Histone Deacetylase 7-Derived Peptides Play a Vital Role in Vascular Repair and Regeneration

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1. Introduction

Cardiovascular disease (CVD) represents a major cause of morbidity and mortality globally. Intimal injury is a common pathological basis of the process of CVD, and the recovery of injured endothelium is essential for protection against CVD.[1,2] Dysfunction of endothelial cells (ECs) is thought to play a pivotal role in injured endothelium, and EC health has been suggested to be maintained by a consistent supply of supporting growth factors and activation of local stem/progenitor cells that are capable of differentiation toward ECs.[3,4] It is well documented that vascular progenitor cells (VPCs), which are mainly located in the adventitia, contribute to the repair of the injured endothelium via migration toward the endothelium and differentiation into an EC lineage.[5,6] These VPCs are also an important source for the regeneration of tissue-engineered vascular grafts.[7,8]

Vascular bypass surgery has been successfully employed for the therapy of vascular diseases. Unfortunately, in the case of small-diameter vascular grafts (diameter less than 6 mm), a high rate of restenosis limits the clinical use of commercially available products. The main reason for this is the poor regeneration of vascular tissue including slow and poor endothelialization.[9,10] In addition, there is a compelling need for novel strategies to revascularize ischaemic tissues, and critical limb ischaemia continues to be a severe health problem globally. In many cases, surgical or graft-based procedures are not possible, necessitating the development of molecular or cell-based therapies to promote angiogenesis.[11,12]

Histone deacetylases (HDACs) are a family of enzymes that remove acetyl groups from N-acetylated lysine residues on histones and are involved in gene transcriptional regulation through modulating chromatin structures. HDAC7, a member of the class II HDACs, is specifically expressed in the vascular endothelium and contributes to the maintenance of vascular integrity during early embryogenesis.[13,14] Recently, we demonstrated that a 7-amino acid (aa)-peptide (7A, MHSPGAD) could be alternatively translated from a short open reading frame within the 5’-terminal untranslated region of mouse Hdac7 mRNA in stem cell antigen-1 positive (Sca1+) VPCs in vitro and in vivo. The serine residue within the 7A peptide could be...
phosphorylated by the activated kinase MEKK1 S393, which in turn could transfer the phosphate group to the Thr145 site of 14-3-3γ, forming a novel MEKK1-7A-14-3-3γ signaling pathway downstream of vascular endothelial growth factor (VEGF). This novel signaling pathway contributed to the activation of the Sca1+VPCs.[20] In the present study, we focused on the functional analysis of this 7A peptide and its translational potential in vascular repair and regeneration by using different disease and transplantation models.

2. Results

2.1. The 7A Peptide Increased VPC Migration and Differentiation toward the EC Lineage

Recently, we found that endogenous 7A peptide could only be detected in in vitro cultured Sca1+VPCs or Sca1+ cells in injured femoral arteries and not in Sca1+ cells in normal vessels. Therefore, it is very important to investigate the relationship between 7A expression and the activation of Sca1+ VPCs. Considering 7A can function as a phosphate carrier, a synthetic phosphorylated 7A (7Ap) (MH[pSer]PGAD) was included in the functional analyses. As controls, scramble 7S (MPHASGD) and 7Aa (MHAPGAD), in which the serine of the 7A is substituted by alanine to totally abolish phosphorylation, were also included in this study.

As shown in Figure 1A, 7A especially 7Ap accelerated Sca1+VPCs migration. As expected, 7Aa retarded Sca1+VPCs migration. Wound healing can be a combined effect of cell migration and proliferation. To assess whether 7A/7Ap-induced migration was derived from the combined effect, a Br-du incorporation assay was performed. Under the serum-free condition, the 7A peptide by itself could not support Sca1+VPC proliferation (Figure S1A, Supporting Information). Further experiments revealed that 7A had no effect on cell survival under oxidative stress (Figure S1B, Supporting Information).

The vessel wall resident Sca1+VPCs can differentiate toward both EC and smooth muscle cell (SMC) lineages.[17,18] When embryonic stem cells differentiate into SMCs, the Hdac7 transcript variant 2 undergoes further splicing, leading to the incorporation of 7A into the far N-terminal end of the HDAC7 protein.[20] Thus, we assumed that 7A might be involved in the cell differentiation process. To test this, Sca1+VPCs were cultured in differentiation medium in the presence of 7-aa peptides and/or VEGF, followed by quantitative RT-PCR analysis of EC and SMC marker expression. As shown in Figure 1C, 7A increased Pecam1 (CD31) and Cdh5 (CD144) mRNA levels, which were significantly enhanced by VEGF, although VEGF alone had only a slight effect. The effect of 7Ap alone on Pecam1 and Cdh5 expression was comparable to the combined effect of 7A and VEGF (Figure 1B left and middle). 7A alone had no effect on Tagln (SM22) expression but significantly decreased Tagln expression in the presence of VEGF, although VEGF had a slight increasing effect. 7Ap alone significantly decreased Tagln expression. These results suggest that 7A, especially 7Ap, favors Sca1+VPC differentiation toward EC but not SMC lineages. The EC differentiation was further confirmed by tube formation assays (Figure 1C).

2.2. The 7aa-Peptide Enhanced Vascular Injury Repair and Angiogenesis in Ischaemic Tissues In Vivo

As described above 7A, especially 7Ap, could facilitate VPC mobilization and differentiation toward the EC lineage, suggesting that the 7A peptide may have therapeutic potential in vascular injury repair and angiogenesis in ischaemic tissues. To test this, the wire-guided femoral artery injury model was created in ApoE−/− mice.[19] Pluronic-127 gel containing 10 ng mL−1 of peptides or PBS was applied surrounding the adventitia of the injured vessels, which were harvested four weeks postsurgery. The hematoxylin and eosin (H&E) staining of cryo-sections revealed that 7A or 7Ap administration significantly reduced neointima formation (Figure 2A).

Furthemore, a hindlimb ischaemia model was introduced in eight-month (Figure 2B) and ten-week (Figure S3, Supporting Information) old C57BL/6j mice, in which Pluronic-127 gel containing 10 ng mL−1 peptides was applied surrounding the injured vessels.[20] Foot blood perfusion was measured by a Doppler Scanner on day 7 and day 14 postsurgery. Young mice showed better foot blood perfusion recovery compared to aged mice (PBS group in Figure S3, Supporting Information, vs Figure 2B). Importantly, 7Ap significantly promoted recovery in both groups of mice, which might be due to the increased formation of a new Sca1+ cell niche in the ischaemic tissues (Figure 2C).

2.3. Fabrication of Peptide-Loaded Tissue Engineered Vascular Grafts (TEVGs)

To evaluate whether the 7A peptide has therapeutic potential, TEVGs were fabricated by a co-electrospinning technique.[20] The
Figure 1. 7A and 7Ap peptides increased migration and differentiation toward the EC lineage. A) 7Ap significantly increased VPC migration in a wound healing model. Wound was introduced into confluent VPCs by tip scratching, and incubated with DMEM medium containing 2% FBS and 1 ng mL$^{-1}$ of 7S, 7Aa, 7A, or 7Ap peptide. Images were taken at 0, 12, 24, and 36 h postscratching (left, Scale bar: 100 µm). The migrated cells in scratched area were counted from three views per scratching, three scratchings per well, and three wells per peptide (right). Data presented are representative images or mean of three independent experiments. *: $p < 0.05$ (7Ap vs 7S) ($n = 6$, one-way-ANOVA followed by Tukey's post hoc analysis). B,C) 7A/7Ap increased VPC differentiation toward the EC lineage. The 3 d spontaneously differentiated VPCs were incubated with differentiation medium containing 1 ng mL$^{-1}$ peptides and 10 ng mL$^{-1}$ VEGF for 4 d, followed by quantitative RT-PCR analysis with GAPDH as house-keeping gene (B) or tube formation assay (C, Scale bar: 200 µm). 1% BSA was included as vehicle control. The data presented are representative images or mean of three independent experiments ($n = 6$, two-way ANOVA followed by Dunnett's multiple comparison tests). D) MEKK1-7A-14-3-3$\gamma$ mediated 7aa-peptide-induced VPC differentiation toward EC lineage. The VPCs were transfected with siRNA and cultured in differentiation medium for 3 d, followed by further differentiation with same protocol described above for 4 d, followed by quantitative RT-PCR analysis of Pecam1, Cdh5, and Tagln mRNA levels with Gapdh as house-keeping gene. 1% BSA was included as vehicle control. The data presented are representative images or mean of three independent experiments. *: $p < 0.05$. **: $p < 0.01$ ($n = 6$, two-way ANOVA followed by Dunnett's multiple comparison tests).
TEVGs were sized at 2.0 mm diameter and 600 µm wall thickness (Figure 3A). They showed a micro/nanohybrid fibrous structure, consisting of PCL microfibers (≈6 µm) to provide mechanical support and structural maintenance and collagen nanofibers (≈600 nm) to act as a carrier for delivery of peptides (Figure 3B). The loading amount is about 417.5 ng for the graft of 1 cm length that has been calculated from the feeding ratio of electrospinning. Both fibers were distributed homogeneously across the graft wall as revealed by scanning electron microscopy (SEM) (Figure 3B middle). The distribution of the two types of fibers could also be clearly identified by double labeling showing PCL in red and collagen in green (Figure 3B right). The peptides (7S, 7A, 7Ap) released from the grafts as a result of the degradation of the collagen fibers exhibited a sustained profile over a period of 30 d. The cumulative release reached \(43 \pm 7\%\) (s.d.) of the theoretical value (Figure 3C) in 7A group, and a similar releasing profile has been observed in 7S and 7Ap groups (Figure S4, Supporting Information). The TEVGs (1.0 cm in length) were further evaluated by in vivo implantation in a rat abdominal artery replacement model (Figure 3D left). The patency of the implanted grafts was first examined by ultrasound at different timepoints (Figure 3D, right). Most of the grafts had good patency without aneurysms or bleeding and patency rates were higher in the 7A/7Ap groups than in the PBS and 7S groups (Figure 3E).[21,38] Further analysis by stereomicroscopy revealed that the luminal surfaces of all groups were smooth, clean and free of thrombi at 12 weeks (Figure S5, Supporting Information).

2.4. 7Ap Promoted Vascular Regeneration and Remodeling in TEVGs

The endothelialization on the lumen side of the implanted TEVGs was first observed by scanning electronic microscopy at three locations selected continuously from the anastomotic site to the midportion. As shown in Figure 4A, the lumen side of the 7A and 7Ap-loaded TEVGs was already covered by a layer of
cells at two weeks postimplantation. In contrast, the PBS and 7S-loaded TEVGs were only partially covered, and bare fibers could still be identified in the midportion. After four weeks post-implantation, all of the grafts were almost fully covered by a confluent layer of neo-tissue. In the 7A group, especially the 7Ap group, the cells oriented along the blood flow, resembling the endothelium of the native artery.

4′,6-diamidino-2-phenylindole (DAPI) staining revealed that a number of cells had incorporated into the wall of all of the TEVGs (Figure 4B). Immunofluorescence staining demonstrated that there was a layer of ECs on the lumen side (Figure 4B). The statistical analysis of the CD144+ cell-covering area via immunofluorescence staining on the longitudinal sections revealed that 7Ap-loaded TEVGs had much higher endothelialization rate compared to other groups (Figure 4C and Figures S6 and S7, Supporting Information).

After 12 weeks postimplantation, all of the grafts were almost fully endothelialized without detectable differences (Figure S8, Supporting Information). These results suggest that 7A especially 7Ap can facilitate endothelialization of TEVGs.

In the native arterial vessels, ECs form a thin layer of intima while one to several layers of SMCs lie in the media. To assess whether the endothelialization of the TEVGs was accompanied by SMC layer formation, immunofluorescence staining was first performed on the cross sections with anti-α-SMA antibody. As shown in Figure 5A, a layer of SMA+ cells could be observed after four weeks, and the SMCs were well organized. The evolution of α-SMA+ layers from 4 to 12 weeks showed a divergent tendency among the four groups, that is, their thickness decreased in the 7A and 7Ap groups and increased in the PBS and 7S groups (Figure 5B). This indicates that 7A, especially 7Ap, could reduce the overproliferation of SMCs that lead to adverse intimal hyperplasia.

The regeneration of mature smooth muscle (SM-MHC+) did not show any significant differences at four weeks post-implantation (Figure 5C). After 12 weeks, the thickness of the SM-MHC+ layer was remarkably higher in the 7Ap group than in the PBS one (Figure 5D), suggesting that 7Ap could modulate the infiltrated SMCs into a more mature contractile phenotype. Furthermore, double immunofluorescence staining with anti-vWF (red) and anti-α-SMA (green) antibodies exhibited better tissue regeneration in the 7Ap group compared to the control group, showing a layer of neo-endothelium and several layers of aligned SMCs beneath (Figure 5E). At the same time, we noticed that some of the vWF-positive cells were SMA-positive as well. To address whether there is endothelial-to-mesenchymal transition, further detailed investigation is required.

Successful vascular grafts need capillary vessels to support cells in the graft wall. Immunofluorescence staining with anti-CD31 antibody on the cross sections of the TEVGs revealed that the addition of 7A especially 7Ap effectively increased capillary vessel formation even at two weeks postimplantation (Figure S9A, Supporting Information). After four weeks, the density of capillary vessels in 7Ap group was significantly higher than that in other groups (Figure S9B, Supporting Information). These results suggest that 7A especially 7Ap can increase capillary vessel formation. The 7Ap-loaded TEVGs resembled native vasculature more closely.
2.5. 7Ap Stimulated Local Vascular Stem/Progenitor Cells Migration and Differentiation into the EC Lineage

As our in vitro studies have shown that 7A, especially 7Ap, could increase Sca1+VPC migration and differentiation toward the EC lineage, we wondered whether the local VPCs contributed to the neo-vasculature formation in the TEVGs. To test this, immunofluorescence staining was performed with anti-Sca1 antibody on sections from two- and four-week implanted TEVGs. As shown in Figure 6A, the infiltration of Sca1+VPCs within the graft walls could be detected at two weeks postimplantation. As time went on, the Sca1+ cell number increased. Very importantly, 7A, especially 7Ap, enhanced this process, since the 7Ap group had the highest number of Sca1+ cells.
Further experiments with double immunofluorescence staining using CD144 (red) and Sca1 (green) antibodies (Figure 6C) revealed that the majority of the CD144+ cells were also Sca1-positive (Figure 6C, magnified image), indicating that these new ECs are derived from the Sca1+VPC differentiation.

It has been established that both local resident and circulating stem/progenitor cells contribute to vascular remodeling following vascular injury. To explore the origin of the infiltrated Sca1+cells and their contribution to the vascular remodeling in TEVGs, a bi-layered vascular graft was employed in this study (Figure 7A, upper panel). In addition to the regular vascular grafts that act as an inner layer, an external layer composed of PCL nanofibers was introduced as an outside barrier (Figure S10, Supporting Information). Due to its low porosity, this external layer could effectively restrict the infiltration of cells from the surrounding tissues after implantation for two weeks (Figure 7Aa,b). Quantitative analysis based on DAPI staining showed that the average cell number was significantly decreased within the graft wall, especially in the area near the lumen (Figure 7Ac). The infiltration of Sca1+VPCs was further compared between the two groups, and statistical data showed that the density of the Sca1+cells was markedly reduced due to the outside barrier at both two and four weeks (Figure 7B). All of these results confirm that surrounding tissues are an important source for the Sca1+VPCs infiltration within the vascular grafts.

Endothelialization of the two types of vascular grafts was compared by SEM observation (Figure S11, Supporting Information) and immunofluorescence staining (Figure 7C, Figures S12 and S13, Supporting Information). The results indicated that the outside barrier slowed down the endothelialization process, and the average coverage ratio of neo-endothelium was significantly
lower in the outside barrier (bi-graft) graft than in the control one (Figure 7C). These results suggest that both the adjacent ECs\cite{21,27} and the circulating and local resident stem/progenitor cells migrate into and contribute to vascular remodeling in the TEVGs.

To further verify the effect of 7Ap peptides on the migration of Sca1\(^+\)-VPCs from surrounding tissues into the grafts, Matrigel containing GFP-Sca1\(^+\)-VPCs was applied surrounding the adventitia of vascular grafts (Figure 8A). Immunofluorescence images of the cross sections showed that after 3 d

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**Figure 6.** 7Ap increased Sca1\(^+\)-VPC differentiation toward ECs contributing to endothelialization of TEVGs. A) Immunofluorescence staining was performed to detect the recruitment of Sca1\(^+\)-VPCs using anti-Sca1 antibody on the implanted TEVGs sections at time indicated, and B) quantification of the infiltrated Sca1\(^+\) cells within 0–100 \(\mu\)m depth from the lumen by randomly selecting six views (40×) from each cross section. C) The contribution of the Sca1\(^+\) progenitor cells-derived ECs to the endothelialization was detected by double immunofluorescence staining with anti-Sca1 (green) and anti-CD144 (red) antibodies on the four weeks implanted TEVGs sections. DAPI was used to counterstain the nuclei. Data are represented as the mean ± SEM for each group (\(n = 6\) at 2 and 4 weeks, and \(n = 4\) at 12 weeks). \(*: p < 0.05; **: p < 0.01\) (one-way-ANOVA followed by Tukey’s post hoc analysis).
Figure 7. Bi-layered PCL TEVGs revealed that the Sca1⁺-VPCs were mainly derived from the surrounding tissues. A) An illustration of the bi-layered PCL fabricated TEVGs. B,C) The outer layer with small pores reduced Sca1⁺-VPCs recruitment into the TEVGs wall and the lumen endothelialization of the TEVGs. B) The single PCL and bi-layered PCL fabricated TEVGs were implanted into rats to replace the abdominal arteries, and harvested at two and four weeks, respectively, postimplantation, followed by immunofluorescence staining with anti-Sca1 antibody to show the recruitment of Sca1⁺-VPCs in the TEVGs wall or C) with anti-CD31 (green, upper) and anti-CD144 (red, lower) antibodies to show the lumen endothelialization. DAPI was used to counterstain the nuclei. Data are represented as the mean ± SEM. *: p < 0.05; **: p < 0.01; ***: p < 0.001 (n = 6, two-way ANOVA followed by Dunnett’s multiple comparison tests).
postimplantation the number of cells migrated to the graft wall was significantly ($p < 0.05$) lower in the control group compared to the 7Ap group (Figure 8B,C). The fate of infiltrated Sca1+VPCs from surrounding tissues has also been investigated by seeding the GFP-Sca1+VPCs into the graft wall directly (Figure 8D). Double immunofluorescence staining using CD144 and Sca1 antibodies demonstrated after 7 d postimplantation some Sca1+VPCs differentiated into ECs, and 7Ap significantly ($p < 0.05$) enhanced the number of double-positive cells on the lumen side compared to the control group (Figure 8E,F). All of these results confirmed that Sca1+VPCs from the surrounding tissues can migrate transmurally and differentiate into the EC lineage in the TEVGs.

3. Discussion

Tissue-engineered vascular grafts prepared by traditional strategies are time-consuming and costly, which often restricts their clinical application. Instead, a new in situ (cell-free) concept has received increasing attention, that is, a neo-artery is completely generated in situ by taking advantage of the regenerative potential of the body. To stimulate or maximize this in situ regeneration capability, vascular grafts have been engineered in terms of physical structure and surface chemistry, as well as the delivery of vasoactive molecules. In this study, we demonstrated that an Hdac7-derived 7-aa peptide can serve as a vasoactive molecule to mobilize local resident progenitor cells, contributing to in situ neo-artery formation.

The origin of cells participating in the regeneration of vascular tissues remains a major obstacle restricting the success of in situ tissue engineering strategies. We tend to believe that vascular cells (ECs and SMCs) migrate from the anastomotic native tissues when implanted a polymer-based graft as an inferior vena cava interposition in mice. However, the results are often limited by many factors, including graft structure, implant site, and animal species. For example, cell migration from other directions (such as surrounding tissues) has been restricted
due to the dense structure of the graft walls. It has been accepted that endothelial cell migration from the anastomotic sites is generally limited to 1 cm, which is far less than the clinical requirement of re-endothelialization of relatively longer vascular grafts. Therefore, there is a need for recruitment of cells (both mature ECs and stem/progenitor cells) from sites beyond the anastomotic border via circulation or through migration from the surrounding tissue.

Vascular tissue-resident stem cells have been discovered to display the capacity to differentiate into vascular cell lineages, which may contribute to the regenerative process. More specifically, the adventitial resident VPCs play a major role in the repair of the injured endothelium via migration toward the endothelium and differentiation into an EC lineage.

HDAC7 plays an important role in the maintenance of endothelium integrity. In a parallel study, we have demonstrated that mouse Hdc7 mRNA can undergo alternative translation to produce a biological active 7-aa peptide, which functions as a phosphate transfer carrier in cellular signal transduction. In this study, we have evaluated its application potential in peptide-loaded tissue engineered vascular grafts by incorporating the 7A/7Ap peptides into the micro/nanofibrous vascular grafts.

In vitro studies demonstrated the 7Ap peptide is very powerful in Sca1+ VPCs-derived EC differentiation. However, we also noticed that 7Ap seemed less effective during EC differentiation in the presence of VEGF compared to 7Ap alone. This phenomenon may be due to the diverse functions of VEGF. In vitro differentiation process, the increased EC cells can be derived from both the newly differentiated ECs and the proliferation of the differentiated mature ECs. When mature ECs undergo proliferation, the EC marker especially those involved in cell-to-cell contact such as CD31 (PECAM1) and CD144 (Cdh5) will be downregulated to relieve the connection among adjacent cells. This will result in an observable decrease in EC marker expression at mRNA levels and tube formation. Although 7Ap had no effect on EC proliferation, VEGF is a mitogenic factor for EC. Thus, the combination of 7Ap and VEGF may show a decrease in EC marker expression and tube formation compared to 7Ap alone.

An intact endothelial cell monolayer has proven to be an antithrombogenic interface to maintain the patency of the implanted vascular grafts. In the meantime, the recruitment of SMCs is vital to stabilize the newly formed endothelium. In this study, we have demonstrated that the inclusion of 7A, especially the phosphorylated version 7Ap, significantly increased endothelialization. The majority of the ECs, especially those in the midportion of the TEVGs, may be derived from the local resident Sca1+ progenitor cells of the surrounding tissues, because 7Ap can serve as a chemoattractant for Sca1+ VPCs and direct their differentiation toward the EC lineage. The ECs on the lumen of TEVGs will secrete relevant cytokines (such as PDGF or TGF-β1) to recruit SMCs to stabilize the newly formed endothelium. The interactions between ECs and SMCs are very important for the function and homeostasis of the regenerated neo-artery. Considering that 7Ap suppresses Sca1+ VPCs differentiation toward SMCs and has no effect on SMC migration and proliferation, the recruitment of SMCs may be mainly mediated by ECs on the lumen of the TEVGs. After all, precise lineage tracing may be required to verify the contribution of Sca1+ VPCs.

Therefore, 7Ap may help with the organization and maturation of SMCs within the TEVGs, which could also explain the discrepancy between in vivo SMC regeneration and in vitro SMC differentiation. Importantly, the Sca1+ progenitor cells also contribute to capillary vessel formation in the TEVGs wall, which is critical for the functional maintenance of the neo-artery following scaffold degradation.

The dose-dependency of the 7A/7Ap peptide is an important factor affecting therapeutic efficacy. 7A/7Ap was effective in EC differentiation at a concentration of 0.1 ng mL−1. For cell migration, it was more effective at a relatively higher concentration, and therefore 1 ng mL−1 was used. In vivo studies, we assumed that there would be a concentration gradient extending from the carrier (F-127 gel or collagen fiber), and the final concentration within the ischaemic tissue reached ≈0.1 ng mL−1. For future studies, it will be necessary to detect whether there is dose dependency.

In summary, the clinical potential of histone deacetylase 7 (HDAC7)-derived 7-amino acid peptides in vascular repair and regeneration has been systematically evaluated. The effect of the 7A peptide, especially the phosphorylated 7A peptide (7Ap), on VPC migration and EC differentiation has been investigated in vitro functional analyses. Local delivery of 7A/7Ap increased re-endothelialization and suppressed neointima formation in the femoral artery injury model, and promoted foot blood perfusion recovery in the hindlimb ischaemia model. Degradeable tissue-engineered vascular grafts loaded with peptides have been successfully prepared and evaluated in an abdominal aorta replacement model of rats. Sustained delivery of 7A, especially 7Ap, from tissue-engineered vascular grafts enhanced endothelialization and intima/media formation in the vascular graft via inducing Sca1+ VPC migration and differentiation. All of these results suggest that 7A/7Ap provides a promising therapeutic strategy for ischaemia diseases and vascular graft bypass surgery. Because the human orthologue of Sca1 has not been identified yet, it will be necessary to investigate whether 7A/7Ap has similar effect on human adult stem/progenitor cells, which will provide direct evidence for its possible application in translational medicine.

Supporting Information
Supporting Information is available from the Wiley Online Library or from the author. All mice experiments were performed according to protocols approved by the Institutional Committee for the Use and Care of Laboratory Animals, UK. The use of experimental rats was approved by the Animal Experiments Ethical Committee of Nankai University and carried out in conformity with the Guide for Care and Use of Laboratory Animals.

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Conflict of Interest
The authors declare no conflict of interest.
Keywords
HDAC7-derived peptide, ischaemia disease, tissue-engineered vascular grafts (TEVGs), vascular progenitor cells (VPCs)

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[1] K. H. Park, W. J. Park, J. Korean Med. Sci. 2015, 30, 1213.
[2] J. Davignon, P. Ganz, Circulation 2004, 109, 1127.
[3] Q. Xu, Z. Zhang, F. Davison, Y. Hu, Circ. Res. 2003, 93, e76.
[4] P. Libby, Arterioscler., Thromb., Vasc. Biol. 2012, 32, 2045.
[5] H. K. Yip, T. H. Tsai, H. S. Lin, S. F. Chen, C. K. Sun, S. Leu, C. M. Yuen, T. Y. Tan, M. Y. Lan, C. W. Liou, C. H. Lu, W. N. Chang, Crit. Care 2011, 15, R40.
[6] A. Zampetaki, J. P. Kirton, Q. Xu, Cardiovasc. Res. 2008, 78, 413.
[7] A. Margariti, B. Winkler, E. Karamariti, A. Zampetaki, T. N. Tsai, D. Baban, J. Raguossis, Y. Huang, J. D. Han, L. Zeng, Y. Hu, Q. Xu, Proc. Natl. Acad. Sci. USA 2012, 109, 13793.
[8] P. Campagnolo, T. N. Tsai, X. Hong, J. P. Kirton, P. W. So, A. Margariti, E. Di Bernardini, M. M. Wong, Y. Hu, M. M. Stevens, Q. Xu, Biomaterials 2015, 60, 53.
[9] Y. Yao, J. Wang, Y. Cui, R. Xu, Z. Wang, J. Zhang, K. Wang, Y. Li, Q. Zhao, D. Kong, Acta Biomater. 2014, 10, 2739.
[10] E. Pektok, B. Nottelet, J. C. Tille, R. Ganz, A. Calonge, S. Moeller, B. H. Waltho, Circulation 2008, 118, 2563.
[11] E. A. Silva, E. S. Kim, H. J. Kong, D. J. Mooney, Proc. Natl. Acad. Sci. USA 2008, 105, 14347.
[12] B. Park, A. Hoffman, Y. Yang, J. Yan, G. Tiej, H. Bagshahi, P. T. Nowicki, L. M. Messina, J. Vasc. Surg. 2010, 51, 165.
[13] W. Fischle, S. Emilian, M. J. Hendzel, T. Nagase, N. Nomura, W. Voelter, E. Verdin, J. Biol. Chem. 1999, 274, 11713.
[14] S. Chang, B. D. Young, S. Li, X. Qi, J. A. Richardson, E. N. Olson, Cell 2006, 126, 321.
[15] D. Ummarino, Y. Li, L. Zeng, Curr. Angiogr. 2013, 3, 60.
[16] L. Zeng, J. Yang, J. Clin. Exp. Cardiol. 2016, 7, 11(suppl).
[17] Z. Tang, A. Wang, F. Yuan, Z. Yan, B. Liu, J. S. Chu, J. A. Helms, S. Li, Nat. Commun. 2012, 6, 875.
[18] A. Margariti, Q. Xiao, A. Zampetaki, Z. Zhang, H. Li, D. Martin, Y. Hu, L. Zeng, Q. Xu, J. Cell Sci. 2009, 122, 460.
[19] Q. Xiao, L. Zeng, Z. Zhang, A. Margariti, Z. A. Ali, K. M. Channon, Q. Xu, Y. Hu, Arterioscler., Thromb., Vasc. Biol. 2006, 26, 2244.
[20] L. Zeng, Q. Xiao, M. Chen, A. Margariti, D. Martin, A. Ivetic, H. Xu, J. Mason, W. Wang, G. Cockerill, K. Mori, J. Y. Li, S. Chien, Y. Hu, Q. Xu, Circulation 2013, 127, 1712.
[21] T. Pennel, D. Beuzidhenu, J. Koehe, N. H. Davies, P. Zilla, Acta Biomater. 2018, 65, 237.
[22] J. M. Daniel, D. G. Sedding, J. Mol. Cell. Cardiol. 2011, 50, 273.
[23] P. J. Psaltis, A. Harbuzaria, S. Delacroix, E. W. Holroyd, R. D. Simari, J. Cardiovasc. Transl. Res. 2011, 4, 161.
[24] C. Zhang, L. Zeng, C. Emanuelli, Q. Xu, Cardiovasc. Res. 2013, 99, 251.
[25] H. Talacua, A. I. Smits, D. E. Muylaert, J. W. van Rijswijk, A. Vink, M. C. Verhaar, A. Driessen-Mol, L. A. van Herwerden, C. V. Bouten, J. Klun, F. P. Baaijens, Tissue Eng., Part A 2015, 21, 2583.
[26] Z. Wang, Y. Cui, J. Wang, X. Yang, Y. Wu, K. Wang, X. Gao, D. Li, Y. Li, X. L. Zheng, Y. Zhu, D. Kong, Q. Zhao, Biomaterials 2014, 35, 5700.
[27] N. Hibino, G. Villalona, N. Pietris, D. R. Duncan, A. Schoffer, J. D. Roh, Y. T. Li, L. W. Dobrucki, D. Mejias, R. Sawh-Martinez, J. K. Harrington, A. Sinusas, D. S. Krause, T. Kyriakides, W. M. Saltzman, J. S. Pober, T. Shin’oka, C. K. Breuer, FASEB J. 2011, 25, 2731.
[28] S. Li, D. Sengupta, S. Chien, Wiley Interdiscip. Rev.: Syst. Biol. Med. 2014, 6, 61.
[29] W. Zheng, Z. Wang, L. Song, Q. Zhao, J. Zhang, D. Li, S. Wang, J. Han, X. L. Zheng, Z. Yang, D. Kong, Biomaterials 2012, 33, 2880.
[30] Q. Ji, S. Zhang, J. Zhang, Z. Wang, J. Yang, Y. Cui, L. Pang, S. Wang, D. Kong, Q. Zhao, Biomacromolecules 2013, 14, 4099.
[31] Z. Wang, Y. Lu, K. Qin, Y. Wu, Y. Tian, J. Wang, J. Zhang, J. Hou, Y. Cui, K. Wang, J. Shen, Q. Xu, D. Kong, Q. Zhao, J. Controlled Release 2015, 210, 179.
[32] L. P. Brewster, D. Buflalino, A. Ucuzian, H. P. Greisler, Biomaterials 2007, 28, 5028.
[33] A. Mahar, S. Sonekawa, N. Kobayashi, Y. Hirano, Y. Kimura, T. Fujisato, T. Yamaoka, Biomaterials 2015, 58, 54.
[34] Y. Hu, Z. Zhang, E. Torsney, A. R. Afzal, F. Davison, B. Metzler, Q. Xu, J. Clin. Invest. 2004, 113, 1258.
[35] S. Hui, K. R. Brun, M. Husain, Circ. Res. 2010, 106, 1180.
[36] A. Margariti, A. Zampetaki, Q. Xiao, B. Zhou, E. Karamariti, M. Martin, X. Yin, M. Mayr, H. Li, Z. Zhang, E. De Falco, Y. Hu, G. Cockerill, Q. Xu, L. Zeng, Circ. Res. 2010, 106, 1202.
[37] J. M. Richardson, A. Dawson, N. O’Hagan, P. Taylor, D. J. Finnegan, M. D. Walkinshaw, EMBO J. 2006, 25, 1324.
[38] S. Row, H. Peng, E. M. Schlai, Q. Koenigsknecht, S. T. Andreadis, D. D. Swiftz, Biomaterials 2015, 50, 115.
[39] M. B. Chan-Park, J. Y. Shen, Y. Cao, Y. Xiong, Y. Liu, S. Rayatpisheh, G. C. Kang, H. P. Greisler, J. Biomater. Release 2009, 88, 1104.
[40] L. Beck Jr., P. A. D’Amore, FASEB J. 1997, 11, 365.
[41] K. K. Hirsch, S. A. Rohovs, P. A. D’Amore, J. Cell Biol. 1998, 141, 805.
[42] W. N. Makow, J. Deng, X. Z. Ruan, Q. Xu, Arterioscler., Thromb., Vasc. Biol. 2017, 37, e41.