Identifying deer antler *uhrf1* proliferation and *s100a10* mineralization genes using comparative RNA-seq

Dai Fei Elmer Ker¹, Dan Wang², Rashmi Sharma³, Bin Zhang¹, Ben Passarelli⁴, Norma Neff³, Chunyi Li⁵, William Maloney¹, Stephen Quake³,⁶,⁷ and Yunzhi Peter Yang¹,³,⁸*

**Abstract**

**Background:** Deer antlers are bony structures that re-grow at very high rates, making them an attractive model for studying rapid bone regeneration.

**Methods:** To identify the genes that are involved in this fast pace of bone growth, an in vitro RNA-seq model that paralleled the sharp differences in bone growth between deer antlers and humans was established. Subsequently, RNA-seq (> 60 million reads per library) was used to compare transcriptomic profiles. Uniquely expressed deer antler proliferation as well as mineralization genes were identified via a combination of differential gene expression and subtraction analysis. Thereafter, the physiological relevance as well as contributions of these identified genes were determined by immunofluorescence, gene overexpression, and gene knockdown studies.

**Results:** Cell characterization studies showed that in vitro-cultured deer antler-derived reserve mesenchyme (RM) cells exhibited high osteogenic capabilities and cell surface markers similar to in vivo counterparts. Under identical culture conditions, deer antler RM cells proliferated faster (8.6–11.7-fold increase in cell numbers) and exhibited increased osteogenic differentiation (17.4-fold increase in calcium mineralization) compared to human mesenchymal stem cells (hMSCs), paralleling in vivo conditions. Comparative RNA-seq identified 40 and 91 previously unknown and uniquely expressed fallow deer (FD) proliferation and mineralization genes, respectively, including *uhrf1* and *s100a10*. Immunofluorescence studies showed that *uhrf1* and *s100a10* were expressed in regenerating deer antlers while gene overexpression and gene knockdown studies demonstrated the proliferation contributions of *uhrf1* and mineralization capabilities of *s100a10*.

**Conclusion:** Using a simple, in vitro comparative RNA-seq approach, novel genes pertinent to fast bony antler regeneration were identified and their proliferative/osteogenic function was verified via gene overexpression, knockdown, and immunostaining. This combinatorial approach may be applicable to discover unique gene contributions between any two organisms for a given phenomenon-of-interest.

**Keywords:** Deer antler, Bone regeneration, Comparative RNA-seq, *uhrf1*, *s100a10*
Background

Deer antlers can account for 28% of the skeletal body weight [1, 2] and are the only known example of a mammalian tissue that regenerates rapidly, easily producing 10 kg or more of bone tissue within a relatively short period of 2 to 3 months [1–7]. Although deer antlers and human bone develop via intramembranous and endochondral modes of ossification [3, 4, 6, 7], deer antlers can grow up to 2 cm per day [3], which sharply contrasts with human femur bone, which grows at 2 cm per year during puberty [8]. Thus, if the molecular components that are involved in this process can be elucidated, this knowledge is expected to advance our understanding of the mammalian bone regeneration and holds promise for rapidly generating large bone volumes for skeletal tissue engineering.

Despite previous efforts [3, 4, 6, 7, 9–20], genes involved in fast antler regeneration remain poorly studied, and functional demonstration of their role(s) in proliferation and bone differentiation is lacking. For example, prior transcriptomic studies of deer antler tissues employed mouse microarrays [12], cDNA-amplified fragment length polymorphism (cDNA-AFLP) [15], or RNA-seq [18–20], but efforts to identify and characterize gene contributions to rapid deer antler growth have been hindered by logistical and technical issues. Such limitations include cross-species hybridization [12], sequence variation [21], the overwhelming number of candidates in transcriptomic datasets [12, 18–20], and the presence of complex spatial and temporal variables among tissue samples [12, 15, 18–20]. Indeed, one of the foremost challenges when working with transcriptomics datasets is determining which detected genes play a role in proliferation, skeletal differentiation, or a completely unrelated cellular process. Thus, previous transcriptomic studies to date have been limited to gene expression profiling with little-to-no characterization of gene contribution to antler proliferation or differentiation.

In this study, we hypothesized that an in vitro comparison of fallow deer (FD) and human RNA-seq data could circumvent several of the aforementioned challenges in identifying FD antler proliferation and mineralization genes. This was based on our reasoning that the fast pace of deer antler regeneration must stem, at least in part, from the rapid proliferation and differentiation of skeletal progenitors. Thus, a comparison of differentially expressed genes obtained from FD- or human-derived cells cultured under identical control and treatment group conditions would eliminate complex in vivo spatial and temporal variables while simplifying bioinformatics analysis to identify proliferation and mineralization gene candidates uniquely expressed in deer (Additional file 1: Figure S1; Additional file 2). Human mesenchymal stem cells (hMSCs) were selected to be compared against as they are a clinically promising therapeutic target for cell-based regenerative medicine [22]. In deer, antlers regenerate from skeletal progenitors found in the periosteum of the cranial appendages called pedicles (pedicle periosteum; PP), which, in turn, generate reserve mesenchyme (RM) cells of the growing antler tip that eventually differentiate into chondrocytes and mineralizing bone cells (Additional file 1: Figure S2) [3, 4, 6, 7]. Based on this, skeletal progenitors were collected from fallow deer (FD) facial periosteum (FP; as a non-antler-derived control), PP, and RM tissues to determine a suitable cell type for comparison with hMSCs. Subsequently, RNA-seq was performed, and genes whose expression were unique to FD antler progenitors were identified based on subtraction analysis between FD and human datasets. Lastly, the function and physiological relevance of these identified genes were determined by immunofluorescence, gene overexpression, and gene knockdown studies.

Results

Characterization of deer antler-derived cells
In this study, cell characterization studies were performed on FD- and human-derived skeletal progenitors to establish an in vitro model for comparative RNA-seq. Of the cells collected, RM cells exhibited the highest alkaline phosphatase (ALP) activity in the presence of osteogenesis-inducing bone morphogenetic protein-2 (BMP-2; Fig. 1a) while surprisingly, FP and PP cells cultured under mineralization conditions with BMP-2 and dexamethasone for 24 days exhibited little-to-no positive Alizarin Red S staining for calcium deposits (Fig. 1b). Also, fluorescence-activated cell sorting (FACS) analysis and immunofluorescence staining indicated that in vitro-cultured RM cells were similar to their in vivo counterparts [16, 17], with a majority (99.3%) of RM cells expressing ALP and a small subset (5.2%) expressing STRO-1 (Fig. 1c). Together, these studies indicated that in vitro-cultured RM cells were ideal models of deer antler skeletal progenitors for comparison with hMSCs.

Establishment of an in vitro model that compares deer antler-derived RM cells and hMSCs
Prior to performing RNA-seq, it was necessary to characterize in vitro-cultured RM cells and hMSCs to determine if there is sharply differential osteogenesis in vitro which may reflect the significant bone growth difference in vivo. Successful demonstration of this phenomenon in vitro would thus justify its use for comparative RNA-seq. Proliferation studies involving cells grown in three different mammalian culture media for 6 days demonstrated that RM cells exhibited increased growth relative to hMSCs. RM cells yielded 10.7–45.3 × 10⁴ cells with a doubling time of 17.9–24.7 h whereas hMSCs yielded 1.2–3.9 × 10⁵ cells with a doubling time of 37.8–60.1 h (Fig. 2a; Bartmann et al. [23] and Schallmoser et al.
As such, RM cells yielded 8.6–11.7-fold increase in cell numbers and 2.1–2.4-fold less doubling time compared to hMSCs when controlled for medium formulation (Fig. 2a). Cell cycle analysis showed similar results with a larger percentage of RM cells undergoing cell division relative to hMSCs (Fig. 2b). Differentiation studies...

**Fig. 1** Characterization of in vitro-cultured FD-derived cells. 

**a** FP and RM cells (isolate 2) cultured with 100 ng/mL BMP-2 for 6 days exhibited increased ALP activity relative to their respective control whereas PP cells (isolate 2) did not. Semi-quantification of ALP activity in FP, PP, and RM cells. 

**b** FP and PP cells (isolate 2) cultured with 100 ng/mL BMP-2 and 100 nM dexamethasone for 24 days did not exhibit increased Alizarin Red S staining relative to their respective control. 

**c** FACS analysis of RM cells. The percentage of cells that were negative for STRO1 and ALP, negative for STRO1 but positive for ALP, negative for ALP but positive for STRO1, and positive for both STRO1 and ALP were 0.28–1.25%, 91.83–97.53%, 0.004–0.10%, and 1.20–7.89%, respectively.

Data were from n = 3 isolates (three independent experiments with nine replicates per isolate for ALP and mineralization studies and one independent experiment with three replicates per isolate for FACs studies). Gray circles indicate observed data points. Error bars indicate standard error of mean or SEM. Statistical significance as indicated (Ker et al. Stem Cell Research & Therapy 2018:9:292, Page 3 of 15).
demonstrated that RM cells did not undergo adipogenic differentiation but underwent chondrogenesis and osteogenesis (Fig. 2c–e). In osteogenic gene expression studies, RM cells (isolate 2) cultured in the presence for BMP-2 for 6 days showed high upregulation of typical osteogenic genes such as alkaline phosphatase (alp), osteocalcin (ocn), osteoblast-specific factor-1 (osf-1), and runt-related transcription factor-2 (runx2) relative to control by 651.1-fold, 1.9-fold, 42.8-fold, and 557.5-fold, respectively (Fig. 2d). In osteogenic mineralization studies, control
groups for RM cells and hMSCs (cultured in the absence of BMP-2 and dexamethasone) exhibited 28.0 μg/mL and 39.6 μg/mL. Alizarin Red S stain, respectively, whereas treatment groups for RM cells and hMSCs (cultured in the presence of BMP-2 and dexamethasone) exhibited 1702.2 μg/mL and 98.0 μg/mL. Alizarin Red S stain, respectively (Fig. 2e).

Remarkably, even though identical culture conditions were used, RM cells and hMSCs demonstrated a 62.3-fold and 1.9-fold increase, respectively, in calcium mineralization relative to their respective controls, with RM cells demonstrating 17.4-fold increased levels of calcium mineralization compared to hMSCs despite similar baseline levels in the control groups (Fig. 2e). In addition, RM cells showed 2.0–18.1-fold increase over hMSC-mediated calcium mineralization despite a 5-fold decrease in BMP-2 concentration, as well as 3.1-fold increase in calcium mineralization over hMSC-like C3H10T1/2 cells despite a 1.6-fold longer cell doubling time (Fig. 2; Additional file 1: Figure S3; Ker et al. [25]). Thus, the sharply contrasting proliferation and mineralization capabilities of RM cells and hMSCs justified the use of this in vitro model and were expected to identify genes that orchestrate the differential growth and mineralization rates observed between deer antlers and human bone.

**Identification of deer antler proliferation and mineralization genes using comparative RNA-seq**

To identify the proliferation and mineralization genes that contribute to the fast growth and differentiation observed in our in vitro model, we focused our attention on uniquely expressed genes in RM cells. This was achieved by independently comparing RNA-seq data of RM cells (isolate 2) and hMSCs (isolate 24268) under proliferation (control group, 0% serum; treatment group, 10% serum) as well as mineralization (control group, 0 nM dexamethasone; treatment group, 100 nM BMP-2 and 0 nM dexamethasone; treatment group, 100 nM BMP-2 and 100 nM dexamethasone) conditions (Figs. 3 and 4). RNA-seq proliferation samples were sequenced to 76,781,962–99,716,096 reads per library with replicates showing a strong correlation of gene expression under serum-free and serum-containing conditions (Fig. 3a and Additional file 1: Table S1). As expected of a non-model organism, a larger percentage of unannotated genes was present in FD (36%) compared to human (8%) RNA-seq data (Fig. 3a). Despite this, Ingenuity Pathway Analysis (IPA) of annotated transcripts showed similar activation of proliferation-associated pathways such as *mitotic roles of polo-like kinase* as well as the expression of typical proliferation genes such as *cdk1, rrm1, cdc7, aurka*, and *plk4* in both datasets (Fig. 3a). Correspondingly, gene ontology analysis showed upregulation of the processes associated with proliferation including mitotic checkpoints and chromosome condensation (Fig. 3b). Also, RNA-seq mineralization samples were sequenced to 62,601,720–86,750,048 reads per library with replicates showing a strong correlation of gene expression under non-mineralization and mineralization conditions (Fig. 4a and Additional file 1: Table S2). Similar to the proliferation dataset, a larger percentage of unannotated genes was present in FD (41%) compared to human (13%) RNA-seq data (Fig. 4a). IPA of annotated transcripts showed similar activation of osteogenic-associated pathways such as *roles of osteoblasts, osteoclasts, and chondrocytes in rheumatoid arthritis* as well as the expression of typical osteogenic genes such as *dlx5, tsc22d3, alpl, klf4, ext1*, and *stc1* in both datasets (Fig. 4a). Correspondingly, gene ontology analysis showed upregulation of processes associated with skeletal catabolism including collagen synthesis as well as face and body morphogenesis (Fig. 4b). Subsequently, subtraction analysis was performed between human and FD datasets for differentially expressed genes. Using the following criteria of highly upregulated (>5-fold) and uniquely expressed FD genes, 40 proliferation and 91 mineralization candidate genes were identified (Figs. 3a and 4a). Thus, in vitro comparative RNA-seq identified gene candidates that were uniquely expressed in RM cells with a presumed role in rapid deer antler regeneration.

**Validation of in vitro comparative RNA-seq**

To validate the physiological relevance and role of uniquely expressed genes identified by in vitro comparative RNA-seq, a proliferation and mineralization gene candidate was each selected and further assessed in immunofluorescence, gene overexpression, and gene knockdown studies (Figs. 5 and 6). Since chemical- and electroporation-mediated transfections of hMSCs were unsuccessful (data not shown), hMSC-like mouse C3H10T1/2 cells were used in gene overexpression studies.

Of the 40 proliferation gene candidates, FD *uhrf1* was chosen due to its role in epigenetic inheritance [26] and high expression in several cancers [27], suggesting a role for this gene in simultaneously controlling stem cell self-renewal [28] and growth in deer antlers. In immunofluorescence studies, regenerating FD antlers obtained from an independent herd showed UHRF1 expression in RM tissue (Fig. 5a) while supplementation of known mitogens such as fibroblast growth factor-2 (FGF-2) and insulin-like growth factor-1 (IGF-1) in RM cells showed a good correlation between UHRF1 expression and RM cell proliferation (Additional file 1: Figure S4). For example, RM cells exhibited increased UHRF1 expression relative to control when cultured in the presence of IGF-1 under serum-free conditions as well as in the presence of FGF-2 under both serum-free and serum-containing conditions (Additional file 1: Figure S4a and S4b). Correspondingly, increased proliferation was only observed under the aforementioned culture conditions but not in the presence of IGF-1 under serum-containing conditions, where there was no upregulation of UHRF1 (Additional file 1: Figure S4c). Together, these data demonstrate the physiological
Fig. 3 RNA-seq analysis of RM cells and hMSCs under proliferation and mineralization conditions. a RNA-seq analysis of RM cells (isolate 2) and hMSCs (isolate 24268) under serum-free (0% serum) and serum-containing (10% serum) conditions identified 40 candidate proliferation genes. Scatterplots indicate the correlation ($r^2$) between replicates for each condition. FPKM, fragments per kilobase of transcript per million mapped reads. b Gene ontology enrichment analysis of RM cells (isolate 2) and hMSCs (isolate 24268) under proliferation conditions. Graphs indicate the top 5 upregulated (red) and downregulated (green) biological processes, cellular components, and molecular functions. Data were from $n = 1$ isolate (one independent experiment with two replicates per group).
Fig. 4 RNA-seq analysis of RM cells and hMSCs under proliferation and mineralization conditions. a RNA-seq analysis of RM cells (isolate 2) and hMSCs (isolate 24268) under control (0 ng/mL BMP-2 and 0 nM dexamethasone) and osteogenic (100 ng/mL BMP-2 and 100 nM dexamethasone) conditions identified 91 candidate mineralization genes. Scatterplots indicate the correlation ($r^2$) between replicates for each condition. FPKM, fragments per kilobase of transcript per million mapped reads. b Gene ontology enrichment analysis of RM cells (isolate 2) and hMSCs (isolate 24268) under mineralization conditions. Graphs indicate the top 5 upregulated (red) and downregulated (green) biological processes, cellular components, and molecular functions. Data were from $n = 1$ isolate (one independent experiment with two replicates per group).
relevance of *uhrf1* in FD RM cell proliferation and verified the expression pattern of this gene relative to our RNA-seq proliferation data (Fig. 5a and Additional file 1: Figure S4). In gene knockdown studies, RM cells (isolate 2) treated with control siRNA yielded $5.1 \times 10^4$ cells, whereas groups treated with siRNAs yielded $2.8 - 4.2 \times 10^4$ cells (Fig. 5b and Additional file 1: Figure S5a). As such, siRNA-mediated knockdown of *uhrf1* inhibited RM cell growth of 17.9–45.5% (Fig. 5b and Additional file 1: Figure S5a). In gene overexpression studies, C3H10T1/2 cells stably transfected with *uhrf1* yielded $13.6 \times 10^4$ cells with a doubling time of 11.8 h (Fig. 5c). As such, overexpression of FD *uhrf1* in C3H10T1/2 cells (Additional file 1: Figure S5b) increased cell proliferation by 1.49–1.76-fold and
Fig. 6 Identification of s100a10 as a uniquely expressed mineralization gene using in vitro comparative RNA-seq. a S100A10 immunofluorescence staining in regenerating deer antler tissue. b RM cells (isolate 2) cultured with 100 ng/mL BMP-2 and 100 nM dexamethasone exhibited increased S100A10 expression relative to control. c C3H10T1/2 cells stably transfected with s100a10 and cultured with 100 ng/mL BMP-2 for 4 h exhibited increased alp gene expression relative to untransfected control and empty plasmid control. C3H10T1/2 cells stably transfected with s100a10 and cultured with 100 ng/mL BMP-2 for 12 days exhibited increased ocn and runx2 gene expression relative to their respective control. d C3H10T1/2 cells stably transfected with s100a10 and cultured with 100 ng/mL BMP-2 for 4 days exhibited increased ALP activity relative to untransfected control. e C3H10T1/2 cells stably transfected with s100a10 and cultured in the presence of 100 ng/mL BMP-2 and 100 nM dexamethasone exhibited increased Alizarin Red S staining relative to untransfected control and empty plasmid control. Scale bars as indicated. Data were from n = 3 isolates (an independent herd for antler immunofluorescence studies), n = 2–3 independent experiments with 4–10 replicates per group for osteogenic gene expression studies, n = 3 independent experiments with 9 replicates per group for s100a10 overexpression ALP studies, and n = 5 independent experiments with 15 replicates per group for s100a10 overexpression mineralization studies. Gray circles indicate observed data points. Error bars indicate SEM. Statistical significance as indicated.
decreased cell doubling time by 13.8–15.6% (Fig. 5c) without compromising contact inhibition (Fig. 5c and Additional file 1: Figure S6) or ALP activity (Fig. 5e). This is particularly noteworthy given that untransfected C3H10T1/2 cells already grow rapidly and double every 13.9–15.1 h (Fig. 5c and Ker et al. [25]). Thus, in vitro comparative RNA-seq identified uhrf1 as contributing towards a previously unknown role in FD antler cell proliferation.

Of the 91 mineralization gene candidates, FD s100a10, which has roles in fibrinolysis and intracellular membrane organization [29], was chosen due to the paucity of data regarding its role in osteogenesis, making it as a novel target for further study. In addition, members of the S100 family of proteins such as S100A4 have been reported to be a negative regulator of osteoblast differentiation and matrix mineralization [30]. In immunofluorescence studies, regenerating FD antlers obtained from an independent herd showed S100A10 expression in the cartilage regions of the antler undergoing mineralization (Fig. 6a) while S100A10 was upregulated during osteogenic differentiation of RM cells (Fig. 6b). Together, these data demonstrate the physiological relevance of s100a10 in FD mineralization and verified the expression pattern of this gene relative to our RNA-seq mineralization data (Fig. 6a, b). In gene overexpression studies, C3H10T1/2 stably transfected with s100a10 (Additional file 1: Figure S7) showed increased alp expression (Fig. 6c) and ALP activity (Fig. 6d) relative to empty plasmid and untransfected controls as well as upregulation of other typical osteogenic genes ocn and runx2 relative to control when cultured in the presence of BMP-2 (Fig. 6c). In addition, C3H10T1/2 stably transfected with s100a10 exhibited increased calcium mineralization relative to untransfected and empty plasmid controls when in the presence of BMP-2 and dexamethasone (Fig. 6e).

Under these culture conditions, C3H10T1/2 cells, C3H10T1/2 cells stably transfected with empty plasmid, and C3H10T1/2 cells stably transfected with s100a10 exhibited 257.2 μg/mL, 341.9 μg/mL, and 807.9 μg/mL Alizarin Red S stain, respectively (Fig. 6e). As such, overexpression of FD s100a10 in C3H10T1/2 cells increased cell-mediated mineralization by 2.4–3.1-fold (Fig. 6e). Thus, in vitro comparative RNA-seq identified s100a10 as contributing towards a previously unknown role in FD antler cell mineralization. Together, these results demonstrated the capability of in vitro comparative RNA-seq analysis to identify uniquely expressed FD proliferation and mineralization genes.

**Discussion**

In this study, deer antler-derived RM cells and human bone marrow-derived mesenchymal stem cells were used to establish an in vitro model for comparative RNA-seq (Additional file 1: Figures S1, S2, and S3; Figs. 1 and 2) and identified uhrf1 and s100a10 as uniquely expressed deer antler proliferation (Figs. 3 and 5 and Additional file 1: Figures S4, S5, and S6) and mineralization genes (Figs. 4 and 6 and Additional file 1: Figure S7), respectively. The approach developed here may be broadly applied towards studying another biological phenomenon, and the genes identified with this approach will not only advance our understanding of mammalian bone regeneration but also offer promising therapeutic strategies for bone tissue engineering.

Our premise for using an in vitro-based approach was based on several reasons. First, an in vitro approach enabled greater experimental control by allowing culture conditions to define and study the phenomenon-of-interest. Second, the use of identical conditions for culturing deer and human cells not only eliminated complex in vivo spatial and temporal variables but also allowed differential gene expression data from each species to serve as a basis for comparison for identifying uniquely expressed deer antler genes via a simple subtraction analysis. For example, serum-free and serum-containing media conditions were used to identify proliferation genes via differential gene expression. These experiments are independently performed for human and deer cells. Thereafter, subtraction analysis between human and deer proliferation genes would yield uniquely expressed deer proliferation genes. In addition, an in vitro approach reduced the logistical burden for long-term housing of a large, non-model organism such as deer while RNA-seq enabled comprehensive and sensitive detection of transcripts [31] with little bias and error even when non-target species reference genomes are used [32]. Thus, an in vitro approach was expected to identify uniquely expressed deer antler proliferation and mineralization genes.

In establishing this model to identify deer antler proliferation and mineralization genes, hMSCs and RM cells were compared. The basis underlying this choice stems from the significant differences in bone growth between human skeletal and deer antler tissues [3, 4, 6–8] as well as and the therapeutic promise of recapitulating such rapid growth in human skeletal tissues. Although hMSCs and RM cells possess different genetic backgrounds and they do not originate from anatomically equivalent tissues (hMSCs were obtained from the iliac crest whereas RM cells were harvested from the cranial pedicle), a transcriptomic comparison of these cells is still expected to provide important insights as to the genes necessary to stimulate fast proliferation and high mineralization, particularly since hMSCs are a clinically promising therapeutic target for bone tissue engineering [22, 33].

In addition, in vitro-cultured RM cells exhibited high osteogenic capability (Additional file 1: Figure S3, Figs. 1 and 2) and similar cell surface markers (ALP and STRO-1) as their in vivo counterparts [16, 17]. When compared to hMSCs under identical culture conditions, RM cells demonstrated 8.6–11.7-fold increased cell growth and 17.4-fold
increased levels of calcium mineralization (Fig. 2), to some extent reflecting the rapid growth and differentiation phenomena observed in regenerating deer antlers. Together, these studies justified the use of in vitro cultured hMSCs and RM cells to identify deer antler proliferation and mineralization genes.

To identify deer antler proliferation and mineralization genes, RNA-seq was performed. Comparison and subsequent subtraction analysis of transcriptomes under proliferation and mineralization conditions identified 40 proliferation and 91 mineralization genes that were uniquely expressed in RM cells (Fig. 3 and Fig. 4). Further bioinformatics analysis using commercial and public databases showed activation as well as enrichment of proliferation and mineralization pathways or keywords, concurring with the culture conditions employed (Figs. 3 and 4). To validate the contribution of identified genes, uhrf1 and s100a10 were chosen for further study based on their potential role in stem cell renewal [28] or novelty, respectively. Although their participation in deer antler biology have not been reported, other studies have indicated that UHRF1 is involved in the proliferation and maturation of colonic Treg cells via epigenetic silencing of CDKNAP21, an inhibitor of cyclin/cyclin-dependent kinase complexes [34] while related members of the S100 family of proteins such as S100A4 are involved in negative regulation of osteoblast differentiation [30]. Overexpression of FD uhrf1 in a mouse cell line with hMSC-like characteristics [25] increased cell proliferation without affecting contact inhibition (Fig. 5, Additional file 1: Figures S5 and S6) whereas siRNA-mediated knockdown of uhrf1 in RM cells decreased cell growth (Fig. 5 and Additional file 1: Figure S5). Similarly, overexpression of FD s100a10 in this mouse cell line increased osteogenic gene expression, ALP activity, and calcium mineralization (Fig. 6 and Additional file 1: Figure S7). In addition, the physiological relevance of these results was confirmed by immunofluorescence staining of regenerating deer antlers (Figs. 5 and 6). Thus, in vitro comparative RNA-seq identified deer antler proliferation and mineralization genes.

The success of in vitro comparative RNA-seq approach is heavily dependent upon several factors. First, it is necessary to use appropriate in vitro culture conditions that closely model the biological phenomenon-of-interest. In this study, we demonstrated that RM cells rapidly proliferated and exhibited increased calcium mineralization levels relative to hMSCs, which mimicked the phenomenon of rapid bone growth in deer antlers (Fig. 2). However, it is possible that the culture conditions do not accurately reflect growth and differentiation stimuli in regenerating deer antlers, resulting in either false positives or negatives. As such, it is vital to ascertain the physiological relevance of these results by determining gene or protein expression in the relevant biological tissue (Figs. 5 and 6). Despite these limitations, our approach was successful in identifying uhrf1 and s100a10 as previously unknown FD antler proliferation and mineralization genes, respectively. However, additional modifications to this study may extend its impact further. These include alternative culture conditions that better mimic physiological conditions such as the use of co-cultures to study antler bone and velvet (skin) paracrine interactions as well as the additional application of more stringent selection criteria such as comparison of in vitro (our current study) and in vivo (deer antler tissue) RNA-seq datasets. Second, it is important to recognize that in vitro comparative RNA-seq performs subtraction analysis between differentially expressed RM genes and its corresponding set of differentially expressed hMSC genes. As such, it is possible that genes which are vital in deer antler proliferation and/or mineralization but are not differentially expressed will not be detected. To address this, our study adopted a strategy of limited biological replication (one isolate, two replicates) with a high number of sequencing reads. This strategy relies on utilizing a large number of sequencing reads to generate increased statistical power for sensitive detection of differential gene expression [35]. Such a strategy would be particularly important for discovering novel, low copy transcripts. Indeed, this approach has been successful in identifying a large number of previously undetected estrogen-related transcripts in breast cancer cells [36]. Third, since the deer genome was only recently sequenced [37], sequencing reads were mapped onto the closely related and well-annotated Bos taurus genome. Although such cross-species mapping can result in sequence and expression data loss as well as increased bias and error, these effect sizes are reported to be small within a 100-million-year window and exhibit better mapping performance relative to de novo transcriptome assembly [32]. Despite this, our deer RNA-seq datasets do contain a large percentage of unannotated genes, and this result may be improved by applying orthology-driven Blast mapping.

**Conclusion**

In conclusion, we have developed an in vitro model for comparative RNA-seq between FD RM cells and hMSCs to simplify analysis of transcriptomic datasets and for the first time to identify unique genes pertinent to deer antler regeneration. The discovery of these genes advances our understanding of deer antler biology and offer promising strategies for rapid bone regeneration. We envisage a similar comparison strategy can be applied to almost any tissue for identifying the contributions of uniquely expressed genes to a phenomenon of interest.

**Methods**

Detailed materials and methods are provided in Additional file 1: Supplementary Information.
Fallow deer
Tissues were harvested from Lazy Arrow Camatta Ranch (Santa Margarita, CA) and Walking Beam Ranch (Santa Paula, CA) in accordance with approved guidelines established by Stanford University’s Administrative Panel on Laboratory Animal Care (APLAC 28057). Antler tissue was identified [14] and harvested using enzymatic digestion.

Cell culture
Mouse C3H10T1/2 mesenchymal fibroblasts (American Type Culture Collection; ATCC, Manassas, VA) were maintained in DMEM, 10% FBS, and 1% P/S and used within the first ten passages from the date of receipt. FD cells were maintained in DMEM, 10% FBS and 1% P/S and used between passages 2 and 8. Human mesenchymal stem cells (hMSCs; Lonza, Switzerland) were maintained according to the manufacturer’s instructions and used between passages 4 and 7. Hoechst staining (Anaspec, Fremont, CA) was used to monitor mycoplasma contamination in cell cultures.

Cell proliferation
Cell proliferation studies were performed in DMEM, 10% FBS, 1% P/S, mesenchymal stem cell growth media (Lonza, Switzerland), and mesenchymal stem cell growth media supplemented with 10 ng/ml fibroblast growth factor-2 (FGF-2; Peprotech, Rocky Hill, NJ). Cells were counted using a Beckman Coulter Z2 Particle Counter (Beckman Coulter; Brea, CA), and cell doubling times were calculated using R-studio (R Studio, Boston, MA, http://www.rstudio.com). Cell cycle studies were performed using propidium iodide/RNAse solution (Cell Signaling Technology, Danvers, MA) on a BD Aria II flow cytometer and analyzed using Flowjo 9.7.5 (Flowjo LLC, Ashland, OR, http://www.flowjo.com).

Cell differentiation
Adipogenic and chondrogenic differentiation were performed using StemPro Adipogenic (Gibco, Thermo Fisher Scientific, Waltham, MA) and StemPro Chondrogenic Media (Gibco, Thermo Fisher Scientific, Waltham, MA), respectively, according to the manufacturer’s instructions. Osteogenic differentiation was performed using either DMEM, 10% FBS, 1% PS, and 100 ng/ml BMP-2 (Infuse Bone Graft, Medtronic, Sunnyvale, CA) or DMEM, 10% FBS, 1% P/S, 50 μg/ml ascorbic acid, 10 mM β-glycerophosphate, 100 ng/ml BMP-2, and 100 nM dexamethasone (Sigma Aldrich, St. Louis, MO). Osteogenic gene expression was performed on cDNA templates (RNA isolation: Qiagen RNeasy Plus Mini kit, Qiagen, Germany; Reverse transcription: Omniscript kit, Qiagen, Germany) for 40 cycles using TaqMan Gene Expression Mastermix (4369016, Applied Biosystems, Thermo Fisher Scientific, Waltham, MA) on an Applied Biosystems HT7200 thermocycler (Applied Biosystems, Thermo Fisher Scientific, Waltham, MA). Gene expression data were analyzed using SDS 2.2.2 (Applied Biosystems, Thermo Fisher Scientific, Waltham, MA, http://www.thermofisher.com/). ALP activity (Kit 86C, Sigma Aldrich, St. Louis, MO) was detected according to the manufacturer's instructions. Where necessary, the average pixel intensity was determined using the image histogram tool in Adobe Photoshop as previously described [38, 39]. Osteogenic mineralization was determined using 2% Alizarin Red S stain (Electron Microscopy Sciences, Hatfield, PA) on solvent-extracted samples. Absorbance values from the samples and standards were read at 405 nm using a Tecan Infinite F50 spectrometer (Tecan Trading AG, Switzerland).

RNA-seq
Proliferation libraries were obtained from cells cultured under DMEM, 0% FBS, and 1% P/S (0% serum) and DMEM, 10% FBS, 1% P/S (10% serum) while mineralization libraries were obtained from cells cultured under DMEM, 10% FBS, 1% P/S, 50 μg/ml ascorbic acid, and 10 mM β-glycerophosphate (control media; without BMP-2 and dexamethasone) and DMEM, 10% FBS, 1% P/S, 50 μg/ml ascorbic acid, 10 mM β-glycerophosphate, 100 ng/ml BMP-2, and 100 nM dexamethasone. RNA was isolated (Qiagen RNeasy Plus Mini kit, Qiagen, Germany), reverse-transcribed into cDNA (Ovation RNA-seq System V2 kit, NuGEN, San Carlos, CA), sheared (S2 Focused-ultrasonicator, Covaris, Woburn, MA), end-repaired, dA-tailed, ligated with custom adaptors, and PCR-amplified for RNA-seq library construction (NEBNext DNA Library Prep Master Mix Set for Illumina, New England Biolabs, Ipswich, MA). Samples were sequenced using 100 base-pair, paired-end RNA-seq technology (HiSeq 2000, Illumina, San Diego, CA), and data were analyzed using several bioinformatics software including Spliced Transcripts Alignment to a Reference (STAR; Version 2.3.0, https://code.google.com/p/rsa-star/) [40], SAMtools (Version 0.1.19 http://www.htslib.org/), the Cufflinks package (Version 2.1.1.1, https://github.com/cole-trapnell-lab/cufflinks) [41], the Cummerbund package (http://combio.mit.edu/cummeRbund/), Ingenuity Pathway Analysis (Qiagen, Germany; https://www.qiagenbioinformatics.com/), and Gene Ontology Enrichment Analysis (http://geneontology.org/page/go-enrichment-analysis). Statistical analyses for RNA-seq were performed as previously described [41]. Gene candidates were identified based on differentially expressed genes that exhibited more than fivefold upregulation in control versus treatment conditions as well as genes that were uniquely expressed in the FD RM cell dataset. From this set of candidates, deer uhrf1 and s100a10 genes were manually identified as genes-of-interest based on a literature search (Pubmed; http://www.pubmed.com) of their known biological functions and novelty within the context of mammalian bone biology.
Gene cloning and transfection
Genes were PCR-cloned (Platinum Blue PCR SuperMix, Invitrogen, Thermo Fisher Scientific, Waltham, MA) from deer cDNA libraries using primers designed from the closely related bovine genome prior to subcloning into a pVitró2-MCS-Blast plasmid (InvivoGen, San Diego, CA). C3H10T1/2 cells were transfected with 2–3 μg plasmid(s) containing the gene-of-interest according to the manufacturer’s instructions (Polyplus, France), and stably transfected cells were selected using 3 μg/mL blasticidin (Invitrogen, Thermo Fisher Scientific, Waltham, MA).

siRNA
Cells were transfected with 30 nM uhrf1 siRNA A and E (siRNAs were custom-designed by Santa Cruz Biotechnology Inc., Dallas, TX, based on bovine uhrf1 sequence) according to the manufacturer’s instructions (Polyplus, France) for 72 h.

Immunofluorescence staining
Cells were fixed in 4% paraformaldehyde (Electron Microscopy Sciences, Hatfield, PA), blocked with 10% donkey serum (Jackson Immunoresearch Laboratories Inc., West Grove, PA), and incubated with primary antibody followed by secondary antibody incubation with appropriate washes in between. Antibodies included 10 μg/mL mouse anti-Stro-1 (MAB1038, R&D Systems Inc., Minneapolis, MN), 4 μg/mL rabbit anti-alkaline phosphatase (Sc-98652, ALP; Santa Cruz Biotechnology Inc., Dallas, TX), 10 μg/mL rabbit anti-UHRF1 (Sc98704, Santa Cruz Biotechnology Inc., Dallas, TX), 1 μg/mL mouse anti-S100A10 (Ab89438, Abcam Inc., Cambridge, MA), 15 μg/mL donkey anti-mouse Alexa 488 (715-545-150, Jackson Immunoresearch Laboratories Inc., West Grove, PA), and 15 μg/mL donkey anti-rabbit Alexa 647 (711-605-152, Jackson Immunoresearch Laboratories Inc., West Grove, PA). Where necessary, antigen retrieval was performed using antigen retrieval buffer solution (IHC World LLC, Woodstock, MD) at 80–90 °C for 30–60 min prior to antibody incubation. Average pixel intensity was determined using the image histogram tool in Adobe Photoshop as previously described [38, 39].

Statistical analysis
Statistical analyses involving RNA-seq were performed by Cufflinks and R-Studio [41]. Statistical significance for differentially expressed genes was established at p ≤ 0.05 and q ≤ 0.05. Statistical analyses not involving RNA-seq were performed using IBM SPSS Statistics for Windows 22 (IBM Corp., North Castle, NY, http://www.ibm.com). Quantitative data was presented as means ± standard error of mean (mean ± SEM) where appropriate. Relative fold changes for PCR data were log transformed in order to make the data distribution more symmetrical [42]. The Shapiro-Wilk test and the Levene test were used to determine whether data were normally distributed and contained equal variances among groups, respectively. For two mean comparisons, p values were computed via the t test. p values were calculated using pooled and separate variance for data with equal and unequal variances, respectively. For more than two mean comparisons, p values were computed via analysis of variance (ANOVA). If the majority of the data were normally distributed or there was an equal variance among groups, p values were calculated using ANOVA followed by Tukey’s honest significant difference post hoc multiple comparison test [43, 44]. Otherwise, p values were calculated using Welch’s ANOVA followed by Games-Howell post hoc multiple comparison test [45]. Statistical significance was established at p ≤ 0.05.

Additional files

**Additional file 1**: Supplementary figures. (PDF 2760 kb)

**Additional file 2**: Comparison of Differentially Expressed Genes from Fallow Deer RM and Human Mesenchymal Stem Cells. (XLSX 13100 kb)

Abbreviations
ALP: Alkaline phosphatase; ANOVA: Analysis of variance; BMP-2: Bone morphogenetic protein-2; cDNA-AFLP: cDNA-amplified fragment length polymorphism; FACS: Fluorescence-activated cell sorting; FD: Fallow deer; FGF-2: Fibroblast growth factor-2; FP: Facial periosteum; hMSCs: Human mesenchymal stem cells; IGF-1: Insulin-like growth factor-1; IPA: Ingenuity Pathway Analysis; osn: Osteocalcin; oif-1: Osteoblast-specific factor-1; PP: Pedicle periosteum; RM: Reserve mesenchyme; runx2: Runt-related transcription factor-2; STAR: Spliced Transcripts Alignment to a Reference

Acknowledgements
We would like to thank Amy Lam and Michael Lin (Stanford University) for providing the mRuby2 gene. We would like to thank Felicia Morrison and her family at Lazy Arrow Ranch as well as Pete Busch and his family at Walking Beam Ranch for providing the fallow deer samples. We would like to thank Chris Brown and John Halford for the assistance with antler harvesting. We would like to thank Gary Mantelas for the assistance with RNA-seq studies. We would like to thank Joseph Wu for providing the manuscript edits. We would like to thank Ivan Cheng and Jill Helms for the generous use of their equipment and facilities.

Funding
This work was supported by NIH grants R01AR057837-06S1 Star Award (YPY), R01AR057387-02 (YPY), and R01DE021468 (YPY); Department of Defense (W81XWH-10-1-0966; YPY); Boswell Foundation (YPY); and AO Startup Grant (Project 5-13-134K; DFEK). Flow cytometry sorting and analysis were performed on an instrument in the Stanford Shared FACS Facility obtained using NIH S10 Shared Instrument Grant S10RR027431-01.

Availability of data and materials
The transcriptome sequencing data for deer and human cells has been deposited in the Gene Expression Omnibus database (https://www.ncbi.nlm.nih.gov/geo/) under accession number GSE89796.

Authors’ contributions
DFEK contributed to the experimental design, collection and assembly of data (antler harvesting and isolation, flow cytometry, cell doubling studies, cell differentiation studies, human and deer RNA-seq studies and bioinformatics, antler immunofluorescence studies, deer gene cloning and deer overexpression studies, siRNA-mediated deer gene knockdown studies and preparation of the figures), data analysis and interpretation, and manuscript writing. DW contributed...
to the experimental design, collection and assembly of data (cell doubling studies, cell differentiation studies, antler immunofluorescence studies, deer gene cloning, and deer overexpression studies, and preparation of the figures), data analysis and interpretation, and manuscript writing. RS contributed to the collection and assembly of data (flow cytometry, cell differentiation studies, human RNA-seq studies and bioinformatics, deer gene cloning, and deer overexpression studies) and data analysis and interpretation. BZ contributed to the collection and assembly of data (antler harvesting and isolation and cell differentiation studies) and data analysis and interpretation. BP contributed to the collection and assembly of data (RNA-seq bioinformatics) and data analysis and interpretation. NFN contributed to the collection and assembly of data (RNA-seq studies) and data analysis and interpretation. CL contributed to the data interpretation. WM and SQO contributed to the conception and experimental design and data analysis and interpretation. YPY contributed to the conception and experimental design, data analysis and interpretation, and manuscript writing. All authors read and approved the final manuscript.

Ethics approval
Ethics approval for the harvesting of deer antlers was obtained from Stanford University's Administrative Panel on Laboratory Animal Care (APLAC 28057).

Consent for publication
Not applicable.

Competing interests
The authors declare that they have no competing interests.

Publisher's Note
Springer Nature remains neutral with regard to jurisdictional claims in published maps and institutional affiliations.

Author details
1Department of Orthopaedic Surgery, Stanford University, 300 Pasteur Drive, Stanford, CA 94305, USA. 2Department of Stomatology, Tenth People's Hospital of Tongji University, 301 Yanchang Road, Shanghai 200072, China. 3Department of Bioengineering, Stanford University, Shriram Center 443 Via Ortega, Stanford, CA 94305, USA. 4Scientific Computing Core, Calico Life Sciences LLC, 1170 Veterans Blvd., South San Francisco, CA 94080, USA. 5State Key Lab for Molecular Biology of Special Economic Animals, 4899 Juyue Street, Changchun 130112, Jilin, China. 6Department of Applied Physics, Stanford University, 348 Via Pueblo Mall, Stanford, CA 94305, USA. 7Howard Hughes Medical Institute, 4000 Jones Bridge Road, Chevy Chase, MD 20815, USA. 8Department of Material Science and Engineering, Stanford University, 496 Lomita Mall, Stanford, CA 94305, USA.

Received: 15 May 2018 Revised: 24 August 2018 Accepted: 30 September 2018 Published online: 31 October 2018

References
1. Gómez IA, Ceacerio F, Landete-Castillejos T, Gaspar-López E, Garcia AJ, Gallego LF. Factors affecting antler investment in Iberian red deer. Anim Prod Sci. 2012;52:867–73.
2. Huxley JS. The relative size of antlers in deer. Proc Zool Soc London. 1931;101:819–64.
3. Goss RJ, Deer antlers. Regeneration, function and evolution. New York: Academic Press; 1983.
4. Kierdorf U, Kierdorf H. Deer antlers - a model of mammalian appendage regeneration: an extensive review. Gerontology. 2011;57:53–65.
5. Li C. Histogenetic aspects of deer antler development. Front Biosci. 2013;5:479–89.
6. Li C, Yang F, Sheppard A. Adult stem cells and mammalian epimorphic regeneration: insights from studying annual renewal of deer antlers. Curr Stem Cell Res Ther. 2009;4:237–51.
7. Price J, Faucheux C, Allen G. Deer antlers - a model of mammalian regeneration. Curr Top Dev Biol. 2005;67:1–48.
8. Sistrom HA, Kember NF. Longitudinal bone growth of the human femur. Postgrad Med J. 1973;50:433–7.
9. Allen SP, Maden M, Price JS. A role for retinoic acid in regulating the regeneration of deer antlers. Dev Biol. 2002;251:409–23.
10. Bartol C, Schams D, Bubenik GA. Testosterone, but not IGF-I-LH, prolactin or cortisol, may serve as antler-stimulating hormone in red deer stags (Cervus Elaphus). Bone. 2009;44:691–8.
11. Daley E, Allford A, Miller JD, Goldstein S. Phenotypic differences in white-tailed deer antlerogenic progenitor cells and marrow-derived MSC. Tissue Eng A. 2013;20:1416–25.
12. Guryanj I Jr, Molnár A, Borsy A, Stéger V, Hackler L, Jr, Zomborszky Z, Papp P, Duda E, Deák F, Lakatos P, et al. Gene expression dynamics in deer antler: mesenchymal differentiation toward chondrogenesis. Mol Gen Genomics. 2007;272:221–35.
13. Hu W, Li T, Hu R, Wu L, Li M, Meng X. MicroRNA let-7a and let-7f as novel regulatory factors of the sika deer (Cervus nippon) IGF-I gene. Growth Factors. 2014;32:227–33.
14. Li C, Clark DE, Lord EA, Stanton JA, Suttie JM. Sampling technique to discriminate the different tissue layers of growing antler tips for gene discovery. Anat Rec. 2002;268:125–30.
15. Molnár A, Guryanj I, Korpus E, Borsy A, Stéger V, Buza L, Kiss I, Zomborszky P, Deák F, Orosz L. Identification of differentially expressed genes in the developing antler of red deer Cervus elaphus. Mol Gen Genomics. 2007;277:237–48.
16. Price JS, Ouyajobi BO, Orefo RO, Russell RG. Cells cultured from the growing tip of red deer antler express alkaline phosphatase and proliferate in response to insulin-like growth factor-I. J Endocrinol. 1994;143:89–96.
17. Rolf HJ, Kierdorf U, Kierdorf H, Schulz J, Seymour N, Schlepake H, Napp J, Niebert S, Wolfel H, Wiese KG. Localization and characterization of STRO-1 cells in the deer pedicle and regenerating antler. PLoS One. 2008;3:e2064.
18. Yao B, Zhao Y, Wang G, Zhang M, Liu M, Liu H, Li J. De novo characterization of the antler tip of Chinese sika deer transcriptome and analysis of gene expression related to rapid growth. Mol Cell Biochem. 2012;364:130–9.
19. Yao B, Zhao Y, Zhang H, Zhang M, Liu M, Liu H, Li J. Sequencing and de novo analysis of the Chinese sika deer antler-tip transcriptome during the ossification stage using illumina Rna-Seq technology. Biotechnol Lett. 2012;34:813–22.
20. Zhao Y, Yao B, Zhang M, Wang S, Zhang H, Xiao W. Comparative analysis of differentially expressed genes in sika deer antler at different stages. Mol Biol Rep. 2012;40:1665–76.
21. Bae KM, Kwon YS, Cho IH, Yi SI. Use of cDNA-AFLP for transcript profiling in narrow genetic pools, for example, cucumber (Cucumis sativus L.). Plant Breed. 2006;125:488–92.
22. Mendicino M, Bailey AM, Wonnacott K, Puri RK, Bauer SR. MSC-based product characterization for clinical trials: an FDA perspective. Cell Stem Cell. 2014;14:141–5.
23. Battmann C, Rohde E, Schallmoser K, Purstner P, Lanzer G, Linkesch W, Strunk D. Two steps to functional mesenchymal stromal cells for clinical application. Transfusion. 2007;47:426–35.
24. Schallmoser K, Battmann C, Rohde E, Reinisch A, Kashofer K, Stadelmeyer E, Drexler C, Lanzer G, Linkesch W, Strunk D. Human platelet lysate can replace fetal bovine serum for clinical-scale expansion of functional mesenchymal stromal cells. Transfusion. 2007;47:1346–46.
25. Ker DFE, Sharma R, Wang ETH, Yang YP. Development of milky2-transfected C3H10T1/2 fibroblasts for musculoskeletal tissue engineering. PLoS One. 2015;10:e0139054.
26. Sharif J, Muto M, S-I, Suzuki, Itakura, A, Endo TA, Shiraya, J, Mizutani, Koseki Y, Toyoda T, Okamura K, et al. The SRA protein NPS5 mediates epigenetic inheritance by recruiting DNMT1 to methylated DNA. Nature. 2007;450:908–12.
27. Moudi M, Hoppner R, Abbady AQ, Monte D, Jeanblanc M, Oudert P, Louis B, Bronner C, KPB90 belongs to a new family of proteins with an expression that is deregulated in cancer cells. Br J Cancer. 2003;89:120–7.
28. Zhao J, Chen X, Song G, Zhang J, Liu H, Liu X. UHRF1 controls the self-renewal versus differentiation of hematopoietic stem cells by epigenetically regulating the cell-division modes. Proc Natl Acad Sci. 2017;114:E142–51.
29. Hajari KA. The biology of Annexin A2 from vascular fibroblast to innate immunity. Trans Am Clin Climatol Assoc. 2015;126:144–55.
30. Duarte WR, Shibata T, Takenaga K, Takahashi E, Kubota K, Ohyama K, Ishikawa I, Yamauchi M, Kasugai S. S100A4: a novel negative regulator of mineralization and osteoblast differentiation. J Bone Miner Res. 2003;18:493–501.
31. Mortazavi A, Williams BA, McCue K, Schaeffer L, Bold B. Mapping and quantifying mammalian transcriptomes by RNA-Seq. Nat Methods. 2008;5:621–8.
32. Honnet EA, Wheat CW. Quantitative RNA-Seq analysis in non-model species: assessing transcriptome assemblies as a scaffold and the utility of evolutionary divergent genomic reference species. BMC Genomics. 2012;13:361.
33. Diederich S, Tuan RS. Functional comparison of human-induced pluripotent stem cell-derived mesenchymal cells and bone marrow-derived mesenchymal stromal cells from the same donor. Stem Cells Dev. 2014;23:1594–610.
34. Obata Y, Furusawa Y, Endo TA, Sharif J, Takahashi D, Atarashi K, Nakayama M, Onawa S, Fujimura Y, Takahashi M, et al. The epigenetic regulator UHRF1 facilitates the proliferation and maturation of colonic regulatory T cells. Nat Immunol. 2014;15:571–9.
35. Tarazona S, Garcia-Alcalde F, Dopazo J, Ferrer A, Conesa A. Differential expression in RNA-Seq: a matter of depth. Genome Res. 2011;21:2213–23.
36. Hah N, Danko CG, Core L, Waterfall JJ, Siepel A, Lis JT, Kraus WL. A rapid, extensive, and transient transcriptional response to estrogen signaling in breast cancer cells. Cell. 2011;145:622–34.
37. Bana NA, Nyiri A, Nagy J, Frank K, Nagy T, Steger V, Schiller M, Lakatos P, Sugar L, Hori P, et al. The red deer Cervus Elaphus genome Cerela1.0: sequencing, annotating, genes, and chromosomes. Mol Gen Genomics. 2018;293:655–84.
38. Ker ED, Chu B, Phillips JA, Gharabeh B, Huard J, Weiss LE, Campbell PG. Engineering spatial control of multiple differentiation fates within a stem cell population. Biomaterials. 2011;32:413–22.
39. Ker ED, Nain AS, Weiss LE, Wang J, Suhan J, Amon CH, Campbell PG. Bioprinting of growth factors onto aligned sub-micron fibrous scaffolds for simultaneous control of cell differentiation and alignment. Biomaterials. 2011;32:8097–107.
40. Dobin A, Davis CA, Schlesinger F, Drenkow J, Zaleski C, Jha S, Batut P, Chaisson M, Gingeras TR. STAR: ultrafast universal RNA-Seq aligner. Bioinformatics. 2012;28:15–21.
41. Trapnell C, Roberts A, Goff L, Pertea G, Kim D, Kelley DR, Pimentel H, Salzberg SL, Rinn JL, Pachter L. Differential gene and Transcript expression analysis of RNA-Seq experiments with Tophat and cufflinks. Nat Protoc. 2012;7:562–78.
42. Derveaux S, Vandesompele J, Hellemans J. How to do successful gene expression analysis using real-time PCR. Methods. 2010;50:227–30.
43. Box GEP. Non-normality and tests on variances. Biometrika. 1953;40:318–35.
44. Box GEP. Some theorems on quadratic forms applied in the study of analysis of variance problems, I. effect of inequality of variance in the one-way classification. Ann Math Stat. 1954;25:290–302.
45. Tomarken AJ, Serlin RC. Comparison of ANOVA alternatives under variance heterogeneity and specific noncentrality structures. Psychol Bull. 1986;99:90–9.