The Co-Culture of ASCs and EPCs Promotes Vascularized Bone Regeneration in Critical-Sized Bone Defects of Cranial Bone in Rats

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Abstract

**Background:** The repair of critical-sized bone defect represents a challenging problem in bone tissue engineering. To address the most important problem in bone defect repair, namely insufficient blood supply, this study aimed to find a method that can promote the formation of vascularized bone tissue.

**Method** The phenotypes of ASCs and EPCs were identified respectively, and ASCs/EPCs were co-cultured in vitro to detect the expression of osteogenic and angiogenic genes. Furthermore, the co-culture system combined with scaffold material was used to repair the critical-sized bone defects of the cranial bone in rats.

**Results** The co-culture of ASCs/EPCs could increase osteogenesis and angiogenesis-related gene expression in vitro. The results of in vivo animal experiments demonstrated that the ASCs/EPCs group could promote bone regeneration and vascularization in the meantime and, then significantly accelerate the repair of critical-sized bone defects.

**Conclusion** It is feasible to replace traditional single seed cells with ASCs/EPCs co-culture system for vascularized bone regeneration. This system could ultimately enable clinicians to better repair the defect of craniofacial bone and avoid donor site morbidity.

**Background**
The repair of critical bone defects represents a challenging problem in bone tissue engineering. At present, bone tissue engineering technology has made great progress, but there are still some obstacles, such as osteoblasts unable to penetrate the center of the tissue-engineered bone, slow speed of ossification, as well as limited amount of bone regeneration[1]. The previous researches found it was mainly because the early nutritional support of tissue engineered bone mostly comes from the fusion of blood and tissue fluid, which severely impedes nutrition from entering the engineered bone center[2]. Thus, oxygen and blood supply, as well as metabolism of the cells in the central area are limited, resulting in insufficient tissue nutrition and poor osteogenesis. If angiogenesis and osteogenesis can be integrated to construct vascularized bone tissue, the vascular system is enable to provide oxygen and nutrition to the entire engineered bone tissue.

The application of endothelial progenitor cells (EPCs) is a potential method to achieve
vascularization[3]. Endothelial progenitor cells (EPCs) are precursor cells of vascular endothelial cells, which have the ability to proliferate, migrate and differentiate into cells arranged along the lumen of blood vessels[4]. It has been found in previous studies that EPCs can enhance the osteogenic activity of pre-osteoblasts[5], and play an important role in bone formation and repair[6, 7]. Adipose-derived mesenchymal stem cells (ASCs) are extracted from adipose tissue with high proliferative capability and multi-differential potential[8, 9]. Compared with stem cells derived from other tissues, ADSCs have the unique advantages of a wide range of tissue sources, easy availability and small damage to donor site[10]. Some studies have demonstrated that ADSCs can secrete vascular endothelial growth factor (VEGF) through the paracrine pathway and promote the formation of new blood vessels [11, 12]. Therefore, the co-culture of ADSCs and EPCs may promote the formation of new bone and blood vessels through cell-cell interactions.

To study the effect of ADSCs and EPCs co-culture on the construction of vascularized bone tissue, our team first identified the phenotypes of ASCs and EPCs, and then determined whether co-culture of ASCs and EPCs could increase the differentiation of osteoblasts and vascular endothelial cells in vitro. We further used the ASCs/EPCs co-culture cell system combined with scaffold materials to repair severe bone defects in the cranial bone of rats and analyzed the results. The above results showed that the co-culture of ASCs and EPCs facilitate bone regeneration and angiogenesis, and significantly promote the repair of critical-sized bone defects. It is feasible to use ASCs/EPCs co-culture cell system instead of traditional single seed cells for bone tissue engineering.

Materials And Methods
Preparation and Culture of ADSCs and EPCs
Animal procedures were conducted in accordance with the protocol approved by IACUC of China Medical University. Sprague-Dawley (SD) rats, 4-week- old, were euthanized by CO₂. The subcutaneous white adipose tissue of the inguinal region was cut off, while the blood vessels and other tissues were removed. The fresh tissue was washed three times with PBS containing 1% penicillin/streptomycin and minced by sterile surgical scissors. The minced tissue was digested with 0.1% collagenase type I at 37°C for 1 h and then centrifuged at 1000 rpm for 5 min. The upper
layer of fat was removed, the remaining cell pellet was resuspended and filtered through a 70 µm filter. After another centrifugation for 5 min, the cell pellet was resuspended in the culture medium consisting of low-glucose Dulbecco’s Modified Eagle Medium (DMEM), 10% fetal bovine serum (FBS), and 1% penicillin/streptomycin. The suspension was transferred to a flask and cultured at 37°C with 5% CO₂ in a humidity atmosphere[13]. The culture medium was replaced 24 h after transferring into flasks[14]. The medium was replenished every 2–3 days, and the cells were passaged after 80% confluence. The ASCs at passage 3 were used for the following experiment.

After 4-week-old SD rats were sacrificed with CO₂, their femurs were taken, and the bone marrow was evenly beaten. Single-cell suspension was obtained by centrifugation with Histopaque-1083, and mononuclear cell layer was separated by mononuclear cell separation solution. CD34+ cells were separated by immunomagnetic bead method. Cells were resuspended in EGM medium and seeded into culture flasks and cultured at 37°C with 5% CO₂ in a humidity atmosphere. The culture medium was replaced 24 h to remove nonadherent cells.

The medium was replenished every 2–3 days, and the cells were passaged after 80% confluence. The EPCs at passage 3 were used for the following experiment.

Flow Cytometry Analysis and Immunofluorescence Staining

1*10⁶ ASCs and EPCs at passage 3 were harvested, washed with 10% FBS/PBS and centrifuged at 1000 rpm, 5 min to gather cell pellets. For flow cytometry analysis, ASCs were stained with FITC-conjugated rat anti-CD70, Cy5.5-conjugated rat anti-CD90, PE-conjugated rat anti-CD45 and Alexa Fluor 647-conjugated rat anti-CD34 antibodies at a concentration of 2 mg/ml at 4°C[15]. EPCs were stained with PE-conjugated rat anti-CD31, Alexa Fluor 647-conjugated rat anti-CD34, FITC-conjugated rat anti-CD45 and FITC-conjugated rat anti-CD133, at a concentration of 2 mg/ml at 4°C. Mouse IgG was served as negative controls. Processed specimens were washed with 2 ml of 10% FBS/PBS for 30 min. After resuspended in 500 µl PBS, the cell pellets were tested by flow cytometry with 10,000 events recorded for each condition. The results were analyzed by FACS Express software. For immunofluorescence staining, EPCs at passage 3 were co-stained with DPBS-E containing 10 mg/ml Dil-labeled acLDL for 60 min at 37°C, then observed under fluorescence microscopy.
Co-culture of ASCs and EPCs in vitro
To determine the optimal ratio of EPCs and ASCs in bone regeneration, six groups were divided for experimental observation, namely ASCs alone, EPCs alone, and EPCs: ASCs at ratios of 1: 1, 1: 2, 1: 5, and 1:10. Cells were seeded in 12-well plates at the density of $1 \times 10^5$ cells per well and induced with EGM/CM media (EGM media: complete media ratio of 1:1) or EGM/OS media (EGM media: OS media ratio of 1:1) for 7 days, which was prepared for ALP activity assay as previously reported.

Alkaline phosphatase activity assay
ALP activity was detected by using an ALP assay kit (Sigma) following the manufacturer’s instructions. In brief, cells were mixed with an alkaline buffer solution (1.5 M, pH10.3) containing 10 mM p-nitrophenyl phosphate as a substrate and NaOH solution (3 M) was used as stop solution. The optical density was measured at 405 nm with a microplate reader. ALP activity was normalized by the DNA content and expressed as nmol of p-nitrophenol produced per minute per mg of total DNA$^{[45]}$. The implanted samples were smashed in liquid nitrogen, and lysed in 1 ml harvest buffer for 1 hour, then homogenized carefully to further lyse cells. After a centrifugation at 2000 rpm for 10 min, 10 ml supernatant were harvested for ALP activity assay.

Quantitative reverse transcription-polymerase chain reaction (qRT-PCR)
Total RNA was isolated using Trizol reagent (Invitrogen) according to manufacturer's instructions. Reverse transcription of total RNA was performed by RT-PCR (Invitrogen) using the reverse transcription first chain synthesis system. Real-time PCR reaction was performed with synthetic cDNA. Specific primers were used for PCR amplification to analyze the expression of osteoblastic marker genes including OCN, Col1a1, BMP2, vascular endothelial growth factor (VEGF), Cadherin (cdh5), and von Willebrand factor (vWF). According to the manufacturer's instructions, real-time PCR was performed using SYBR GREEN PCR Master Mix on ABI PRISM 7500 sequence detection system. PCR conditions were 94℃ 1 min, 95℃ 30 s, and 58℃ 40 s, with a total of 35 cycles. All reactions were repeated three times and normalized to GAPDH. Comparative ct was used to calculate the relative difference of PCR results.
Table 1
The primer sequences used for qRT-PCR

|          | Forward sequence | Reverse sequence |
|----------|------------------|------------------|
| OCN      | 5'-tcttttccttgccgcg-3' | 5'-caccgtctcaacttctccc-3' |
| Col1a1   | 5'-ggaagacagcctctccatac-3' | 5'-caatgctccaggagccacat-3' |
| BMP2     | 5'-ccgctccacatgagaa-3' | 5'-aaaggcatqatatgcccccgag-3' |
| VEGF     | 5'-ggccgaaactacgttttc-3' | 5'-aatctctctctcctgc-3' |
| cdh5     | 5'-gccgaaactacgttttc-3' | 5'-aatctctctctcctgc-3' |
| vWF      | 5'-ccggaagcgaccct-3' | 5'-cggctgcaatgagaa-3' |

Von Kossa staining

After ASCs/EPCs induced with osteogenic medium for 3 weeks, the formation of mineralized nodules in vitro was detected by Von kossa staining. Briefly described below, 1% silver nitrate solution was added to the culture medium for 45 min under ultraviolet light, rinsed with distilled water, and treated with 3% sodium thiosulfate for 5 minutes. After another rinse, specimens were re-stained and washed with ethanol, then finally performed microscopic observation, image acquisition and analysis.

Rat critical-sized cranial bone defect model

The in vivo experimental protocol was approved by IACUC of China Medical University. Thirty-two male SD rats at eight-week old underwent surgery under general anesthesia, induced by 5% isoflurane/O₂ gas inspiration and maintained by 1–2% isoflurane/O₂ by a facial mask. The scalps were shaved cleanly, disinfected with iodophors, and infiltrated with 0.1–0.5 ml of a local anesthetic agent of 2% lidocaine with epinephrine (0.01 mg/ml). The skin and periosteum were incised along the midline to expose the cranial bone surface[16]. A trephine bur with diameter of 8 mm was used to create a standardized, round, segmental defect around the sagittal suture, maintaining the underlying dura mater intact. A single implant of 1*10⁶ cells mixed with the scaffold was inserted into each defect. The periosteum and skin were sutured in layers with non-absorbable 4 – 0 prolene sutures. For 2 days after surgery, the rats were treated with carprofen for analgesia and penicillin for prevention of infection. Animals were divided into 4 groups randomly: group 1, blank group; group 2, hydroxyapatite/collagen scaffold only (HA/Col); group 3, HA/Col + ASCs, and; group 4, HA/Col + ASCs + EPCs. At the end of the eighth week after surgery, animals were euthanized with CO₂ and the repaired calvaria bones were harvested for the following analyses.

Analysis of bone regeneration

Micro-CT (Latheta LCT200) was used to scan the harvested samples for 3D imaging analysis.
Meanwhile, Bone Mineral Density (BMD, g/cm²) were performed with LUNAR PIXImus bone densitometer and analyzed by LUNAR PIXImus software according to the manual book of the equipment. A total of 6 samples were analyzed in each group. On the computerized scan, 5 regions of interest (ROI) of each slide were selected to measure the BMD of the defect area, and the average of these values was taken as the final result.

For histological analysis, six samples per group were decalcified and cut into 5 mm sections, half of which were used for hematoxylin-eosin (HE) staining and the other half for immunohistochemistry analysis. Digital images of each slide were acquired using a digital camera mounted to a microscope. Newly formed bone areas in the total defect area were calculated manually at 10 × magnification by using NIH ImageJ software.

Analysis of Blood Vessel Ingrowth
VEGF was detected by immunohistochemistry in paraffin embedded and decalcitonized bone sections. As previously mentioned, VEGF is largely present in the subendothelial matrix. Half of the samples in each group were analyzed independently. The primary antibody of goat VEGF was diluted at a ratio of 1:300, and the hrp-conjugated rabbit anti-goat antibody was diluted at a ratio of 1:500 to 1% BSA. The peroxidase activity was observed with diaminobenzidine. Both the negative control group and the positive control group were included under the same conditions. Negative control staining was performed on the same bone plates without primary antibodies. The kidney sections as positive group were stained with the same primary and secondary antibodies. AxioImager software was used for image acquisition. Using the NIH Image J software, the number of blood vessels in the entire implant area of each sample displayed by VEGF staining was calculated manually in a 10x magnification manner.

Statistical analysis
The test results were expressed as mean ± standard deviation, and statistical analysis was performed by SPSS 22.0 software (IBM SPSS Statistics, IBM, Armonk, NY). T-test was used for the comparison of the data of two groups and one-way ANOVA was for the data of multiple samples. P < 0.05 was considered as statistically significant.

Results
Detection and Analysis of ASCs and EPCs Surface Markers

As shown in Fig. 1, ASCs are positive for CD73 and CD90 and negative for CD45 and CD34. Meanwhile, the immunofluorescence staining results of ASCs cell surface markers CD73 and CD90 (Fig. 2) also suggested that CD73 and CD90 were positive. The flow cytometry results of EPCs are shown in Fig. 3, indicating that EPCs were positive for CD133 and CD34 and negative for CD11b and CD31. To confirm the EPC phenotype, Dil-ac-LDL and lectin staining of EPCs were performed after the cells were cultured for 7 days. As shown in Fig. 4, Dil-ac-LDL (red) and lectin (green) staining both were positive.

ASCs / EPCs co-culture results

Co-culture of ASCs and EPCs at the optimal ratio

In the previous work, ASC and EPC in different ratios were compared respectively for ALP activity after cultured with osteogenic induction medium and whole medium (Fig. 5). The comparison results showed that the ALP activity of ASCs/EPCs at 1: 1 ratio was significantly higher than that of other groups after induced by osteogenic induction culture. With the same co-culture ratio, the ALP activity of the osteogenic induction medium was higher than that of the whole medium group.

Co-culture of ASCs and EPCs increases osteogenesis and angiogenesis-related gene expression

To further investigate whether the co-culture of ASCs and EPCs affected the expression of osteogenic and angiogenic genes, OCN, Col I, and BMP2 for osteogenesis, and VEGF, cdh5, and vWF for angiogenesis were analyzed by using real-time PCR. The results showed that the expression levels of osteogenic genes OCN, Col I and BMP2 in the ASCs/EPCs group were significantly higher than those in the ASCs or EPCs groups (Fig. 6), suggesting that the co-culture of ASCs and EPCs can increase the expression of osteogenesis-related genes. The mRNA levels of angiogenic genes, including VEGF, cdh5 and vWF were also dramatically higher in the ASCs/EPCs groups than in the single cell groups (Fig. 6), suggesting that the co-culture can increase the expression of angiogenesis-related genes. It was indicated in these results, that co-culture of ASCs and EPCs can enhance both of osteogenesis and angiogenesis-related gene expression. After the osteogenic medium induction was continued for 3 weeks, the formation of in vitro mineralized nodules was detected by Von kossa staining. The results showed that the number of in vitro mineralized nodules of ASCs / EPCs was significantly
increased and the osteogenesis ability was significantly enhanced (Fig. 7).

Co-Culture of ASCs and EPCs Promotes Vascularized Bone Regeneration in Critical-Sized Bone Defects of Cranial Bone in Rats

To evaluate the potential of ASCs/EPCs for bone and vascular regeneration in vivo, 8-week-old SD rats with critical-sized bone defect (d = 8 mm) of the cranial bone were prepared as animal models.

ASCs/EPCs were combined with hydroxyapatite/collagen scaffolds to repair the critical-sized bone defects, and the repaired bones were harvested at the 8th week after surgery. As shown in the three-dimensional imaging by Micro-CT scan (Fig. 8), the group 4 (HA/Col + ASCs + EPCs) had remarkably stronger osteogenic activity. Compared with the other three groups, the bone defect area in the group 4 was almost completely closed, which indicated that the co-culture of ASCs and EPCs enable the promotion of bone regeneration. Quantitative analysis of BMD showed that bone density in the implantation area of the group 4 was dramatically higher than the other three groups (p < 0.05) (Fig. 9).

HE staining sections showed no residual material or inflammatory infiltrating cells in the defect area after 8 weeks of implantation. In all samples, the amount of new bones in the blank and HA/Col groups was significantly smaller when compared with other groups, and most of the defect area was covered with fibrous tissue (Fig. 10A, B). In the group 2 (HA/Col + ASCs), new bone was formed, partially covering the defect area, and still part of the defect area was covered by fibrous tissue (Fig. 10C). While the group 4 showed robust osteogenic activity, and regenerated bone continuously covered almost all defect areas (Fig. 10D). Histomorphometric analysis showed that the amount of newly-formed bone (BA) to the total implant area (TA) in the group 4 was significantly greater than that in the other four groups (p < 0.05) (Fig. 11), indicating that EPCs could promote bone regeneration.

The ingrowth of blood vessels in the regenerated bone

To confirm the formation of blood vessels within the implants, immunostaining with VEGF antibodies was performed, which could specifically stain and identify vascular endothelial cells. As shown in Fig. 12A-D, much more blood vessels were found in HA/Col + ASCs + EPCs group compared to the other three groups. By quantifying the blood vessel density (BVD) of the entire implant area, it was
confirmed that the BVD of the ASCs/EPCs group was significantly higher than that of the other groups (Fig. 13). These results demonstrated that EPCs could promote vascular growth, and that the co-culture of ASCs and EPCs dramatically enhances vascularized bone regeneration.

**Discussion**

For the repair of critical-sized bone defect, the difficulties are mainly focused on the formation of vascularized bone, especially the center of the defect. To construct bone tissue accompanying vascular system, our team co-cultured ASCs and EPCs to establish a dual stem cell system. In this present study, it was found that the ASCs/EPCs co-culture system can enhance the expression of osteogenic and angiogenic genes in vitro, and furthermore, by supporting vascularized bone regeneration, it can significantly accelerate bone healing of critical-sized bone defects in vivo.

EPCs are precursor cells of vascular endothelial cells, which have the ability to proliferate, migrate and differentiate into cells arranged along the lumen of blood vessels and can be isolated from peripheral blood and spleen. ASCs also have high proliferative growth characteristics and multi-differential potential, can be extracted from autologous subcutaneous fat, and have a wide range of tissue sources. But to date, specific markers for each cell type are still lacking. A number of surface proteins have been used to identify rat adipose stem cells, including CD73, CD90, CD105, CD44, etc[17–20]. Here, CD73 and CD90 are used as positive markers to identify ASCs. By using hematopoietic stem cell (HSCs) marker CD45 and EPCs marker CD34 to identify ASCs free of HSCs and EPCs, our results demonstrated that ASCs expressed a cell-surface protein profile positive for CD73 and CD90 and negative for CD45 and CD34. For EPCs, we detected the cell markers CD34 and CD133 (AC133) [21], which are highly expressed in EPCs, but not expressed after the EPCs differentiated into mature vascular endothelial cells. Meanwhile, we also detected CD31, which is not expressed in EPCs, but highly expressed in mature endothelial cells, as well as, CD11b, which is expressed in monocytes, but not in EPCs[22–25]. Our results showed that EPCs expressed a surface protein profile positive for CD133 and CD34 and negative for CD11b and CD31. These results confirmed the ASCs and EPC phenotypes, suggesting that the above markers can be used for the identification of ASCs and EPCs.
To determine the effect of these two kinds of cells in osteogenic and angiogenic differentiation, ASCs and EPCs were co-cultured in vitro, and then the osteogenesis-related genes (OCN, Col I and BMP2) and angiogenesis-related genes (VEGF, cdh5 and vWF) were analyzed by real-time RT-PCR. The results showed that the ASCs/EPCs co-culture system can increase gene expression of osteogenesis and angiogenesis. Von kossa staining was used to detect the formation of mineralized nodules in vitro, which also confirmed that ASCs/EPCs co-culture can improve the osteogenesis of this cell system.

Based on the in vitro results, we further performed an in vivo experiment on the repair of critical-sized bone defects in rats. The Micro-CT scans of the rat's cranial bones were performed after 5 weeks after surgery. The results clearly showed that the amount of new bone in the defect area was much greater in the HA/Col + ASCs + EPCs group than in the other groups, including HA/Col + ASCs group. In addition, BMD also showed that there was more bone tissue formation in the HA/Col + ASCs + EPCs group. Moreover, as confirmed by immunohistochemical analysis, the blood vessel density in the defect area was higher in the HA/Col + ASCs + EPCs group than in other three groups. The possible reason for this phenomenon is that the lack of EPCs in the ASCs group resulted in a relative decrease in vascularization and bone formation, which relied solely on the inward growth of the host blood vessels. However, the distance between the blood vessels of the host tissue and the center of the bone defect is far from sufficient to achieve bone regeneration, especially for critical-sized defect. As a result, nutrients, metabolites and other molecules cannot be delivered to the center area of defect, which severely impedes bone regeneration[26]. In the ASCs/EPCs group, EPCs directly increased the invasion of blood vessels and promoted the differentiation of ASCs to osteogenesis and, meanwhile, pre-osteogenic ASCs could promote the recruitment of EPCs and enhance the ability of EPCs to form blood vessels. The interaction accelerates vascularization and bone formation in the meantime, making nutrients, cytokines, and other molecular factors involved in the bone healing process more accessible. Taken together, these results indicated that the ASCs/EPCs co-culture system can promote the formation of vascularized bone during bone regeneration and achieve the repair of the critical-sized bone defect.
Conclusion

In summary, this study provides evidence that the co-culture of ASCs and EPCs promotes osteogenic and angiogenic differentiation in vitro and significantly accelerates bone regeneration in critical-sized cranial bone defect of rat in vivo by supporting vascularization and bone formation in the meantime. On the basis of the data presented here, it was concluded that it is feasible to replace traditional single seed cells with ASCs/EPCs co-culture system for vascularized bone regeneration. This system could ultimately enable clinicians to better repair the defect of craniofacial bone and avoid donor site morbidity.

Abbreviations

EPCs: Endothelial progenitor cells.
ASCs: Adipose-derived mesenchymal stem cells.
BMP2: Bone morphogenic protein 2
VEGF: Vascular endothelial growth factor
Cdh5: Cadherin,
vWF: von Willebrand factor
HA/Col: Hydroxyapatite/collagen scaffold
BMD: Bone Mineral Density
BVD: Blood vessel density
HSCs: Hematopoietic stem cell

Declarations

Acknowledgements

Not applicable

Authors’ contributions

Yuanjia He did the analysis and interpretation of data and manuscript drafting.
Shuang Lin did the data analysis.
Qiang Ao helped in the critical discussion.
Xiaoning He conceived and designed the experiments and directed the manuscript drafting, financial support, and final approval of the manuscript.
All authors read and approved the final manuscript.

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**Availability of data and materials**

The datasets used and/or analyzed during the current study are available from the corresponding author on reasonable request.

**Ethics approval and consent to participate**

All experiments were performed according to standard protocols, in compliance with the Guide of the Animal Ethics Committee of China Medical University. Consent to participate is not applicable.

**Consent for publication**

Not applicable

**Competing interests**

The authors declare that they have no competing interests.

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Figures

![Figure 1](image-url)

**Figure 1**

Surface markers of ASCs flow cytometry.
ASCs immunofluorescence staining. (a) CD73; (b) DAPI; (c) CD73 + DAPI; (d) CD90; (e) DAPI; (f) CD90 + DAPI. It was shown that the ASCs markers CD73, CD90 are positive.

Surface markers of EPCs flow cytometry.
EPCs immunofluorescence staining. (a) VEGF; (b) DAPI; (c) VEGF + DAPI; (d) Dil-ac-LDL; (e) lecin; (f) merge of d and e.
Figure 5

Changes in ALP of ASCs and EPCs in different ratios (A: ASCs, E: EPCs). * p < 0.05.
Figure 6

RT-PCR detection of mRNA expression of osteogenic and angiogenic genes, osteogenic related genes OCN, Coll, BMP2; angiogenic related genes VEGF, cdh5, vWF. (A: ASCs, E: EPCs). #p < 0.05.

Figure 7

Von Kossa staining. The number of calcified nodules in the ASCs / EPCs co-culture group was greater than that in the ASCs and EPCs groups.
Three-dimensional Micro-CT imaging of the cranial bone at 8 weeks following surgery. (A: blank group, B: HA/Col group, C: HA/Col+A group, D: HA/Col+A+E group)

Figure 9

Quantitative analysis of BMD in the implantation area, * p < 0.05.
Figure 10

Hematoxylin and eosin staining analysis of new bone formation (A-D low magnification, bar = 1mm; A'-D 'high magnification, bar = 0.5mm. A, A': blank control group, B, B ': HA/Col group, C, C ': HA / Col + ASCs group, D, D ': HA / Col + ASCs + EPCs group)
Figure 11
Quantitative analysis of bone area in implanted region. BV[bone area in implant; TV, total implant area, * p <0.05.

Figure 12
Immunostaining of VEGF in the newly-formed bone (A: blank control group, B: HA / Col group, C: HA / Col + ASCs group, D: HA / Col + ASCs + EPCs group)
Figure 13

Quantitative analysis of blood vessel density.