microRNA-320/RUNX2 axis regulates adipocytic differentiation of human mesenchymal (skeletal) stem cells

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The molecular mechanisms promoting lineage-specific commitment of human mesenchymal (skeletal or stromal) stem cells (hMSCs) into adipocytes (ADs) are not fully understood. Thus, we performed global microRNA (miRNA) and gene expression profiling during adipocytic differentiation of hMSC, and utilized bioinformatics as well as functional and biochemical assays, and identified several novel miRNAs differentially expressed during adipogenesis. Among these, miR-320 family (miR-320a, 320b, 320c, 320d and 320e) were ~2–3-fold upregulated. Overexpression of miR-320c in hMSC enhanced adipocytic differentiation and accelerated formation of mature ADs in ex vivo cultures. Integrated analysis of bioinformatics and global gene expression profiling in miR-320c overexpressing cells and during adipocytic differentiation of hMSC identified several biologically relevant gene targets for miR-320c including RUNX2, MIB1 (mindbomb E3 ubiquitin protein ligase 1), PAX6 (paired box 6), YWHAH and ZWILCH. siRNA-mediated silencing of those genes enhanced adipocytic differentiation of hMSC, thus corroborating an important role for those genes in miR-320c-mediated adipogenesis. Concordant with that, lentiviral-mediated stable expression of miR-320c at physiological levels (~1.5-fold) promoted adipocytic and suppressed osteogenic differentiation of hMSC. Luciferase assay validated RUNX2 (Runt-related transcription factor 2) as a bona fide target for miR-320 family. Therefore, our data suggest miR-320 family as possible molecular switch promoting adipocytic differentiation of hMSC. Targeting miR-320 may have therapeutic potential in vivo through regulation of bone marrow adipogenesis.

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Bone marrow fat is increasingly recognized as an important component of the bone marrow microenvironment with potential role in regulating bone formation, hematopoiesis and the whole body’s energy metabolism.1,2 During aging and in a number of skeletal diseases, an inverse relationship between bone marrow trabecular bone mass and fat mass has been reported, suggesting a common regulatory genetic program.3–6 Based on a large number of in vitro studies, bone marrow adipocytes (ADs) and osteoblasts originate from a common progenitor cells within the bone marrow stroma known as mesenchymal (skeletal or stromal) stem cells (MSCs).7 It is thus envisaged that controlling MSC fate into osteoblasts or AD can be a target for intervention with the aim of enhancing bone formation in bone loss disorders.8 To achieve this goal, molecular mechanisms controlling MSC commitment to ADs versus osteoblasts need to be identified.

MicroRNAs (miRNAs) are double-stranded noncoding RNA molecules of ~22 nucleotides that function as post-transcriptional regulators of gene expression and are found in a wide variety of organisms, from plants, insects to humans.9,10 miRNAs have been identified to affect multiple biological functions including stem cell differentiation, neurogenesis, hematopoiesis, immune response, skeletal and cardiac muscle development.11–17 Several previous studies have identified a number of miRNAs as important regulators of MSC differentiation into osteoblasts (for review, see Taipaleenmaki et al.,18 Eskildsen et al.19 and Zeng et al.20) and chondrocytes.21 On the other hand, few studies have examined miRNA regulation of MSC differentiation into ADs.22,23 In the present study, we carried out a comprehensive analysis of miRNA expression profiling and bioinformatics analyses of human MSC (hMSC) during in vitro AD differentiation. We identified several novel pro-adipogenic miRNAs, and found that miR-320 to be an important regulator of adipocytic differentiation of hMSC.

Results

Identification of differentially expressed miRNAs during adipocytic differentiation of hMSCs. Using standard AD-induction medium (AIM), hMSC differentiated readily into

**Abbreviations:** 3′UTR, 3′-Untranslated region; C/EBP, CCAAT-enhancer-binding protein; DMEM, Dulbecco’s modified Eagle’s medium; FACS, Fluorescence-activated cell scan; FBS, Fetal bovine serum; HEK-293, Human embryonic kidney 293 cells; hMSCs, Human mesenchymal stem cells; hTERT, Human telomerase reverse transcriptase; miRNAs, MicroRNAs; MSCs, Mesenchymal stem cells; PBS, Phosphate-buffered saline; PPARγ, Peroxisome proliferator-activated receptor-γ; qRT PCR, Quantitative real-time reverse transcription PCR; RUNX2, Runt-related transcription factor 2; MIB1, Mindbomb E3 ubiquitin protein ligase 1; PAX6, Paired box 6

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mature lipid-filled ADs as demonstrated by positive staining for Oil Red O (Figure 1a) and increased expression of several AD-specific genes (Figure 1b). Global miRNA expression profiling carried out on AD-differentiated hMSC revealed 38 miRNAs to be differentially expressed on day 13 compared with day 0 ($P<0.05$; Figure 1c and Table 1). Eleven miRNAs were found to be differentially expressed on day 7 compared with day 0 ($P<0.05$; Figure 1d and Supplementary Table 1). The expression levels of selected group of miRNAs identified in the microarray experiment, miR-374–5p, –30b, –222, –320c, –186, –320a, –320e and –29c, were validated using quantitative real-time reverse transcription PCR (qRT-PCR) that confirmed the microarray results and showed upregulation of miR-30b, miR-320 family (320a/320c/320e) on day 7 post-AD differentiation induction and further increase in expression levels of the same miRNAs in addition to miR-186 on day 13 (Figure 1e). miR-222 was found to be downregulated on day 13 (Figure 1e). Among the identified miRNAs, several members of the miR-320 (miR-320a, 320b, 320c, 320d and 320e) family were differentially expressed and were chosen for further investigation, as they have not previously been implicated in regulating the adipogenic differentiation of MSCs.

Overexpression of miR-320c and miR-30b promote adipogenic differentiation of hMSCs. To examine for the potential role of selected miRNAs, miR-320c and -30b in regulating the adipogenic differentiation of hMSCs, cells were transfected with pre-miR-320c, pre-miR-30b or pre-miR-negative control and subsequently were exposed to AIM. qRT-PCR revealed significant increase in mRNA expression in transfected cells (data not shown). As shown in Figure 2a, cell transfected with pre-miR-320c and -30b exhibited enhanced formation of lipid-filled mature ADs. Concordant with those data, Nile red staining and fluorescence-activated cell scan (FACS) analysis revealed increased number of Nile Red High population in hMSC cultures transfected with pre-miR-320c and -30b compared with the controls (Figures 2b and c). As miR-320 family was the most novel family of miRNAs identified in current study as a possible regulator of adipocytic differentiation of hMSCs, all subsequent experiments focused on miR-320c member. In order to confirm that the enhanced adipocytic differentiation mediated via miR-320c was specific and not because of nonspecific effect as a result of transfection, we generated hMSCs stably expressing miR-320c using lentiviral-mediated transduction. As shown in Figures 2d and e, stable expression of miR-320c indeed led to enhanced adipocytic differentiation of hMSCs compared with cells transduced with control lentivirus. Representative images of Oil Red O staining are shown in Figure 2e, while quantification of Oil Red O staining demonstrated enhanced adipogenesis in LV miR-320c cells compared with control cells (Figure 2f). Similarly Nile red staining and quantification also demonstrated enhanced lipid droplet accumulation in miR-320c compared with control cells (Figures 2g and h). We observed no significant difference in cell viability on day 7 post-AD differentiation induction between LV miR-320c and LV control cells (Figure 2i), therefore the difference in Nile red staining is not due to difference in cells numbers. Concordant with that, the expression of AD-specific genes was higher in LV miR-320c cells compared with control cells (Figure 2j). Taken together, our data indicated enhanced adipocytic differentiation of hMSCs overexpressing miR-320c.

Identification of bona fide mRNA targets for miR-320c. In order to identify possible gene targets of miR-320c that regulate MSC differentiation into ADs, we used the following approach. hMSC transfected with miR-320c or control miRNA were cultured in presence or absence of AIM, total RNA was extracted from baseline and following 7 days of AD induction and subjected to microarray analysis. As shown in Figures 3a and b, hMSC overexpressing miR-320c did cluster together compared with cells transfected with miRNA control. Supplementary Tables 2 and 3 contain a list of genes that were downregulated (–1.3-fold, $P<0.05$) in hMSCs transfected with miR-320c compared with control cells at 72 h post-transfection and in day 7 adipocytic induced hMSC, respectively. In addition to these genes that were experimentally determined, a third list of predicted miR-320c gene targets were curated from TargetScan database. The intersection in Venn diagram between three lists, identified 210 common genes, that is, genes that were downregulated upon miR-320c overexpression, downregulated during adipocytic differentiation of hMSCs and that were predicted to be targeted by miR-320c in silico. Gene ontology and pathway analysis of the identified 210 common genes showed strong enrichment for genes involved in regulating the cell cycle and cell differentiation (Figure 3d and Supplementary Table 4). Concordant with those data, we observed that un-induced hMSC stably expressing miR-320c had slower proliferation rate compared with control cells (Figure 3e). Several genes involved in cell cycle regulation and cell differentiation that were common to the three lists (MIB1, PAX6, ZWILCH, YWHAH and SEMA5A) or genes that were common to Exp. determined and in silico (TGFBR1, TGFBR2, NRP1, RASA1, ULK1, RUNX2 (Runt-related transcription factor 2), BMPR1A and KITLG) were chosen for further investigation. As confirmation of the microarray data, qRT-PCR showed good concordance between microarray and qRT-PCR except for one gene KITLG (Figure 3f).

Validating the role of selected miR-320c targets in regulating adipocytic differentiation of hMSC. To confirm whether the identified miR-320c targets are indeed involved in regulating adipocytic differentiation of hMSCs, we used siRNA approach to knock down the expression of selected genes. As shown in Figure 4a, siRNA-mediated silencing of MIB1, PAX6, RUNX2, YWHAH and ZWILCH led to increased number of adipocytic cells differentiated from hMSC. Concordant with that, qRT-PCR indicated upregulation of several adipocytic markers, AdipoQ, PPARγ, PPARδ (peroxisome proliferator-activated receptor-δ) and FABP4, in cells transfected with YWHAH, MIB1, RUNX2 and ZWILCH siRNAs, suggesting a plausible role for these genes in miR-320-mediated effects on adipocytic differentiation of hMSC.

miR-320c suppresses osteogenic differentiation of hMSCs. Interestingly, RUNX2, which is a key transcription factor (TF) involved in osteogenesis, was among the novel
miRNA expression profiling during adipocyte (AD) differentiation of hMSC. hMSC-TERT cells were induced to AD differentiation. (a) Oil Red O staining of lipid-filled mature ADs on days 7 and 13. (b) qRT-PCR analysis of AD marker genes (peroxisome proliferator-activated receptor-γ, PPARγ; AD protein 2, AP2; leptin, LEP; adiponectin, AdipoQ). Gene expression was normalized to GAPDH and β-actin. Data are presented as mean ± S.E. of fold changes compared with non-induced controls, n = 6 from two independent experiments. ***P < 0.0005 between non-induced and induced samples. hMSC were induced to AD differentiation and on days 0, 7 and 13, miRNA expression profiling was done using the miRCURY LNA miRNA Array (6th GEN). (c) Heat map and unsupervised hierarchical clustering were performed on the top 30 miRNAs differentially expressed on AD day 13 (D13) versus AD D0; the color scale illustrates the relative expression level of miRNAs (log2). Red color represents an expression level below the reference channel, and green color represents expression higher than the reference. (d) Venn diagram depicting the overlap in miRNAs that were differentially expressed on AD D13 and AD D7. (e) Validation of selected miRNAs identified in c using Taqman miRNA qRT-PCR. Data are presented as mean ± S.E., n = 6. *P < 0.05, ***P < 0.0005.

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gene targets identified for miR-320c family in current study. Therefore, we sought to assess the effect of miR-320c expression on osteogenic differentiation of hMSC. Data presented in Figure 5a showed lower ALP staining in LV miR-320c cells compared to LV controls (Figure 5c). There was no difference in cell viability of OB-differentiated LV miR-320c and LV control hMSC (Figure 5d).

**Identification of RUNX2 as bone fide target for miR-320c during adipogenesis.** Among the miR-320c-identified gene targets, RUNX2 was the most prominent. In addition to TargetScan, another database (HOCTAR, http://hoctar.tigem.it/), which examines for an inverse relationship between the expression of miRNA host gene and a potential target, was also used. We found that miR-320 family to be among the top 10% miRNAs predicted to regulate RUNX2, which further supports a role for miR-320 family in regulating RUNX2 expression during AD differentiation (Supplementary Table 5). Interestingly, RUNX2 had four predicted miR-320 family binding sites on it 3' untranslated region (3'UTR) located between nucleotides 1175 and 3142 (Figure 6a). Over-expression of pre-miR-320c led to significant reduction in RUNX2 expression in hMSC (Figure 6b). To confirm that RUNX2 is indeed a direct target for miR-320 family, we constructed reporter vector carrying the predicted binding site(s) of RUNX2 downstream of a firefly luciferase gene in the pMIR-REPORTER miRNA Expression Reporter vector (Figure 6c). A mutant version of RUNX2 UTR reporter vector with mutations in the predicted miR-320 seed region(s) in the 3' UTRs was also generated using the primer combination listed in Table 2. The pRL-SV40 (encoding for renilla luciferase) was used for normalization. Co-transfection experiments in HEK-293 (human embryonic kidney 293) cells using two different RUNX2 UTR reporter clones demonstrated significant regulation of the RUNX2 reporter by miR-320c miRNA (~50%; Figure 6d). The regulation of RUNX2 UTR by miR-320c was specific, as mutating the seed region completely abrogated this effect.

**Discussion**

In the present study, we have identified miR-320 family as novel regulator of bone marrow-derived hMSC differentiation into ADs. Our data corroborate an increasing number of studies demonstrating the role of miRNAs in regulation of
Figure 2  Forced expression of miR-320c- and miR-30b-enhanced AD differentiation of hMSCs. hMSCs were transfected with 30 nM of pre-miR-320c, pre-miR-30b, and pre-miR-Neg, then were subjected to AD differentiation. (a) AD differentiation was assessed on day 7 using Oil Red O staining. (b) The percentage of Nile red-positive cells was enumerated using flow cytometry. (c) Quantitative presentation of the data obtained in b. Data are presented as mean ± S.E, n = 3. (d) Stable expression of miR-320c in hMSC cells using lentiviral expression vector led to significant increase in miR-320c expression. (e) Oil Red O quantification in LV miR-320c and LV control cells after 7 days of adipocytic differentiation. (f) Nile red staining of LV miR-320c and LV control cells on day 7 adipocytic induction. The level of Nile red staining was quantified using molecular devices M5 microplate reader using fluorescence well-scan mode (f). Data are representative of three independent experiments, n = 36. Cell numbers was quantified using the alamarBlue assay on LV miR-320c and LV control cells (i), n = 9. (j) Expression of adipo-specific markers in LV miR-320c or LV control cells after 7 days of adipocytic differentiation, n = 6, *P < 0.05, **P < 0.005, ***P < 0.0005
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Figure 3  Identification of miR-320c bona fide gene targets during adipogenic differentiation of hMSCs. (a) Hierarchical clustering of hMSC transfected with miR-320c or control miRNA 72 h post transfection based on mRNA expression levels. Each column represents one replica. Expression level of each gene in a single sample is depicted according to the color scale. (b) Hierarchical clustering of control MSCs or MSCs differentiated into ADs (day 7) based on mRNA expression levels, where each column represents one replica. Expression level of each gene in a single sample is depicted according to the color scale. (c) Venn diagram demonstrating the overlap between experimentally determined miR-320c targets at baseline or following adipocytic differentiation of hMSC, and the in silico-predicted miR-320c targets based on TargetScan database. (d) Pie chart illustrating the distribution of the top 20 GO categories for the 210 predicted miR-320c gene targets. The pie section size corresponds to fold enrichment. (e) Quantification of cell viability of control hMSC or hMSC stably transduced with miR-320c LV on day 4 using alamarBlue assay. (f) qRT-PCR validation (blue) of selected miR-320c gene targets identified from microarray (red) (c). Data are presented as mean ± S.E., n = 6 from two experiments, *P < 0.05, **P < 0.005.
hMSC cell lineage fate as well as increasing number of other types of stem cells. Although several miRNA candidates that control osteoblastic differentiation of MSC have been described, only few miRNA have been reported to regulate their adipocytic differentiation. Among the reported miRNAs are miR-143, miR-138 and miR-637 that were implicated in regulating AD differentiation via modulation of ERK5, EID-1 and Osterix, respectively.

In current study, we used an integrated analysis of miRNA expression profiling combined with bioinformatics analyses. Interestingly, several of the identified differentially regulated miRNAs during AD differentiation of hMSC have previously been reported to regulate hMSC differentiation (e.g., miR-222, miR-138 and miR-30 family), indicating the importance of the regulatory network controlled by miRNAs as they are preserved across different cellular models of MSC.

Figure 4  Functional validation of the identified miR-320c targets in regulating adipocytic differentiation of MSCs. hMSC were transfected with the indicated siRNA or control siRNA and were subjected to adipocytic differentiation induction for 7 days. (a) Oil Red O staining of mature lipid-filled ADs on day 7. (b) qRT-PCR analysis of AD marker genes (AdipoQ, PPARγ and FABP4). Gene expression was normalized to GAPDH and β-actin. Data are presented as mean fold change compared with cells transfected with control siRNA ± S.E., n = 6 from two independent experiments. *P < 0.05, **P < 0.005, ***P < 0.0005
We identified miR-320 family as the most prominent novel regulator of hMSC differentiation into ADs. Using Tri-Pronged approach combined with functional and biochemical assays, we identified several novel gene targets for miR-320 family during adipocytic differentiation of hMSC. Among these MIB1, PAX6, YWHAH, ZWILCH and RUNX2 were most relevant to adipogenesis. Interestingly, several of the identified genes are known to have a role in regulating cell proliferation and stem cell differentiation. For example, MIB1 and PAX6 were implicated in regulating neural stem cell differentiation.\(^{31,32}\) YWHAH has recently been implicated in regulating cell division during meiosis.\(^{33}\) while ZWILCH has been shown to be essential for kinetochore functions during cell division.\(^{34}\) However, our data revealed an additional role of these proteins in bone marrow adipogenesis. RUNX2 was one target identified in this study that is known for its being a master TF for inducing osteoblast differentiation. Therefore, it is plausible that miR-320 family promote adipogenesis via blocking other MSC differentiation pathways (i.e., osteoblast; Figure 6d).

Bioinformatics analysis revealed that RUNX2 3’ UTR harbors four potential binding sites for miR-320 family. Regulation of RUNX2 expression by miR-320 was subsequently confirmed using qRT-PCR and luciferase assay. The interaction between miR-320 and RUNX2 3’ UTR was found to be specific, as mutating miR-320 seed region in the 3’ UTR of RUNX2 completely abrogated its regulatory effects. RUNX2 is osteoblast-specific TF that has an important role in MSC differentiation to osteoblasts.\(^{35-37}\) Previous studies have demonstrated that adipocytic differentiation of MSC is suppressed by RUNX2 and RUNX2\(^{-/-}\) calvarial cells exhibited an enhanced AD differentiation.\(^{38}\) Our data corroborate that osteoblast differentiation is the primary default differentiation pathway for bone marrow-derived hMSC and thus

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**Figure 5** Overexpression of miR-320c suppressed ALP activity in hMSC. (a) Representative hMSC were stably transduced with a lentivirus containing miR-320c or control miRNA and were subjected to osteoblast differentiation induction for 10 days. ALP staining for control hMSC (LV control) or hMSC stably expressing miR-320c (LV miR-320c) is shown. (b) qRT-PCR analysis of osteoblast gene markers (RUNX2 and osteonectin). (c) Quantification of ALP activity on cells from a. Data is presented as relative ALP activity compared to cells transduced with LV control. Data are presented as mean ± S.E. from five independent experiments, \(n = 50\) (d) Quantification of cell viability of LV control or LV miR-320c cells on day 10 post-osteogenic induction showing no significant difference between the two groups, *\(P < 0.05\), ***\(P < 0.0005\)
inhibition of RUNX2 activity is needed to promote AD differentiation. Interestingly, previous studies have also identified RUNX2 as a bona fide target for pro-adipocytic miRNAs such as miR-30a, 30d and miR-204/211. Therefore, RUNX2 appears to be a key negative regulator of adipogenesis that seems to be targeted by several miRNA families, including the miR-320 family in our study. This is not surprising as RUNX2 has relatively large 3′ UTR (3.777 kb) that makes it a likely target for several groups of miRNAs. Interestingly, lentiviral-mediated stable expression of miR-320c at physiological levels (~1.5-fold; Figure 2d) also promoted adipocytic differentiation of hMSC, thus further supporting that the observed effects have physiological relevance.

MSC commitment to specific lineage, AD or osteoblast, was shown to be regulated by the expression of different TFs, which are involved in different cellular pathways. MSCs express several adipogenic TFs, for example, CCAAT-enhancer-binding protein (C/EBP) and PPARγ, as well as osteoblastic TF, for example, RUNX2, MSX2, DLX5 and Osterix.39–46 It is plausible that the undifferentiated state of MSC is maintained by suppression of lineage-specific TFs. Most of the studies focused on the ability of miRNA to induce
humidified atmosphere containing 95% air and 5% CO₂ at 37 °C, medium was
in vivo
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and the cells were cultured in the same culture medium as above. All reagents were
pyruvate, 10% fetal bovine serum (FBS), 1% penicillin
dexamethasone, 0.45 mM isobutyl methyl xanthine (Sigma), 3
hMSCs. Adipogenic differentiation of hMSCs. Cells were cultured as above then were exposed to osteogenic induction medium (DMEM containing 10% FBS, 1% Pen–Strep,50 μg/ml L-aspartic acid (Wako Chemicals GmbH, Neuss, Germany), 10 mM β-glycerophosphate (Sigma) and 10 nM calcitriol (1α,25-dihydroxy vitamin D₃; Sigma) and 10 nM Dexametha-
sone (Sigma)).
Lentiviral transduction. Lentiviral particles encoding for has-miR-230c-1
LP-000302-0200) or control lentiviral particles (LP-000302-0200) were purchased from Genecopoeia (Genecopoeia Inc., Rockville, MD, USA). Hundred thousand hMSCs were seeded in complete DMEM in 24-well plate. Forty-
eight hours later (~80 confluency), media was removed and then 20 μl of crude lentiviral particles in 500 μl of DMEM+5% heat-inactivated serum (Invitrogen) and 1% Pen–Strep supplemented with polybrene (8 μg/ml; Sigma) was added to the cells. Seventy-two hours later, media was removed and transduced cells were selected with puromycin (1 μg/ml; Sigma) for 1 week until stably transduced cells were generated.

Total RNA isolation and quantification of miRNA and mRNA expression. Total RNA containing the small RNA fraction was isolated from hMSC-TERT cells using Total RNA Purification Kit (Norgen-Biotek Corp., Thorold, ON, Canada) according to the manufacturer’s instructions. The concentrations of total RNA were measured using NanoDrop 2000 (Thermo Scientific, Wilmington, DE, USA). qRT-PCR analysis was performed as previously described29 to assess the expression levels of miRNAs using TaqMan miRNA Assays (Applied Biosystems Inc., Foster City, CA, USA). In brief, 10 ng of total RNA was subjected to reverse transcription using miRNA-specific primers supplied by ABI. Subsequently, second set of miRNA-specific primers was used to amplify each miRNA according to the manufacturer’s recommendations. Expression levels of adipogenic-related genes, PPARγ, LEP, AP2 and AdipoQ, were assessed using qRT-PCR. Reverse transcription was performed on 500 ng of total RNA using High Capacity Reverse Transcriptase Kit (Applied Biosystems Inc.) according to manufacturer’s specifications. qRT-PCR was done using FAST-SYBR Green Master Mix (Applied Biosystems Inc.) and the StepOne Plus Real-Time PCR Detection System (Applied Biosystems Inc.). Primers used for gene expression analysis are listed in Table 1 and were either previously published or were designed using NCBI Primer-BLAST (http://www.ncbi.nlm.nih.gov/tools/primer-blast/). The Z score value method was used to calculate relative expression of miRNAs and mRNAs.31

miRNA expression profiling. All miRNA microarray experiments and analyses were conducted by Exiqon (Copenhagen, Denmark). hMSC-TERT cells were differentiated into ADs, and on days 0, 7 and 13 RNA was extracted as described above.19 The samples were labeled using the miRCURY LNA microRNA Hi-Power Labelling Kit, Hy3/Hy5, and hybridized on the miRCURY LNA microRNA Array (6th GEN) as outlined in Supplementary Figure 1. Samples that did not meet quality requirements were excluded from the data analyses. Data were normalized using the global Lowess regression algorithm, then were subjected to unsupervised analysis using functional-annotation tools, as we have previously described.52 miRNA expression data sets were deposited to the Gene Expression Omnibus (GEO), accession number GSE59684.

miRNA and siRNA transfection experiments. To investigate the role of selected miRNAs in regulating hMSC differentiation into ADs, hMSC cells were transfected with the indicated miRNA precursors (pre-miR-Neo, pre-miR-320c and pre-miR32, Ambion, Foster City, CA, USA) or siRNAs (Ambion) using reverse transfection protocol and Lipofectamine 2000 (Invitrogen) as we previously described.32 In brief, 0.05 × 10⁶ cells were reverse-transfected with 30 nM of the indicated pre-miR or siRNAs complexed with 1.5 μl of Lipofectamine 2000 in a 24-well tissue culture plate. Transfection cocktail was subsequently replaced after 4 h with normal DMEM without antibiotics. On day 3, medium was replaced with DMEM+AIM as described above and fresh induction medium was replaced every 3 days.
Cloning of RUNX2 3′ UTR and luciferase assay. A reporter vector (pMR-Report, ABI) carrying the predicted miR-320 binding site(s) from RUNX2 3′ UTR was constructed using partially complementary primer pairs listed in Table 1. Amplification was conducted as we previously described\textsuperscript{24} and using AmpliTaq gold DNA polymerase (Applied Biosystems Inc.). As positive control, we constructed a vector carrying the full-length complementary sequence to Let-7b miRNA. A mutant version of RUNX2 3′ UTR reporter plasmid was generated by mutating the seed region for the miR-320 miRNA family using the indicated primers in Table 1. All regions were subsequently cloned into the SpeI and HindIII sites downstream of the firefly lucerase gene in the pMR-REPORT vector (Applied Biosystems Inc.). To assess the direct interaction between miR-320 miRNA family and the 3′ UTR from RUNX2, HEK-293 cells were co-transfected with 100 nM of pre-miR-Neg or pre-miR-320c and 100 ng of pMR-REPORT carrying either wt or mutant 3′ UTR sequences, along with 20 ng of pRL-SV40 vector (Promega, Madison, WI, USA) carrying the Renilla lucerase gene. Transfection experiments were conducted using Lipofectamine 2000 (Invitrogen). At 48 h post transfection, luciferase activity was measured using the Dual-Glo lucerase assay system (Promega). Firefly lucerase activity was then normalized to that of Renilla lucerase.

Gene expression microarray. hMSC were differentiated into ADs as described above. On day 7, total RNA was extracted using Total RNA Purification Kit (Thermo Scientific). Extracted RNA was labeled and then hybridized to the Agilent Human SurePrint G3 Human GE 8×60 k microarray chip (Agilent Technologies, Santa Clara, CA, USA). All microarray experiments were conducted at the Microarray Core Facility ( Stem Cell Unit, King Saud University College of Medicine, Riyadh, Saudi Arabia). Data analyses were conducted using GeneSpring 12.0 software (Agilent Technologies) and DAVID bioinformatic tool as described before.\textsuperscript{52,53} Percentile Shift was used for data normalization while Benjamini-Hochberg false discovery rate method was used for multiple testing corrections. The gene expression profiling in hMSCs transfected with miR-320c and the gene expression profiling during adipogenic differentiation of hMSCs data sets were deposited to the GEO accession numbers GSE59458 and GSE59450, respectively.

Oil Red O staining for ADs. At the indicated time points, adipogenic differentiation was determined by Oil Red O staining for lipid-filled mature ADs. Cells were washed with phosphate-buffered saline (PBS), fixed with 4% paraformaldehyde for 10 min and were incubated with a newly made and filtered (0.45 μm) Oil Red O staining solution (Sigma; 0.5% in 60% isopropanol) for 1 h at room temperature. Photomicrographs were acquired using inverted Zeiss microscope (Thornwood, NY, USA).

AD enumeration by flow cytometry. Nile Red Staining was performed as we described previously.\textsuperscript{17} In brief, following trypsinization, the cells were washed with calcium and magnesium-free PBS. Subsequently, Nile Red dye (N3013; Sigma) was added at a final concentration of 100 ng/ml. Following 5 min incubation at 4 °C, the cells were washed in PBS, centrifuged and re-suspended in 500 μl PBS and were analyzed using BD FACS Calibur flow cytometer (BD Biosciences, Franklin Lakes, NJ, USA). Staining was detected in the green fluorescence channel (FL1); the gating strategy is presented in Figure 2. Data were analyzed using FlowJo software (Tree Star, Ashland, OR, USA).

Nile red fluorescence determination and quantification of adipogenesis using microplate reader. Stock solution of Nile red (1 mg/ml) in DMSO was prepared and stored at −20 °C protected from light. Staining was performed on unfixed cells. Cultured undifferentiated and differentiated cells (were grown in Corning polystyrene flat bottom 96-well TC-treated black microplates, Corning, NY, USA) were washed once with PBS. The dye was then added directly to the cells (5 μg/ml in PBS), and the preparation was incubated for 10 min at room temperature then washed twice with PBS. Fluorescent signal was measured using SpectraMax/M5 fluorescence spectrophotometer plate reader ( Molecular Devices Co., Sunnyvale, CA, USA) using bottom well-scan mode where nine readings were taken per well using Exc (485 nm) and Em (572 nm) spectra. Furthermore, fluorescence images were taken using FLoid cell imaging station (Life Technologies Inc., Grand Island, CA, USA).

ALP activity quantification. To quantify ALP activity in control and differentiated hMSC, we used the BioVision ALP activity colorimetric assay kit (BioVision, Inc., Milpitas, CA, USA) with some modifications. Cultures were collected in 96-well plates under normal or osteogenic induction conditions, then on day 10, wells were rinsed once with PBS and were fixed using 3.7% formaldehyde in 90% ethanol for 30 s at room temperature. Subsequently, fixative was removed and 50 μl of pNPP solution was added to each well and incubated for 1 h in the dark at room temperature. Reaction was subsequently stopped by adding 20 μl stop solution and gently shaking the plate. O.D. was then measured at 405 nm using SpectraMax/M5 fluorescence spectrophotometer plate reader.

RUNX2 quantification. For quantification of RUNX2 protein, hMSC were transfected with pre-miR-Neg or pre-miR-320c (30 nM), and 72 h later cells were collected and washed with PBS. Cells were lysed in 100 μl PBS containing protease inhibitors using five freeze-thaw cycles. Cell lysate was subsequently spun down at maximum speed for 10 min, and supernatant was collected and stored at −80 °C. Subsequently, RUNX2 was quantified using the RUNX2 ELISA kit according to the manufacturer’s recommendation (Uscn Life Science Inc., Wuhan, PRC).

AlamarBlue cell viability assay. Cell viability was measured using alamarBlue assay according to the manufacturer’s recommendations (AbD Serotec, Raleigh, NC, USA). In brief, we cultured cells in 96-well plates in 100 μl of the appropriate medium and at the indicated time point, and 10 μl of alamarBlue substrate was added and plates were incubated in the dark at 37 °C for 1 h. Reading was subsequently taken using fluorescent mode (Ex 530 nm/Em 590 nm) using BioTek Synergy II microplate reader (BioTek Inc., Winoski, VT, USA).

Statistics. Statistical analyses and graphing were performed using Microsoft excel 2010 and GraphPad Prism 6.0 software (Graphpad software, San Diego, CA, USA). P-values were calculated using the two-tailed t-test.

Conflict of Interest
The authors declare no conflict of interest.

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