SOX2 has a crucial role in the lineage determination and proliferation of mesenchymal stem cells through Dickkopf-1 and c-MYC

SB Park1,2,5, KW Seo1,2,5, Ay So1,2, MS Seo1,2, KR Yu1,2, SK Kang4 and KS Kang1,2,3

SOX2 is a well-known core transcription factor inembryonic stem cells (ESCs) and has an important role in the maintenance of pluripotency. Recently, SOX2 expression has also been reported in adult stem cells (ASCs), but the role of SOX2 in ASCs remains unknown. In this study, we examined the molecular mechanisms of SOX2 in human mesenchymal stem cells (hMSCs), a type of ASCs, by performing inhibition studies. SOX2 inhibition resulted in altered cell growth and differentiation capabilities. These changes coincided with a decrease in Dickkopf-1 (DKK1), a soluble inhibitor of WNT signaling. Chromatin immunoprecipitation and luciferase assays showed that SOX2 binds to DKK1 and has a positive regulatory role in transcription. The enforced expression of DKK1 in SOX2-inhibited hMSCs reversed the differentiation deformities, but could not abrogate the cell proliferation defect. Proliferation was regulated by c-MYC, whose expression can also be controlled by SOX2. Our study shows that SOX2 directly regulates DKK1 expression and, as a consequence, determines the differentiation lineage of hMSCs. Moreover, SOX2 also regulates proliferation by affecting c-MYC. Therefore, these results suggest that SOX2 might have a specific function by regulating DKK1 and c-MYC in the differentiation and growth of ASCs, which is separate from its roles in ESCs.

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The signaling pathways in the core transcriptional network of embryonic stem cells (ESCs) have previously been well described.1 Several core ESC transcription factors are also expressed in human mesenchymal stem cells (hMSCs).2,3 Octamer-binding transcription factor 4 (OCT4) is a POU domain-containing transcription factor, known to be essential for the stemness and pluripotency of ESCs.4 OCT4A has conserved roles in ESCs and in human umbilical cord blood-derived MSCs (hUCB–MSCs).5 OCT4 interacts with sex-determining region Y-box 2 (SOX2), and genome-wide mapping of OCT4- and SOX2-binding sites in human ESCs shows that they co-target multiple genes.6 The cis-regulatory element to which the SOX2–OCT4 complex is bound consists of neighboring SOX (5′-CATTGTT-3′) and OCT (5′-ATGCAA AT-3′) elements.7 Indeed, SOX2 was reported to be indispensable for the maintenance of ESC pluripotency;8 and ESCs from Sox2-null mice primarily differentiate into trophoderm-like cells.9 Microarray analysis has shown that SOX2 regulates multiple transcription factors that affect the OCT4 expression.10 SOX2 expression has been detected in ESCs and in several tissues derived from hMSCs, including adipose tissue, dermis, heart, and neural tissue.11 An integrated expression profile and chromatin immunoprecipitation (ChIP)–sequencing analysis showed that SOX2 is involved in the BMP signaling pathway, steroid metabolic processes, histone modifications, and many receptor-mediated signaling pathways, such as IGF1R and ITPR2.12 The well-known oncogene c-MYC is also an important transcription factor in ESCs. Many genes are regulated by c-MYC and consequently induce cell proliferation. Deregulation of c-MYC contributes to the genesis of most human tumors.13 Our previous study has shown that c-MYC is closely connected with the proliferation and epigenetic regulation of hMSCs.14 Moreover, a recent study has demonstrated a hypothetical relationship between c-MYC and SOX2.15 WNT genes, a family of 19 genes in humans and mice, produce secreted proteins that are involved in cell proliferation, differentiation, and apoptosis. These genes are also crucial for embryonic tissue development and adult tissue regeneration.16 Canonical WNT/β-CATENIN signaling is important for the stemness and differentiation of stem cells.17 The addition of Dickkopf-1 (DKK1), an antagonist of WNT signaling, allows hMSCs to re-enter the cell cycle by inhibiting the canonical WNT/β-CATENIN-signaling pathway.18 The secretion of DKK1 by hMSCs inhibits the growth of breast cancer cells.19 The presence of DKK1 triggers a disruption in the canonical WNT cascade and results in the inhibition of hMSC osteogenesis.20 DKK1 is transiently upregulated in the

1Adult Stem Cell Research Center, College of Veterinary Medicine, Seoul National University, Seoul, Republic of Korea; 2Laboratory of Stem Cell and Tumor Biology, Department of Veterinary Public Health, College of Veterinary Medicine, Seoul National University, Seoul, Republic of Korea; 3Research Institute for Veterinary Science, College of Veterinary Medicine, Seoul National University, Seoul, Republic of Korea and 4Department of Veterinary Biotechnology, Laboratory of Veterinary Biotechnology, College of Veterinary Medicine and BK21 Program for Veterinary Science, Seoul National University, Seoul, Republic of Korea
5These authors contributed equally to this work.

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early stages of hMSC adipogenesis, which indicates that the WNT cascade is important for the adipogenic differentiation of hMSCs. SOX factors appear to modulate β-Catenin/TCF activity through a variety of mechanisms, including protein–protein interactions. DNA binding, recruitment of cofactors, and protein stability. SOX2 and β-Catenin act synergistically in the transcriptional regulation of CCND1 in breast cancer cells. Recent studies have demonstrated that SOX2 contributes to the inhibition of WNT signaling and regulates osteoblast differentiation. However, SOX2 does not bind to TCF/LEF in human cells (Ingenuity Systems, Redwood City, CA, USA), and how WNT signaling is regulated by SOX2 is still unknown.

Previous studies have shown that SOX2 is expressed in several types of adult stem cells (ASCs), including neural stem cells (NSCs), dermal stem cells, and germline stem cells. However, the only known role of SOX2 is in NSCs, and relates to the self-renewal and neuronal differentiation of NSCs. The transcriptomes of NSCs, ESCs, and ASCs are quite different from each other. Therefore, SOX2 may have different roles in each type of stem cell.

Results

SOX2 is expressed in hUCB–MSCs, and cell growth is altered after SOX2 inhibition in hUCB–MSCs. To investigate the role of SOX2 in hMSCs, we examined SOX2 expression. Three well-known types of hMSCs (hUCB–MSCs, human adipose tissue-derived MSCs (hAD–MSCs), and human bone-marrow derived MSCs (hBM–MSCs)) were tested. Tera-1, an embryonic carcinoma cell line, was used as a positive control for SOX2 expression. SOX2 expression was found in hUCB–MSCs, although the levels were lower than in the tera-1 cells (Figure 1a). To quantify the SOX2 expression levels in hMSCs, quantitative reverse transcription PCR (RT-PCR) was performed (Figure 1b). hUCB–MSCs exhibited higher SOX2 expression than the other hMSCs analyzed. Furthermore, immunocytochemistry confirmed the nuclear localization of SOX2 in tera-1 cells and hUCB–MSCs (Figure 1c). Fibronectin staining was performed to determine the morphology of the cells. In the tera-1 cells, SOX2 was highly expressed in the nucleus and colocalized with Hoechst. SOX2 expression was also detected in the nuclear region of hUCB–MSCs (Supplementary Figure 1a).

To investigate the role of SOX2, knockdown was performed in hUCB–MSCs using short hairpin RNA-producing lentivirus. The sh-SOX2 construct targets the 3′-untranslated region of SOX2. A non-targeting, random sequence-inserted lentivirus was used as a control. SOX2 knockdown (sh-SOX2) in hUCB–MSCs caused a decrease in cell proliferation compared with the sh-control (Figures 1d and g). These proliferation rates correlated with SOX2 expression levels after lentiviral infection (Figure 1e). The cell cycle was also analyzed by fluorescence-activated cell sorting (FACS) in both the sh-SOX2- and sh-control-treated cells (Figure 1f). After sh-SOX2 treatment, the proportion of cells in G0/G1 increased, and the portion of cells in S phase decreased, compared with the sh-control. To confirm this phenotype, another SOX2 knockdown study was designed using a commercially available, specific siRNA for SOX2 inhibition (si-SOX2) and a non-targeting random sequence-inserted siRNA as a control (si-control). At 48 h after siRNA transfection, the cells treated with si-SOX2 displayed growth retardation compared with si-control-treated cells (Supplementary Figure 2a). By FACS analysis and MTT assays, the si-SOX2 treatment caused a decrease in S-phase composition (Supplementary Figure 2c) and a decrease in the proliferation rate (Supplementary Figure 2d).

Differentiation ability is altered after SOX2 inhibition. To validate the role of SOX2 in the differentiation of hUCB–MSCs, adipogenic, osteogenic, and chondrogenic differentiation studies were performed. After 2 weeks of adipogenic induction, a decreased number of oil droplets were found in sh-SOX2 compared with the sh-control-treated cells (Figure 2a). The optical density of oil red O after elution confirmed the morphological observations (Figure 2b). Molecular markers of adipogenic differentiation, PPARγ and C/EBP-β, were analyzed by RT-PCR (Figure 2c). The relative expression of PPARγ and C/EBP-β decreased in sh-SOX2 cells, compared with their levels in sh-control cells. An adipogenic differentiation assay was also conducted using cells transiently inhibited with siRNAs. Differentiation was initiated 48 h after of siRNA transfection and continued for 2 weeks. The cells treated with si-SOX2 manifested a decreased level of lipid droplet staining (Supplementary Figures 3a–c). However, the decrease in adipogenesis was greater in the hUCB–MSCs treated with the sh-SOX2 lentivirus. There was no difference in the elution of oil red O between the si-control and si-SOX2 conditions (Supplementary Figure 3b). The molecular markers of adipogenic differentiation were decreased. These results show that the long-term inhibition of SOX2 by lentivirus led to more severe effects on the adipogenesis of hUCB–MSCs than transient inhibition with siRNA.

After 2 weeks of osteogenic induction, alizarin red S staining was performed (Figures 2d and e). In contrast to adipogenesis, the osteogenic differentiation ability was increased by the sh-SOX2 treatment, compared with the sh-control. The expression of osteogenic molecular markers, COL1A2 and VDR, was increased in sh-SOX2 compared with sh-control-treated cells (Figure 2f). In the transient SOX2 inhibition study, there were small differences between the si-control and si-SOX2 conditions after 2 weeks of osteogenic differentiation. Alizarin red S staining was nearly identical (Supplementary Figures 3d–f), but the levels of COL1A2 and VDR were increased in si-SOX2-treated cells compared with the si-control (Supplementary Figure 3f).

After 2 weeks of chondrogenic induction, hUCB–MSCs aggregated and formed a pellet (Figure 2g). The pellet formed
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by the sh-SOX2-treated cells was larger than the sh-control. For histological evaluation, the pellets were cut into serial 3 μm sections and stained with toluidine blue (Figure 2h). Toluidine blue-stained sections showed that sh-SOX2 cells formed larger pellets than the sh-control. The expression of chondrogenic markers, SOX9 and AGGRECAN, was increased in sh-SOX2 compared with the sh-control-treated cells (Figure 2i). The same results were obtained using transient siRNA inhibition as in the lentivirus-mediated inhibition study (Supplementary Figures 3g–i). These results

**Figure 2**  Differentiation abnormalities after SOX2 knockdown in hUCB–MSCs. (a–c) Adipogenic differentiation analysis after SOX2 knockdown with the lentiviral vector. (a) The number of oil red O-stained lipid droplets decreased after SOX2 knockdown. The scale bar represents 100 μm. (b) Elution of oil red O manifested reduced numbers of lipid droplets in SOX2-knockdown hUCB–MSCs. ***P < 0.001. (c) The expression of adipogenic markers decreased after SOX2 knockdown. (d–f) Osteogenic differentiation analysis after SOX2 knockdown with the lentiviral vector. (d) The alizarin red S-stained hUCB–MSC population increased after SOX2 knockdown. The scale bar represents 100 μm. (e) The elution of alizarin red S increased after SOX2 knockdown. ***P < 0.001. (f) The expression of osteogenic markers increased after SOX2 knockdown. (g–i) Chondrogenic differentiation after SOX2 knockdown with the lentiviral vector. (g and h) The maximal diameters of sh-SOX2 cells were larger than those of the sh-control cells. The scale bar represents 100 μm. (i) The expression of chondrogenic markers increased after SOX2 knockdown

**Figure 1**  Analysis of SOX2 expression and proliferation in hMSCs. (a) RT-PCR of SOX2 in tera-1, hUCB–MSCs (UCB), hAD–MSCs (AD), and hBM–MSCs (BM). SOX2 expression in hUCB–MSCs was lower than in tera-1 cells. (b) Real-time PCR of SOX2 in hMSCs. The expression of SOX2 in hUCB–MSCs was higher than in other hMSCs. (c) Immunocytochemistry of SOX2. SOX2 expression in the nucleus was found in tera-1 cells and hUCB–MSCs. The scale bar represents 10 μm. (d) The hUCB–MSC population decreased after SOX2 knockdown. The scale bar represents 100 μm. (e) SOX2 expression decreased by 10% of the sh-control value after SOX2 knockdown by lentivirus infection. (f) The proportion of cells in S and G0/G1 phase decreased and increased, respectively, after SOX2 knockdown. (g) Cell proliferation significantly decreased after SOX2 knockdown, as indicated by the MTT assays. *P < 0.05; **P < 0.01; ***P < 0.001
demonstrated that SOX2 inhibition accelerated osteogenesis and chondrogenesis in hUCB–MSCs.

DKK1 expression decreases in SOX2-inhibited hUCB–MSCs. To determine the molecular targets of SOX2 in hUCB–MSCs, a PCR array was performed (Figures 3a, b and Supplementary Table 2). The expression of the 88 most-cited genes in stem cells was surveyed. Among them, the expression of AXIN1 (one of the canonical WNT/β-CATENIN-signaling genes) was increased by 5.2-fold after sh-SOX2 treatment, compared with the sh-control. To understand the regulation of WNT-signaling genes by SOX2 in greater detail, the levels of key components of the WNT signaling pathway (DKK1, DKK4, FZD1, FZD2, FRAT1, FRAT2, β-CATENIN, LRP5, and LRP6) were examined (Figure 3c). The levels of most WNT signaling-related genes did not change or were only slightly altered after sh-SOX2 treatment, compared with the sh-control. However, the most significant change was found in DKK1 expression; expression was significantly decreased by sh-SOX2 compared with the sh-control. Levels of phosphorylated β-CATENIN were reduced by sh-SOX2 compared with the sh-control (Figure 3d). To confirm the regulation of WNT by SOX2, a TOP/FOP FLASH assay was performed (Figure 3e). The sh-SOX2 treatment resulted in an eight-fold increase in TOP FLASH luminescence, compared with the sh-control. In case of transient inhibition, the expression levels of WNT signaling-related genes were similar to those observed upon lentivirus-mediated inhibition of SOX2 (Supplementary Figure 3).
Figures 4a and b). Moreover, the expression of TOP FLASH was increased by the si-SOX2 treatment (Supplementary Figure 4c). Therefore, DKK1 is suggested as a strong candidate gene that is regulated by SOX2 in hUCB–MSCs.

SOX2 directly binds to the DKK1 promoter region. The SOX family proteins possess a DNA-binding HMG domain and additional activation or repression domains that are implicated in transcriptional regulation.31 The evolutionarily conserved region of DKK1 was determined using database searches (UCSC genome browser, human genome March 2006 assembly, http://genome.ucsc.edu/). The human genome sequence surrounding the DKK1 transcriptional start site, from 5 kbps upstream to 5 kbps downstream, was searched. We found that the exons and the promoter region around 1 kbps upstream are highly conserved in mammalian genomic DNA (Figure 4a). SOX2 typically binds 5’-CTTTGTT-3’ or 5’-CATTGTT-3’ sequences,31 and one SOX2-binding motif was identified at −76 bps (Figure 4a).

To determine whether SOX2 binds to the DKK1 promoter region, a ChIP assay was performed. Tera-1 was used as a positive control, because SOX2 is highly expressed in tera-1 cells. SOX2 bound DKK1 in sh-control-treated cells. The signal disappeared in sh-SOX2-treated cells, and the same result was observed using siRNA-mediated inhibition (Figure 4b and Supplementary Figure 5a).

To examine SOX2 activity at the DKK1 promoter, the human genome fragment from −974 bps to +119 bps around the DKK1 transcription start site, which contains an HMG-binding domain, was cloned into the luciferase enhancer vector (Figure 4c). This region contains a predicted SOX2-binding site (B), which was confirmed by ChIP. The sh-SOX2 treatment resulted in significantly lower luciferase activity than the sh-control when transfection and luciferase activities were normalized to the SV40-promoter control vector (Figure 4d). To verify these results, SOX2 inhibition by siRNA in hUCB–MSCs was also tested. The si-SOX2 treatment resulted in significantly lower luminescence than the si-control (Supplementary Figure 5b). These results provided evidence that DKK1 expression is positively regulated by SOX2, and that DKK1 is a direct SOX2-target gene.

To confirm the SOX2 binding at the DKK1 promoter, the luciferase assay was performed using mutated constructs (Figure 4e). Plasmid M1 has a single point mutation;
second 'T' in the HMG domain-binding region was substituted with an 'A'. M1 is also the SOX2-binding sequences. Plasmid M3 has three point mutations in which the middle 'TGT' was substituted with 'CAG'. Normal hUCB–MSCs and tera-1 cells, which were not treated with lentivirus or siRNA, were used for detecting luciferase activity (Figures 4f and g). M1-transfected cells had similar luminescence activity compared with the non-mutated DKK1 promoter vector-transfected cells. However, M3-transfected hUCB–MSCs and tera-1 cells showed significantly decreased luminescence, compared with the control. These results demonstrated that the binding of SOX2 to the HMG domain of DKK1 directly regulates the transcription of DKK1.

**Ectopic expression of DKK1 in SOX2-knockdown hUCB–MSCs recovers the differentiation malformation.** A DKK1 rescue experiment in SOX2-inhibited hUCB–MSCs was performed to validate the role of DKK1 in SOX2-expressing hMSCs. The adipogenic differentiation ability increased after the ectopic expression of DKK1 in sh-SOX2-treated cells (Figure 5a). Elevated optical density after elution of oil red O was also increased by the ectopic expression of DKK1 in sh-SOX2-treated cells (Figure 5b), and the expression of adipogenic marker genes was altered. The expression of PPAR-γ and C/EBP-β increased after ectopic DKK1 expression in sh-SOX2-treated cells (Figure 5c). The osteogenic and chondrogenic ability of SOX2 knockdown hUCB–MSCs was decreased after the ectopic expression of DKK1, and this result was confirmed by morphology (d–f), alizarin red S elution (e), and marker gene analysis (f). The scale bar represents 100 μm. **P < 0.01. (d–f) The ability of hUCB–MSCs to undergo osteogenic differentiation upon SOX2 knockdown by lentivirus was decreased by the ectopic expression of DKK1, and this result was confirmed by morphology (d), alizarin red S elution (e), and marker gene analysis (f). The scale bar represents 100 μm. ***P < 0.001. (g and h) The ability of hUCB–MSCs to undergo chondrogenic differentiation upon SOX2 knockdown by lentivirus was decreased by the ectopic expression of DKK1, and this result was confirmed by pellet size (g) and marker gene analysis (h). The scale bar represents 100 μm.
hUCB–MSCs decreased after the ectopic expression of DKK1 (Figures 5d–h). The number of alizarin red S-stained cells decreased after ectopic expression of DKK1 in sh-SOX2-treated cells, and these results were confirmed by the elution of alizarin red S (Figures 5d and e). The expression of osteogenic markers (COL1A2 and VDR) decreased after the ectopic expression of DKK1 in sh-SOX2-treated cells (Figure 5f). Chondrogenesis also decreased after ectopic DKK1 expression in SOX2-knockdown hUCB–MSCs (Figures 5g and h). Therefore, DKK1 expression, which is regulated by SOX2, affects the differentiation ability.

c-MYC, not DKK1, is responsible for the proliferation defect of SOX2-inhibited hUCB–MSCs. Cellular proliferation was examined to determine whether DKK1 could rescue not only the differentiation changes, but also the proliferation defect in SOX2-knockdown hUCB–MSCs. However, unlike the differentiation results, the decreased proliferation in sh-SOX2-treated cells was not recovered by the ectopic expression of DKK1 (Figures 6a and b). Moreover, identical results were obtained using both lentiviral and siRNA inhibition studies (Supplementary Figures 6a and b). To determine whether the addition of WNT ligand could rescue the growth inhibition observed in hUCB–MSCs when SOX2 is downregulated, WNT3A and WNT5A, regarded as canonical and non-canonical WNT ligands, respectively, were added to the growth media. The portion of the SOX2-inhibited cells treated with WNT ligand in S phase of the cell cycle increased after 24 h, but there was no change in cell proliferation in response to the WNT3A or WNT5A (Figures 6c, d and Supplementary Figures 6c, d). To confirm the reason of discordance between cell cycle and proliferation test, cell cycle was checked in a time-dependent manner with WNT3A. The results showed that WNT3A influenced cell cycle in a moment, but could not maintain the increased level of S phase consistently until 3 days (Supplementary Figure 6e).

One of the candidate genes was the transcription factor c-MYC. Although c-MYC is regulated by β-CATENIN/TCF signaling,32 the expression of c-MYC was dramatically reduced in SOX2-knockdown hUCB–MSCs (Figure 6b and Supplementary Table 2). SOX2-knockdown hUCB–MSCs exhibited decreased c-MYC expression in western blot (Figure 7a and Supplementary Figure 7). To evaluate the role of c-MYC in SOX2-knockdown hUCB–MSCs, enforced expression of c-MYC using a retroviral vector was performed in sh-SOX2-treated cells (sh-SOX2 + c-MYC). A GFP expression retrovirus was used as a vector control (VC). To quantify c-MYC expression, RT-PCR was performed (Figure 7b). The enforced expression of c-MYC in sh-SOX2 cells resulted in higher expression of c-MYC than in the VC-infected sh-SOX2 cells, and the proliferation was increased after enforced c-MYC expression (Figure 7c). Cell cycle analysis showed that the enforced expression of c-MYC in sh-SOX2 cells caused an increase in S-phase composition (Figure 7d). Moreover, the MTT assay showed that the enforced expression of c-MYC in sh-SOX2 cells resulted in higher proliferation than the VC-infected sh-SOX2 cells (Figure 7e). Therefore, it was suggested that c-MYC expression, regulated by SOX2, was responsible for the growth of hUCB–MSCs.

Discussion

SOX2 is expressed in hUCB–MSCs although the levels were very low compared with the levels in tera-1 cells. Nevertheless, hUCB–MSCs exhibited stronger expression of SOX2 compared with hBM–MSCs and hAD–MSCs, which have been previously reported to express SOX2.2,33 SOX2 inhibition in hMSCs resulted in retarded cell growth, depressed adipogenic differentiation ability, and enhanced osteo-chondrogenic differentiation ability. In hMSCs, SOX2 bound to the promoter region of DKK1 and positively regulated DKK1 transcription. Ectopic expression of DKK1 in SOX2-inhibited hMSCs rescued the differentiation defects, but could not revert the proliferation defects caused by SOX2 inhibition. This proliferation defect was reduced by the additional expression of c-MYC in SOX2-inhibited hMSCs (Figure 8). The levels of SOX2 expression were variable among the three types of hMSCs. Our previous studies have proven that all three types of hMSCs exhibited self-renewability and multi-lineage differentiation potential.3,14,34 There were no
characteristic differences among these hMSCs. The reason for the differential SOX2 expression might arise from the hMSCs source. The hUCB–MSCs were obtained from the blood of newborns, but the hAD–MSCs and hBM–MSCs were collected from adult donors that were greater than 20 years old. Therefore, the hUCB–MSCs were considered more primitive than the other hMSCs.

The differentiation of hMSCs can be divided into two steps: the differentiation fate-decision step (early) and the fully differentiated step (late). Transient inhibition of SOX2 with siRNA permitted us to determine the effect of SOX2 on the early fate-decision step, and the lentivirus-mediated SOX2 knockdown allowed us to determine the effect of SOX2 on both steps of the hMSC differentiation process. In this study, the differentiation defects observed in SOX2 siRNA-inhibited hMSCs were less severe than the defects observed in hMSCs treated with sh-SOX2 lentivirus. Therefore, the effect of SOX2 on the differentiation of hMSCs occurred during the full differentiation process. Canonical WNT signaling is a mesenchymal regulatory factor that provides instructive cues for the recruitment, maintenance, and differentiation of MSCs.35 Once MSCs are committed to the osteogenic lineage, canonical WNT signaling stimulates osteogenic differentiation. Canonical WNT signaling is also likely to promote the initiation of chondrogenesis via crosstalk with TGF-β signaling.36 WNT/β-CATENIN signaling inhibits adipogenic differentiation and enhances chondrogenic differentiation.37 Therefore, the decreased adipogenic ability of SOX2-inhibited hMSCs after adipogenic induction was due to the activation of canonical WNT signaling, which occurred through DKK1 suppression in response to SOX2 knockdown.

SOX2 has an essential role in maintaining ESC pluripotency.38 A previous study has shown that SOX2-null ESCs differentiated into trophectoderm-like cells within 72 h.9 These previous results are similar to our current data using hUCB–MSCs in that, SOX2 regulates the differentiation of the stem cells. However, there are several differences between ESCs and hJBC–MSCs. SOX2 is not highly expressed in hUCB–MSCs compared with ESCs. Therefore, SOX2 inhibition alone could not induce the differentiation of hUCB–MSCs. When hUCB–MSCs are induced toward a specific differentiation pathway, the hUCB–MSCs differentiate into the induced lineages specifically.
SOX2 is also important for ESC proliferation. SOX2 promotes proliferation by facilitating the G1/S transition and by transcriptional regulation of the CCND1 gene. DKK1 is a negative regulator of the WNT signaling pathway that enhances the proliferation of many types of stem cells. In this study, the knockdown of SOX2 in hMSCs caused a reduction in the proliferation rate, although DKK1 levels were also decreased. Ectopic treatment of SOX2-knockdown hMSCs with DKK1 caused more severe proliferation defects. These results indicated that the growth of hUCB–MSCs was controlled by factors beyond WNT signaling. c-MYC is a well-known transcription factor and serves as a growth regulator in cell lines, including the hMSCs. The current study provides substantial evidence that c-MYC regulates proliferation and is regulated by SOX2 expression, although it is not clear whether SOX2 directly regulates c-MYC expression.

This study highlights two facets of SOX2 function, which are the differentiation and proliferation of hMSCs. The molecular mechanism by which the loss of SOX2 induces differentiation has been explored in these studies. DKK1 was shown to be a direct transcriptional target of SOX2 in hMSCs, which had never before been determined. The expression of SOX2 in hMSCs is also important for the proliferation and multipotency of hMSCs, largely through the regulation of DKK1 and c-MYC expression, which is quite different from its role in ESCs. This mechanism may be sustained in other ASCs, which exhibited similar levels of SOX2 expression. These results provide a novel function and mechanism for SOX2 in ASC differentiation, and will serve as an important guide for further clinical research on hMSCs and for the developmental study of ASCs.

Materials and Methods

Cell culture. HUCB–MSCs, hAD–MSCs, and hBM–MSCs were isolated and cultured as previously described. Briefly, hUCB–MSCs were obtained from umbilical cord blood immediately after full-term delivery with written consent from 20- to 30-year-old mothers. The hUCB–MSCs were maintained in endothelial cell culture growth medium-2 (Lonza, Basel, Switzerland) containing 10% fetal bovine serum (FBS). hAD–MSCs were isolated from freshly excised mammary fat tissue acquired from the BaRam Plastic Surgery Hospital. The tissues were obtained from 20- to 30-year-old women during reduction mammoplasty. The hAD–MSCs were maintained in K-SFM medium supplemented with 2 mM N-acetyl-L-cysteine (Sigma-Aldrich, St. Louis, MO, USA) and 0.2 mM L-ascorbic acid (Sigma-Aldrich). hBM–MSCs were isolated from healthy donors and were cultured in low glucose Dulbecco’s modified Eagle’s medium (DMEM) supplemented with 10% FBS, without any additional growth factors. The isolation and research use of hUCB–MSCs, hAD–MSCs, and hBM–MSCs were approved by the Boramae Hospital Institutional Review Board with written consent. All procedures were approved by the Institutional Review Board of Seoul National University (HUCB–MSC number 0603/001–002; hAD–MSC number 0600/001–001; hBM–MSC number 0910/001–003). The tera-1 cell line was purchased from ATCC (HTB-105, Manassas, VA, USA) and maintained in DMEM low-glucose medium ( Gibco BRL, Grand Island, NY, USA) containing 10% FBS ( Gibco BRL).

Transfection. Lentiviruses were generated using the ViraPower Lentiviral packaging mix ( Invitrogen, Carlsbad, CA, USA). Lipofectamine 2000 ( Invitrogen) was used to transfect 293FT cells (Invitrogen) with SHDNC-TRCN0000003252 (sh-SOX2) and SHC022 (sh-control, random sequence inserted; Sigma). The Cells were transduced with SOX2 short hairpin RNA-producing lentiviruses at an multiplicity of infection of 5–10. Polybrene (Sigma) was added to the cell culture medium at a final concentration of 6 μg/ml, and the cell culture medium was replaced with fresh culture medium the day after transfection. For selection, puromycin was added to the culture at a final concentration of 5 μg/ml for 5 days. Retroviruses were used for c-MYC overexpression. The 293FT (Invitrogen) cells were transfected with VSV-G, gag/pol (Sigma), and the c-MYC expression vector (c-MYC, pMxs-hc-MYC; Addgene, Cambridge, MA, USA) or the GFP VC (Cell biolabs, San Diego, CA, USA) using Lipofectamine 2000 ( Invitrogen). Forty-eight hours later, the supernatant was collected and filtered. The sh-control and sh-SOX2 cells were transfected with virus containing the c-MYC expression or GFP control vector with polybrene at a final concentration of 6 μg/ml ( Sigma).

Transient transfection assays were performed using commercially available, specific siRNAs against SOX2 (si-SOX2) and a non-targeting control siRNA (si-control) (ON Target plus SMARTpool, Dharmacon, Lafayette, LA, USA). The siRNA transfections were performed according to the manufacturer’s instructions. The cells were incubated with 30 nM siRNA for 48 h. After inhibition, sequential experiments were performed for genetic and characterization analyses. The DKK1 recovery assay was conducted using a DKK1 plasmid vector (number SC303946; OriGene Technologies, Rockville, MD, USA) and a control vector (number PCM3VXLS; OriGene Technologies). The plasmid transfections were performed according to the manufacturer’s instructions. The cells were incubated with 2 μg of plasmid per well of a six-well plate for 48 h. After infection, sequential experiments were performed for genetic and differentiation analyses.

RT-PCR and real-time PCR. Total RNA was isolated from cells using TRIzol reagent following the protocol provided by the manufacturer (Invitrogen). Total RNA was used for cDNA synthesis using the Superscript III First-Strand Synthesis System ( Invitrogen). For the PCR array, the RT2 Profiler Stem Cell PCR Array (catalog number PAHS-405; SuperArray, Frederick, MD, USA) was used according to the manufacturer’s protocol. For RT-PCR, cDNA and primers were combined with a PCR premix (Bioneer, Sung Nam, Republic of Korea). Real-time PCR was performed by mixing cDNA with primers, and Power SYBR Green PCR Master Mix (Applied Biosystems, Foster City, CA), Real-time PCR was performed using an ABI 7500 Real-time PCR System with supplied software (Applied Biosystems), according to the manufacturer’s instructions. RNA expression levels were compared after normalization to endogenous GAPDH. The primer sequences used in this study are listed in Supplementary Table 1.

Immunocytochemistry. The cells were fixed in 4% paraformaldehyde for 10 min at room temperature. The cells were permeabilized by exposure to 0.5% Triton X-100 for 10 min at room temperature and were blocked for 2 h with 10% normal goat serum (Zymed, San Francisco, CA, USA) at room temperature. Primary antibodies were used at the manufacturer’s recommended dilution in 5% normal goat serum. Rabbit anti-SOX2 (number ab56603, Millipore, Billerica, MA, USA) and mouse anti-fibronectin (number ab6328, Abcam, Cambridge, MA, USA) were used for immunostaining. Fibronectin staining was utilized for cytological observation. The secondary fluorescence conjugated antibodies, goat anti-mouse Alexa Fluor 488, and goat anti-rabbit Alexa Fluor 594 (Invitrogen), were each used at a 1:1000 dilution. Hoechst (Sigma-Aldrich) was used for counterstaining.

MTT cell proliferation assay and FACS assay. The proliferation potential was measured using the MTT assay, which is based on the ability of live cells to convert tetrazolium salt into purple formazan. Briefly, the cells (1 × 10^6 cells per well) were seeded in 24-well microplates in 450 μl media. After 48 h, 50 μl of MTT stock solution (5 mg/ml; Sigma) was added to each well, and the plates were further incubated for 4 h at 37 °C. The supernatant was removed, and 200 μl DMEM was added to each well to solubilize the water-insoluble purple formazan crystals. The absorbance at a wavelength of 540 nm was measured with an EL800 microplate reader (Bio-TEK Instruments, Winooski, VT, USA).

The DNA content was determined by propidium iodide staining intensity. The cells were fixed with ice-cold 70% ethanol at 4 °C for 1 h. Following fixation, 50 μg/ml propidium iodide (Sigma-Aldrich) and 100 μg/ml RNase A (Invitrogen) were added at 37 °C for 1 h. Cell cycle analysis was performed using a FACS Calibur instrument (Becton-Dickinson Pharmingen, San Jose, CA, USA) equipped with ModFit LT (Verity Software House, Topsham, ME, USA).

In-vitro differentiation assays. To induce adipogenic and osteogenic differentiation, hMSCs were treated as previously reported. Briefly, the cells were treated with an adipogenic induction medium (DMEM supplemented with 10% FBS, 200 μM indomethacin, 1 μM dexamethasone, 0.5 mM isobutyl methylyxanthine, and 0.5 μg/ml insulin) or an osteogenic induction medium (DMEM containing 10% FBS, 100 mM dexamethasone, 50 μM ascorbic acid 2-phosphate, and 10 mM β-glycerophosphate; Sigma-Aldrich). After 2 weeks of induction, the cells were stained to confirm that differentiation occurred. For adipogenic differentiation, the cells were stained with oil red O (Sigma-Aldrich).
Bound stain was solubilized using 10% isopropanol, and the released solution was measured at 500 nm, using a spectrophotometer. For osteogenic differentiation, alizarin red S staining was performed. The release of solubilized alizarin red S was measured at 570 nm using a spectrophotometer. To induce chondrogenic differentiation, 1 × 105 cells was added to a 15 ml polypropylene tube and pelletted by centrifugation. The pellet was cultured at 37°C in a 5% CO2 incubator in 1 ml of chondrocyte differentiation medium (Lonza) for 2 weeks. After 2 weeks, the round pellets were embedded in paraffin and cut into 3 μm sections. For histological evaluation, the sections were stained with toluidine blue.

**Western blotting.** Whole-cell protein lysates were extracted with a solution containing 1% Triton X-100, 20 mM Tris HCl (pH 8), 137 mM NaCl, 10% glycerol, and 2 mM EDTA (Sigma-Aldrich). Nuclear and cytoplasmic lysates were extracted using NE-PER reagents (Thermo, Rockford, IL, USA). The protein concentrations were determined using a DC assay kit (Bio-Rad, Hercules, CA, USA), and proteins were separated by 10% SDS-polyacrylamide gel electrophoresis. The separated proteins were then transferred to nitrocellulose membranes at 100 V and 350 mA for 2 h, and probed using the following primary antibodies: β-CATENIN (number 9562; Cell Signaling Technology Inc., Danvers, MA, USA), p-β-CATENIN (number 9561; Cell Signaling Technology Inc.), and β-ACTIN (number 3700; Cell Signaling Technology Inc.). Secondary antibodies were used according to the manufacturer’s instructions. Secondary-antibody binding was detected using an enhanced chemiluminescence detection kit (Amersham, Piscataway, NJ, USA) according to the manufacturer’s instructions. Protein levels were normalized to β-ACTIN using Image J analysis software.

**ChiP and luciferase reporter assays.** ChiP assays were performed according to the manufacturer’s protocol (Upstate Biotechnology, Waltham, MA, USA). Chromatin was immunoprecipitated using rabbit anti-human SOX2 antibodies (number ab158530, Abcam). PCR was performed at a final template dilution of 1:50. The primer sequences used in this study are supplied in Supplementary Table 1. The luciferase assays, the DKK1 promoter was cloned into the pGL3-enhancer vector (Promega, Madison, WI, USA), and the pGL3-SV40-control vector was used as a transfection control. The luciferase reporter assays were performed according to the manufacturer’s instructions. Briefly, 0.75 μg of pGL3-enhancer vector containing the DKK1 promoter, or the pGL3-control vector was transfected using Tfx-50 transfection reagent (Promega) at a 2:1 ratio of Tfx-50 plasmid per well, using the Tfx-50 transfection reagent system (Promega). The ratio of the target gene to the housekeeping gene was calculated and expressed as 2-ΔCT. Statistical analyses were conducted using 2-ΔCT. This ratio was then used to evaluate the expression of each target gene in SOX2-inhibited hUCB-MSCs and control hUCB-MSCs. To determine the changes in gene expression, the normalized expression of the target genes in SOX2-inhibited hUCB-MSCs was divided by the normalized expression of the same target gene in the control samples. All experiments were conducted in triplicate, and the results are expressed as the mean ± S.D. Statistical analyses were conducted via analysis of variance, followed by Duncan’s multiple range tests or Student’s t-test. P < 0.05 was considered to be significant.

**Conflict of Interest**

The authors declare no conflict of interest.
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Supplementary Information accompanies the paper on Cell Death and Differentiation website (http://www.nature.com/cdd)