoblast development between the two conditions, the METTL7A-transfected hBMSCs showed even more enhanced categories (Fig. 2C). Based on the literature review, we selected 30 of those genes associated with cell survival or differentiation, and directly tested their gene expression in hBMSCs, and that the inhibition of METTL7A by shMETTL7A reduced osteogenic differentiation and cell viability of hBMSCs in glucose-supplemented osteogenic medium.

**METTL7A overexpression enhance osteogenic differentiation and viability of hBMSCs in glucose-free conditions**

To demonstrate the effects of METTL7A on osteogenic differentiation and cell survival of hBMSCs, the METTL7A plasmid vector was constructed. Figure 4A shows the overall scheme for the generation of METTL7A-transfected hBMSCs (METTL7A). Since the region containing bacterial skeleton is naturally degraded and disintegrated during the proliferation process, a minicircle vector containing only introduced genes was obtained as the final product. The RFP gene was also inserted into plasmids to monitor the transfection efficiency. The minicircle plasmid carrying the RFP gene was used as the control vector. The generated minicircle plasmid vector was then introduced into hBMSCs via electroporation (Fig. 4A). The fluorescent image obtained 1 day after electroporation demonstrated successful transfection of the plasmid vector as confirmed by RFP signal in both METTL7A-transfected (METTL7A) and control vector-transfected hBMSCs (MiniCircle; MC) (Fig. 4B). Western blotting confirmed the increased expression of METTL7A gene and protein in METTL7A-transfected hBMSCs (Fig. 4C). METTL7A transfection significantly enhanced the viability of hBMSCs in glucose-free condition throughout the culture period (Fig. 4D). As a next step, we tested whether METTL7A overexpression reversed the reduced osteogenic differentiation and cell viability of hBMSCs cultured under glucose-free conditions. Alizarin Red staining showed significantly enhanced formation of calcified matrix in METTL7A compared with MiniCircle after 7 days of culture in glucose-free conditions (Fig. 4E, F and S4B). Addition of human recombinant METTL7A protein led to comparable enhancement of calcified matrix formation in treated hBMSCs after 7 days of culture in glucose-free condition (Fig. S1). Also, under glucose-supplemented conditions, the METTL7A-transfected hBMSCs showed even stronger Alizarin red staining than the control vector-transfected hBMSCs after 14 days of culture (Fig. S2).
METTL7A-transfected hBMSCs enhance new bone formation in vivo in a critical-size segmental bone defect model

The in vivo effectiveness of METTL7A-transfected hBMSCs in bone regeneration was investigated in a rat model of critical-size segmental long-bone defects. 106 METTL7A-transfected hBMSCs were implanted along with fibrin glue into 4 mm-sized segmental defects created in the radii of immunosuppressed rats (Figs. 4A and 5A; schematic diagram of the experiment process). The healing of bone defects was assessed radiographically on days 28 and 56, and via on day 56 after implantation (Fig. 5B). Bone regeneration was
significantly better when implanted with METTL7A-transfected hBMSCs (METTL7A; MC-M7 group) compared with control vector-transfected hBMSCs (MiniCircle; MC group), or without cell implantation (control group) (Fig. 5B). In segmental long-bone defect model, the METTL7A group showed substantially better quality of bone regeneration, and complete healing of the segmental radial defect, while other groups had insufficient healing or gross nonunion of critical-size defects, demonstrated radiographically (Fig. 5B) and via

Fig. 2 Identification of the novel gene associated with enhanced survival and osteogenic differentiation in hBMSCs. A RNA-seq data analysis of hBMSCs. Analysis of hierarchical clusters of genes significantly related to each sample via RNA-seq data analysis. A heatmap was generated to visualize transcriptomic differences among undifferentiated hBMSCs cultured in control medium with 5.5 mM glucose [G(+)CM], osteo-induced hBMSCs cultured in osteogenic medium containing 5.5 mM glucose [G(+)OM] or no glucose [G(−)OM] (fold-change > 2, \( p < 0.05 \)). Red indicates upregulation, while blue indicates downregulation. B Pathway analysis in Functional Annotation for significant probe list was performed using DAVID for G(−)OM and G(+)OM. An analysis of difference in signaling pathways related to metabolic process, stem cell proliferation, cell differentiation, and osteoblast development. C A selected heatmap was generated to visualize relative transcriptomic differences among transcripts from G(+)CM, G(+)OM, and G(−)OM (fold-change > 2, \( p < 0.05 \)). D Relative gene expression of METTL7A was determined by RT-qPCR, and E Western blot. Data were presented as mean ± standard deviation (SD, \( N = 3 \)). NS: no significant difference, *\( p < 0.05 \), **\( p < 0.01 \). G(+)CM: undifferentiated hBMSCs cultured in control medium with 5.5 mM glucose; G(+)OM: osteo-induced hBMSCs cultured in osteogenic medium containing 5.5 mM glucose; G(−)OM: osteo-induced hBMSCs cultured in osteogenic medium without glucose.
MicroCT imaging (Fig. 5C). The bone mass (BV, mm³), BV/TV (bone mass/total volume, %) and bone mineral density (BMD) were evaluated by MicroCT. BV and BV/TV of METTL7A group was about 2-fold higher than that of the MC group. BMD of METTL7A group was similar to that of normal bone, and also significantly greater than that of MC group (Fig. 5D). Histological findings also demonstrated that the METTL7A group had much better quality of bone regeneration than in MC group or control group (Fig. 5E). These results indicated that METTL7A overexpression enhances the in vivo regenerative potential of hBMSCs in bone healing.

DNA methylation profiles differed significantly in hBMSCs depending on glucose availability

DNA methylation profiles of G(+)CM, G(+)OM, and G(−)OM were investigated using deepTools 3.3.0. Dividing the "gene body length" by the size of "bin" confirms the number of bins to calculate the per-base value of each region. The average of bins for each region was determined and the heatmap was created by stacking them vertically. The heat maps are represented by transcription start site (TSS) and transcription end site (TES). The corresponding formula for each of the three comparative pairs was determined according to the scatter plot of the β-value of DNA methylation sites in the three pairs of cells. Curve fitting showed consistent DNA methylation profiles of hBMSCs under the three conditions. The methylation level in general was significantly higher in G(+)-OM than in G(-)-OM (Fig. 6A). A osteogenesis-related gene peak heatmap using MeV 4.9.0 was generated to visualize genes with relative differences in methylation between G(+)OM and G(−)-OM, which showed significant differences in promotor methylation status of osteogenesis-related genes. Methylation of

Fig. 3 The effect of METTL7A knockdown on cell viability and osteogenic differentiation of hBMSC. A Construction of shMETTL7A and scramble control vector. Each vector contains RFP fluorescent genes. B Fluorescent, bright-field and merge images of scramble control vector- and shMETTL7A-transfected hBMSCs (RFP-tagged) after 1 day of transfection. C Western blot of METTL7A in scramble control vector- (shControl) and shMETTL7A-transfected hBMSCs (shMETTL7A). D Alizarin Red staining images and E quantification of shControl and shMETTL7A in osteogenic medium containing 5.5 mM glucose after 14 days. F Cell viability of shControl and shMETTL7A cultured under osteogenic medium containing 5.5 mM glucose on days 7, 14, and 21. Data were presented as mean ± standard deviation (SD, N = 3). NS: no significant difference, *p < 0.05, **p < 0.01. shControl: control vector-transfected hBMSCs; shMETTL7A: shMETTL7A-transfected hBMSCs.
Fig. 4 Cell viability and osteogenic differentiation of METTL7A gene-transfected hBMSCs. A Construction of METTL7A minicircle plasmid vector. Vector also contains RFP fluorescent genes. B Fluorescent, bright-field and merge images of minicircle control vector (MiniCircle)- or METTL7A-transfected hBMSCs (METTL7A; RFP-tagged). C Western blots of METTL7A in hBMSCs. D Cell viability of hBMSCs cultured in glucose-free osteogenic medium on days 7, 14, and 21. E Alizarin Red staining images and F quantification of hBMSCs cultured in glucose-free osteogenic medium after 7 days of culture. Data were presented as mean ± standard deviation (SD, N = 3). NS: no significant difference, *p < 0.05, **p < 0.01. hBMSCs: untransfected hBMSCs control; MC: minicircle plasmid vector (pMC)-transfected hBMSCs control; METTL7A: hMETTL7A- transfected hBMSCs.
**Fig. 5 In vivo new bone formation in segmental bone defect models.**

A. Creation of critical-size segmental long-bone defect in immunosuppressed rat. Length of the defects was 4 mm. hBMSCs were implanted with fibrin glue into segmental defects of rats created in the radius of right forelimb. B. Radiographic images of segmental defects on days 28 and 56 after the implantation of hBMSCs. C. MicroCT images of segmental long-bone defects at 56 days after cell implantation. D. After 56 days, bone volume (BV, mm³), BV/TV (bone volume/total volume, %), and BMD (bone mineral density, mg/cc) were evaluated by MicroCT. E. Goldner's trichrome staining of healed segmental defects on day 56 post implantation of hBMSCs. Statistical processing was performed in comparison with the control group. Data were presented as mean ±standard deviation (SD, N = 4). NS: no significant difference, *p < 0.05, **p < 0.01. Normal: no surgery and no cell implantation; control: surgery only without cell implantation; MiniCircle (MC): minicircle control vector-transfected hBMSCs implantation; MC-METTL7A (MC-M7): METTL7A-transfected hBMSCs implantation.
promotor regions in these genes was generally greater in G(−)OM than in G(+)-OM (Fig. 6B). Figure 6C is a heatmap representing the promotor methylation peaks of specific genes selected from hBMSCs (BMP2, METTL7A, Runx2, and BCL2). BMP2 and Runx2 are associated with osteogenic differentiation while BCL2 gene is related to anti-apoptosis. The methylation peaks of those genes varied strongly between the two conditions (Fig. 6C). Gene peak heatmap and graph using MeV 4.9.0 was generated to visualize the relative differences in promotor methylation between METTL7A-transfected hBMSCs and control vector-transfected
hBMSCs. After 3 days of transfection, there was a significant difference in methylation between the two types of cells (Fig. 6D and E). Methylation of osteogenesis-differentiation-related genes (Runx2, SOX8, COL6A1, and AKT) was greater in control vector-transfected hBMSCs than in METTL7A-transfected hBMSCs. The methylation status of apoptosis-related genes significantly differed between the two types of cells. Methylation in the promoter regions of genes related to apoptosis was greater in METTL7A-transfected hBMSCs than in control vector-transfected hBMSCs (Fig. 6F). As a result, methylation between the conditions with or without glucose at the time of osteogenic differentiation shows a marked difference, and the METTL7A gene appears to regulate this methylation. Figure 7 is a schematic diagram illustrating a new relationship between methylation and a gene that induces bone regeneration in METTL7A transfected bone marrow stem cells. The METTL7A gene is shown to induce bone regeneration by regulating the methylation of genes involved in osteogenic differentiation and cell survival.

**DISCUSSION**

While most bone defects heal by themselves or aided by conventional methods including autologous bone grafting, allografts, or bone substitutes, such lesions as large long-bone defects and osteonecrosis of femoral head are not amenable to such interventions. Therefore, regenerative medicine for bone was mainly dedicated to the possible treatment of these recalcitrant conditions. The application of stem cells, most commonly the use of mesenchymal stem cells (hBMSCs), has emerged as a potentially game-changing therapy for several medical conditions that have been refractory to conventional interventions. However, contrary to original expectations, most healing effects of implanted stem cells are attributed to paracrine effects. Implanted hBMSCs largely undergo massive cell death, failing to engraf and differentiate into host tissue [1, 2]. The massive death of grafted cells upon transplantation in the ischemic site is caused by metabolic stress due not only to hypoxia but also because of reduced supply of critical metabolic nutrients and inadequate metabolic waste removal [18].

Glucose is a primary component of metabolic homeostasis, and is a major energy source used for the synthesis of DNA, RNA, proteins, and lipids [4, 19–21]. Upon transport into the cell via the Glut family of transporters, a glucose molecule is metabolized in the cytoplasm via glycolysis to generate two pyruvate molecules, 2 ATP, and 2 reducing equivalents in the form of nicotinamide adenine dinucleotide (NAD). In hBMSCs survival, glucose is rapidly utilized or depleted, whereas amino acids and other required nutrients are used sparingly. Serum starvation or nutrient depletion appears to have a less notable effect on cells than glucose does [18, 20]. While the role of oxygen is pivotal to hBMSCs survival, several studies showed that hBMSCs endure sustained near-anoxia condition in the presence of exogenous glucose. Protein expression of Hif-1a and angiogenic factors was upregulated by glucose. Ectopically implanted tissue constructs supplemented with glucose exhibited 4- to 5-fold higher viability and vascularization compared with those without glucose [14]. hBMSCs survive exposure to long-term (12 days) and severe (PO2 < 1.5 mmHg) hypoxia in the presence of glucose. hBMSCs remained functionally viable after exposure to long-term, severe hypoxia under glucose supplementation [2]. Under low-glucose conditions, cells initiate adaptation followed by apoptosis responses using PERK/AKT and MEK1/ERK2 signaling, respectively [19]. Normally, normal blood glucose levels are defined as 4–5 mM and low glucose concentrations below 2.2 mM. Several studies also set 0–1 mM glucose as the starting level of low glucose condition [14–17]. We set the glucose-supplemented condition as the normal range of 5.5 mM and the glucose-free condition as 0 mM. In this study, similarly, the pAKT (S473) level decreased and p-ERK increased in glucose-free medium, reflecting apoptotic conditions due to metabolic stress.

While the signaling pathways involving Runx2, Wnts, and BMPs have been extensively studied in osteogenic differentiation of progenitor cells [22], the role of metabolic environment in bone regeneration has yet to be investigated. Glucose uptake induces osteoblast differentiation by suppressing the AMPK-dependent proteosomal degradation of Runx2 and promotes bone formation by inhibiting another function of AMPK. Runx2 also favors Glut1 expression, and this feedforward regulation between Runx2 and Glut1 determines the onset of osteoblast differentiation during development and the extent of bone formation throughout life [9]. In the first step of current study, we reconfirmed that glucose was essential for survival and osteogenic differentiation of hBMSCs under the osteogenic medium. Although the medium contains other nutrients including serum, a markedly reduced cell viability as well as increased expression of apoptosis-related proteins (BAX and p-ERK) and decreased expression of survival-related proteins (BCL-XL and p-AKT) suggested the need for glucose in the cell survival. Also, glucose-free conditions led to an extremely low expression of osteogenic marker proteins (Runx2, BSP) and scanty calcified matrix.

Next, we explored molecules that may replace the role of glucose in enhancing the survival and osteogenic differentiation of hBMSCs by DEG analysis. Among the 48 genes that may be differentially expressed in hBMSCs with or without glucose, METTL7A was identified as a novel factor that showed significant difference in expression in response to glucose and the most prominent osteogenic effect in the absence of glucose. METTL7A, also known as AAM-B, was initially identified as a lipid droplet-associated protein in Chinese hamster ovary K2 cells via proteomic analysis [23]. It was reported as an integral membrane protein anchored into the endoplasmic reticulum membrane that recruit cellular proteins for lipid droplet formation. Interestingly, the intermediate region of METTL7A plays a putative role as S-adenosyl methionine-dependent methyltransferase [12, 24]. The methylation level of METTL7A gene was downregulated in thyroid cancer compared to normal thyroid cells [13]. It is also known to be a tumor suppressor gene with multiple editing sites at its 3′UTR [25]. However, the role of METTL7A in cell survival or osteogenic differentiation and its regulation by glucose have yet to be reported.

We tested the function of METTL7A in cell survival and osteogenesis of hBMSCs in two ways: (1) suppression of METTL7A
gene expression by shRNA reduces the effect of glucose supplementation; and (2) METTL7A gene overexpression mitigates the problem of glucose deficiency. The in vitro results confirmed that METTL7A was necessary for cell survival and osteogenesis of hBMSCs, and that METTL7A overexpression can partially replace the effect of glucose. Also, the implantation of METTL7A-transfected hBMSCs showed significantly greater in vivo osteogenic potential in critical size long-bone defects in immunocompromised rats. These results suggested that METTL7A mediates the effect of glucose in cell survival and osteogenic induction and can be used to promote stem cell-based bone regeneration.

As METTL7A acts as a methyltransferase, we finally investigated if the osteogenic induction and availability of glucose affected the methylation status of promotor regions in genome and also if METTL7A transfection altered the epigenetic status of hBMSCs. DNA methylation patterns have increasingly been implicated in transcriptional regulation and efficiency, due to advances in genome-wide DNA methylation profiling studies [26]. Current advances in transcriptional regulation by DNA methylation mostly focus on the promoter region where hypomethylated CpG islands are present with transcriptional activity, as hypermethylated CpG islands generally result in gene repression [27–29]. Environmental factors induce specific phenotypic changes in genomic DNA through epigenetic modification [30]. Although DNA methylation is known to be the most important epigenetic regulator of mammalian development [31], studies involving altered methylation status due to metabolic stress in osteogenic differentiation of progenitor cells have yet to be reported.

Our results from DNA methylation profiling of promotor regions suggested large differences in methylation status of osteogenesis (BMP2, Runx2) or survival (BCL2)-related genes in hBMSCs depending on glucose availability. METTL7A transfection appears to decrease the methylation of osteoblast differentiation-related genes while increasing that of apoptosis-related genes. Therefore, it is possible that glucose supplementation increased intracellular expression of METTL7A, which in turn leads to altered methylation status of related genes to favor osteogenic differentiation and cell survival. To the best of our knowledge, this is the first reported evidence supporting the role of METTL7A in cell survival and osteogenic differentiation by changing the promoter methylation status of related genes. However, it is not known from the results of this study and awaits further investigation how the increased glucose level leads to increased METTL7A gene expression and how METTL7A differentially methylate certain gene promoters.

MATERIALS AND METHODS

Cell culture

Human bone marrow stem cells (hBMSCs) were isolated from bone marrow of seven patients (mean age: 74 years; range: 60–87 years). Informed consent was obtained from all donors. All experiments were performed in accordance with the relevant guidelines and regulations. The collection of human samples was approved by the Institutional Review Board at Dongguk University Ilsan Hospital (IRB file no. DUIH 2012-01-034). Bone marrow obtained from the human body was diluted with Dulbecco’s phosphate-buffered saline (DPBS: Welgene, Cat. LB 001-02, Republic of Korea) and transferred to a conical tube, supplemented with Lymphoprep (Alere Technologies AS, Cat. 1114544, Norway) in a 1:1.25 ratio. The oil layer was removed by centrifuging at 2000 rpm, brake 0, 30 min, and the cell layer was diluted 1:2 with DPBS after collecting separately in a new tube. The hBMSCs pellet was obtained by centrifuging at 1500 rpm for 8 min from diluted cell layer. The hBMSCs were incubated at 1 × 10^5 cells per 100 mm dish in αMEM (Gibco, Cat. 12571-063, Grand Island, NY, USA) supplemented with 10% FBS (Gibco, Cat. 16000-044, Grand Island, NY, USA) and 1% penicillin–streptomycin 100X (Welgene, Cat. LS 202-02, Republic of Korea) at 37 °C under 5% CO2. It was tested for mycoplasma contamination and was confirmed to be free from contamination.

To further investigate osteogenic differentiation, 2.5 × 10^5 cells were placed in each well of a 6-well plate. Osteogenic differentiation was induced by replacing the osteogenic medium (OM: DMEM with 10% FBS,
The supernatant was removed by centrifuging hBMSCs at 1000 rpm for 1% penicillin–streptomycin, 10 mM glycerolphosphate, 50 mM L-ascorbic acid-2-phosphate, 100 nM dexamethasone) at 37 °C under an atmosphere of 5% CO2, DMEM with low glucose (Welgene, Cat. LM 001-06, Republic of Korea) was used to make 5.5 mM glucose-supplemented osteogenic medium, and DMEM without glucose (Gibco, Cat. 11966-025, USA) to make glucose-free osteogenic medium (OM). The maintenance medium for hBMSCs also contained 5.5 mM glucose, and was used as control medium (CM), designated as glucose-supplemented control medium.

### Construction of MiniCircle Vector
pMC-CMV-MCS (MNS01A-1), a vector to produce MiniCircle (MC), was purchased from Systemic Biosciences (SBI, Palo Alto, CA, USA) and used as a basic framework to prepare the plasmid vector. Sequences capable of expressing METTL7A genes were inserted downstream of the CMV promoter using the BamHI and NheI restriction sites of the parent plasmid pCMC (pCMC vectors). The METTL7A Human shRNA Plasmid Kit (Origene, Cat. TF311498, MD, USA) was used. The pRFP-C-RS Vector (Origene, Cat. TR30014, MD, USA) was used. The pMC control vector-transfected hBMSCs were cultured in α-MEM (Gibco, Cat. 12571-063, Grand Island, NY, USA) without antibiotics. Transfection efficiency was confirmed by counting the fluorescent cells.

### Preparation and transfection of shRNA vector
The METTL7A Human shRNA Plasmid Kit (Origene, Cat. TF311498, MD, USA) and pRFP-C-RS Vector (Origene, Cat. TR30014, MD, USA) was used. The plasmid contains the sequence that inhibits METTL7A mRNA and the restriction enzyme sites that are responsive to chloramphenicol and puromycin. To confirm the transfection efficiency, the RFP gene was inserted into the vector. The pRFP-C-RS shRNA vector was used as the control. Transfection was performed using 2.5 µg of TurboFect transfection reagent (Origene, Cat. TF81001, MD, USA) mixed with 2 x 10⁵ cells according to the manufacturer’s instructions. Transfected cells were cultured for 2 days at 37 °C under an atmosphere of 5% CO2 and selected with 2 µg/mL puromycin for 3 days. The transfection efficiency was confirmed through fluorescence expression.

### RT-qPCR analysis
Total RNA was isolated from hBMSCs using Direct-zol™ RNA MiniPrep Plus (200 preps, Zymo Research, Cat. R2072, Irwin, CA, USA). The cDNA was synthesized using RT-qPCR with Maxime RT PreMix (Oligo dt primer: iNTRON, Cat. 25081, Republic of Korea) in a SimpliAmp Thermal Cycler (Applied Biosystems, Foster City, CA, USA). The primers used are shown in Table 1.

### Alizarin Red staining
To evaluate the osteogenic differentiation of hBMSCs from the calcified matrix, cells were stained with Alizarin Red S (Sigma-Aldrich, Cat. A5533-

### 25G, Saint Louis, MO, USA). After osteogenic differentiation for 7 days, the cells were fixed with 4% paraformaldehyde, then stained with 2% Alizarin Red S and dried. Alizarin Red staining area (%) was measured using the ImageJ program (NIH).

### Cell viability analysis
Cell viability analysis is used to determine the degree to which tetrazoil salt is reduced by mitochondrial NADH dehydrogenase in cells and converted into a colored formazan. In this assay, 1 x 10⁵ hBMSCs were dispensed to each well of 96-well plate, suspended in 100 µL of each culture medium, and incubated at 37 °C under an atmosphere of 5% CO2 for 24 h. Analysis was performed using the EZ-Cytox kit (DoGEN, Cat. EZ-1000, Republic of Korea) as described by the manufacturer’s instructions. The reagent contained in the above kit was added to the medium in the ratio of 1:10 in micro-level quantities. After 1 h, the optical density was measured at 450 nm wavelength with a VersaMax Microplate Reader ( Molecular Devices, Sunnyvale, CA, USA).

### Western blotting analysis
Cells were collected in a radioimmunoprecipitation assay buffer (Thermo Scientific, Cat. 89900, Waltham, MA, USA) containing Halt™ Protease & Phosphatase Inhibitor Cocktail (Thermo Scientific, Waltham, MA, USA). Protein concentration was measured using a bichinchoninic acid assay, and the dissolved proteins were separated by SDS-polyacrylamide gel electrophoresis for 2 h, and transferred to a nitrocellulose membrane (Whatman™, Cat. E06-07-111, UK), followed by blocking with 5% skimmed milk in 1× Tris-buffered saline with 2% Tween-20. Thereafter, the cells were reacted overnight at 4 °C with the primary antibodies (1:1000, Table 2), followed by the secondary antibody (1:1000) (anti-rabbit IgG horseradish peroxidase (HRP)-linked antibody, #7074, Cell Signaling™/anti-mouse IgG HRP-linked antibody, #7076, Cell Signaling™) at room temperature (RT) for 2 h. Antibody-antigen complexes were detected using SuperSignal™ West Femto Maximum Sensitivity Substrate (Thermo Scientific, Cat. 34095, Waltham, MA, USA), and signal intensities were measured using the ChemiDoc™ XR+ Imaging System (Bio-Rad, Hercules, CA, USA).

### Critical-sized segmental long-bone defect in rats
An 11-week-old male Sprague-Dawley (SD) rat was used in the bone defect model. Five animals were assigned to each of the following four groups randomly: normal (no surgery, no cell implantation), control (defects only without cell implantation), MiniCircle (pMC control vector-transfected hBMSCs implantation), and MC-METTL7A (METTL7A-transfected hBMSCs implantation). All animal management procedures, anesthesia, and surgeries were performed in compliance with the ARRIVE guidelines and the protocol of Dongguk University Institutional Animal Care and Use Committee. Segmental long-bone defects were created by drilling to a depth of 4 mm in the proximal radius of SD rats. 1 x 10⁷ METTL7A-transfected hBMSC or control vector-transfected hBMSCs mixed with 20 µL fibrin/thrombin (Green Cross, Republic of Korea) were implanted into the segmental long-bone defects. The rats were injected with immunosuppresive agents (CIPOL; Chong Kun Dang, Republic of Korea) daily, 2.5 mg/300 µL in the first week and 0.83 mg/100 µL from 2 to 8 weeks. After 8 weeks, bone regeneration was confirmed by visual observation, X-ray, and MicroCT. Bone volume (BV, mm³), BV/TV (bone volume/total volume, %), and BMD (bone mineral density, mg/cc) were evaluated by MicroCT.

### GoldeR’s triamcine staining
The segmental bone tissue was frozen into blocks using Tissue-Tek O.C.T. Compound (Sakura Finetek, 4583, Torrance, CA, USA). A freeze-cutting machine (Leica CM1950 Cryostat; Leica Microsystems, Wetzlar, Germany), tissue blocks were sectioned to 20 µm in thickness. Tissue slices were washed with distilled water and treated with a 1:1 mixture of hematoxylin solution II (Merck Millipore, Cat. HX384856, Billerica, MA, USA) and III (Merck Millipore, Cat. HX303261, Billerica, MA, USA). After washing with flowing water, GoldeR I, II, and III solutions (Carl Roth, Cat. Art-Nr. 3469.1, 3470.1, 3473.1, Karlsruhe, Germany) were used sequentially for 5 min each. After washing with distilled water, the stained tissue slice was mounted, and observed under a microscope.

### Glucose uptake assay
Glucose uptake in living cells was observed using 2-NBDG. hBMSCs were seeded at 1.25 x 10⁵ cells per well in 12-well plates and cultured in
Table 2. Primary antibodies used in western blotting analysis.

| Target    | Antibody                           | Company       | Catalog No. |
|-----------|------------------------------------|---------------|-------------|
| METTL7A   | Anti-METTL7A antibody              | Abcam         | Ab79207     |
| BSA       | Anti-Bone Sialoprotein antibody     | Abcam         | Ab30322     |
| Runx2     | Mouse-Runx2 antibody               | Abcam         | Ab76956     |
| BMP2      | Anti-BMP2 antibody                 | Abcam         | Ab14933     |
| Glut1     | Glut1 Antibody (A-4)               | SCBT          | sc-377228   |
| Glut4     | Anti-Glut4 Transporter GLUT4 antibody | Abcam         | Ab216661    |
| LDHA      | LDHA (C485) Rabbit mAb             | Cell Signaling| C485        |
| BAX       | Bax Antibody                       | Cell Signaling| #27725      |
| BCL-XL    | Bcl-xL (54H6) Rabbit mAb           | Cell Signaling| #27645      |
| p-AKT(473)| Phospho-Akt (Ser473) (D9E) XP® Rabbit mAb | Cell Signaling| #40605      |
| p-AKT(380)| Phospho-Akt (Thr380) (D25E6) XP® Rabbit mAb | Cell Signaling| #130385     |
| AKT       | Akt (pan) (C67E7) Rabbit mAb       | Cell Signaling| #46915      |
| p-ERK     | Phospho-p44/42 MAPK (Erk1/2) (Thr202/Tyr204) Antibody | Cell Signaling| #91015      |
| ERK       | p44/42 MAPK (Erk1/2) (137F5) Rabbit mAb | Cell Signaling| #4695       |
| Cleaved caspase3 | Cleaved Caspase-3 (Asp175) (SA1E) Rabbit mAb | Cell Signaling| #9664       |
| GAPDH     | Rabbit-GAPDH antibody              | BETHYL       | A300-641A-M |

**Table 2.** Primary antibodies used in western blotting analysis.

**RNA-seq assay**

Total RNA was extracted from the samples using the TRizol reagent (Invitrogen, Carlsbad, CA, USA). The concentration and purity were determined in terms of optical density at A260 and A260/A280, respectively, using a spectrophotometer (Bio-Tek Instruments). RNA sequencing library of each sample was prepared using the TruSeq RNA Library Prep Kit (Illumina, San Diego, California, USA). RNA-seq experiments were performed in hBMSCs cultured under different conditions each. RNA-seq analysis was performed using the Illumina HiSeq 2000 system. Gene expression was normalized to RPKM/FPKM (reads of paired end fragments per kb of exon model per million mapped reads/fragment per kb of transcript per million mapped reads). The quality of the sequencing reads generated in the RNA-seq experiment was confirmed using the Excel-based Differentially Expressed Gene Analysis (ExDEGA) tool. A heatmap was generated to visualize transcriptomic differences between hBMSCs cultured under control medium containing 5.5 mM glucose [G(+)OM], osteo-induced hBMSCs were cultured under osteogenic medium supplemented with 5.5 mM glucose [G(+)-OM] and under osteogenic medium without glucose [G(−)OM].

**MBD-seq assay**

MBD-seq experiments were performed in groups of hBMSCs [undifferentiated hBMSCs with 5.5 mM glucose; G(−)ICM], (+) glucose [osteogenic differentiation hBMSCs with 5.5 mM glucose; G(+)ICM] and G(−)ICM glucose [osteogenic differentiation of hBMSCs without glucose; G(−)ICM]. For each sample, DNA fragments of less than 1000 bp were prepared. Analysis was performed using the Methylated DNA Enrichment Kit (New England Biolabs., Cat. E2605, USA) according to the manufacturer’s instructions. MBD2-Fc protein and Protein A Magnetic Beads were suspended. Methylated CpG DNA was captured by adding 1 μg of fragmented sample DNA, DNase-free water and 5× Bind/Wash Buffer. After incubating the DNA and MBD2a-Fc/Protein A Magnetic Beads, the supernatant was removed to wash off unbound DNA. The magnetic bead pellet sample and DNase-free water were mixed and incubated in a heat block at 65 °C for 15 min to obtain a supernatant, which contains enriched methyl CpG containing DNA. Cluster generation and 200 bp paired-end sequencing were performed on an Illumina HiSeq2500 instrument (Illumina, San Diego, CA, USA). We performed peak plot heatmap using deepTools 3.3.0. This tool creates a heatmap for scores associated with genomic regions. We performed gene peak heatmap using MeV 4.9.0. The value input into the analysis represents the raw read value of each peak.

**Statistical analysis**

The investigators were blinded to the group allocation during the experiments of the study. The in vitro experiments were performed in triplicate, with similar results obtained. Data were presented as mean ± standard deviation (SD). In this study, the significance of the differences was determined using the Student T-test to analyze multiple groups. P values less than 0.05 were considered to indicate statistical significance.

**CONCLUSION**

We mined a novel factor METTL7A based on glucose metabolic regulation and identified previously unknown roles of METTL7A in enhancing cell survival and inducing osteogenic differentiation of hBMSCs. It is expected that METTL7A overexpression possibly protects implanted cells against metabolic stress due to glucose deficiency. Accordingly, this factor can be used in stem-cell based bone tissue engineering in refractory conditions including large bone defects or osteonecrosis of femoral head.

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COMPETING INTERESTS

The authors declare no competing interests.

ADDITIONAL INFORMATION

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