Abstract. Previous studies have shown that the tetraspanin CD151 is essential for pathological or physiological angiogenesis. However, the cellular signaling mechanism and the role of the CD151 YRSL sorting motif in \textit{in vitro} vasculogenesis remains unknown. In this study, the results showed that both CD151 and CD151-ARSA gene delivery were capable of increasing the expression of CD151 at the protein level in human umbilical vein endothelial cells (HUVECs). Moreover, there was no significant difference in CD151 protein expression between the CD151 group and the CD151-ARSA group. Overexpression of CD151 promoted HUVEC cell proliferation, migration and capillary network formation \textit{in vitro}. However, in the CD151-ARSA group, the abilities of cell proliferation, migration and capillary network formation were all decreased, compared with the CD151 group. Furthermore, the activation of PI3K, Akt and ERK signaling pathways was attenuated in the CD151-ARSA mutant group compared with the CD151 group. This study suggests that the YRSL motif of CD151 plays a key role in CD151-induced angiogenesis. Our observations provide insights into a new mechanism of CD151 regulating angiogenesis via vesicle trafficking.

Introduction

CD151, a tetraspanin superfamily protein, contains two extra-cellular loops, four hydrophobic transmembrane domains and two short cytoplasmic tails (1,2). This tetraspanin is expressed broadly in various tissues and is particularly abundant in endothelia, epithelia, smooth muscle and megakaryocytes (3,4). At the cellular level, CD151 is characteristically localized in intracellular vesicles and at cell-cell junctions in endothelial cells (ECs) (4). Moreover, the association of CD151 with integrins stands out as a prominent feature (3,5,6). A ‘CD151-integrin’ complex model has been proposed and this model is functionally linked to CD151-induced biological processes (6-9).

As previously shown, vesicle trafficking is a fundamental membrane trafficking event in cell migration, and endocytic trafficking pathways have become established in recent years as being important in the regulation of intracellular signaling pathways during processes such as cell division, migration and angiogenesis (18-21). Recently, several studies noted that vesicle trafficking was also one characteristic feature of CD151 and observed that the YRSL sequence of CD151 was necessary for internalization and vesicle trafficking of CD151 (22). It was revealed that the C-terminal cytoplasmic domain of CD151 contains a YRSL sequence or YXXφ sorting motif, in which Y is tyrosine, X is any amino acid, and φ represents the amino acid residue with a bulky hydrophobic
side chain (23, 24). Mutation of this CD151 YRSL motif markedly attenuated internalization and vesicle trafficking of CD151 (22). The YRSL motif-mediated internalization of CD151 was thought to promote cell migration by modulating the endocytosis and/or vesicular trafficking of CD151 (22). Hence, we hypothesized that the YRSL motif of CD151 may be responsible for the regulation of CD151-related human umbilical vein endothelial cell (HUVEC) migration and capillary network formation in vitro, and CD151 may affect angiogenesis via vesicle trafficking.

In this study, we mutated CD151 YRSL→ARSA, which was capable of impairing the vesicle trafficking of CD151 (22). We then examined the roles of CD151 and the CD151-ARSA mutant in the cell proliferation, migration and capillary network formation of HUVECs, with a recombinant adeno-associated virus (rAAV) construct encoding CD151 and the CD151-ARSA mutant. The purpose of the present study was to investigate the critical role of the YRSL sequence of CD151 during angiogenesis in vitro and the mechanism(s) involved.

Materials and methods

Materials and reagents. All cell culture reagents were obtained from Gibco BRL Life Technologies, Inc. (Grand Island, NY, USA), including Dulbecco's modified Eagle's medium (DMEM/F12), trypsin and fetal bovine serum (FBS). Apigenin was supplied by Calbiochem-Novabiochem (Darmstadt, Germany). Rabbit anti-ERK1 and anti-phospho-ERK1 antibodies were purchased from New England Biolabs (Darmstadt, Germany). Antibodies against PI3K (Beverly, MA, USA). The restriction enzymes were purchased from Thermo Scientific (Waltham, MA, USA) (BamHI and NotI restriction sites). Proper ligation was confirmed by sequencing.

Preparation of recombinant adeno-associated viruses (rAAVs). The rAAV vector pXXU1, packaging plasmid pX2, adeno-virus helper plasmid pHelper, and a rAAV plasmid containing GFP cDNA was obtained from Dr Xiao Xiao (University of North Carolina, Chapel Hill, NC, USA). The packing and production of rAAV-GFP, rAAV-CD151 and rAAV-CD151-ARSA were carried out using a triple-plasmid cotransfection method in human embryonic kidney cells [293 cells, American Type Culture Collection (ATCC), Manassas, VA, USA] (12, 13). For purification, a single-step gravity-flow column was applied (25). The titers of vector particles were determined.

HUVEC culture and transfection. HUVECs were obtained from ATCC and grown in DMEM/F12 medium supplemented with 10% FBS (Gibco), streptomycin 100 µg/ml and penicillin 100 U/ml (all obtained from Sigma) at 37°C under 5% CO2 and 95% air. Only cells passaged less than five times were used for experiments. Cells were grown to 50–60% confluence and transfected with rAAV-GFP, rAAV-CD151 and rAAV-CD151-ARSA, as described previously (14). For the control group, PBS or HD-Fugene was added. Cells were incubated in the conditions above for 3 days and then subjected to the following assays.

Cell proliferation assay. Assessment of cell viability was performed using the Cell Counting Kit-8 (CCK-8) assay. At 12, 24 and 48 h after rAAV transfection, HUVECs were incubated in 10% CCK-8 (Beyotime Institute of Biotechnology, Nantong, China) diluted in normal culture medium at 37°C until visual color conversion occurred in 96-well culture plates. The number of viable cells was assessed by measurement of absorbance at 450 nm using a microplate reader.

Cell migration assay. Assessment of cell migration was performed using a cell wound-healing assay. Briefly, transfectant cells were cultured in 60-mm diameter dishes and synchronized in 0.5% FBS. After HUVECs were grown to confluence, wounds were generated by scraping the monolayers with sterile pipette tips. After 0, 12, 24 and 48 h culture at 37°C, respectively, images of the wound in each well were captured using an inverted microscope (Nikon TE 2000; Nikon, Tokyo, Japan).

Capillary network formation assay. Assessment of cell migration was performed using capillary network formation on Matrigel. Briefly, Matrigel (0.5 ml) was polymerized on 24-well plates, and 5x10^5 transfectant cells were then plated in full-growth medium for 1 h. Once the cells were seeded, the medium was replaced with medium containing 0.5% serum. After incubation at 37°C for 12, 24 and 48 h, the capillary network formation was visualized using an inverted microscope (Nikon TE 2000) equipped with digital imaging. For each treatment, 10 field images were captured, and the area containing endothelial tubes and networks that had formed was quantified using the Scion Image Analysis System (Windows version of Scion Image, NIH) with background subtraction.
Protein extraction and western blotting. HUVEC proteins were extracted and used for western blot analysis (12,14), with specific primary antibodies against CD151, PI3K, Akt, phospho-Akt, ERK1, phospho-ERK1 and β-actin. The HRP-conjugated secondary antibodies were used respectively to reveal the specific protein bands with ECL detection reagents. The intensities of protein bands were quantified by densitometry. In addition, inhibitors of MAPK (apigenin) and PI3-kinase (LY294002) were added to cultured HUVECs. The protein levels of PI3K, Akt, phospho-Akt, ERK1 and phospho-ERK1 were observed.

Statistical analysis. Data were analyzed using SPSS 18.0 statistical software (Chicago, IL, USA). Data were presented as the means ± standard deviation (SD) unless otherwise specified. Statistical comparisons between two groups were carried out using the Student’s t-test or one-way ANOVA. *p<0.05 was considered to indicate a statistically significant result.

Results

Expression of CD151 protein after transfection. In the present study, we mutated the YRSL motif in the human CD151 molecule. Gene sequencing analysis showed that the Y and L residues in the CD151 YRSL motif were replaced with alanine (A) residues simultaneously, and the resulting YRSL → ARSA mutant was designated as the CD151-ARSA mutant (Fig. 1A). As shown in Fig. 1B, HUVECs transfected with rAAV-GFP observed using an inverted fluorescence microscope 7 days after transfection (the same field). (C) Western blot analysis. (D) Quantitative analysis of CD151 protein expression. β-actin was used as an internal loading control. The mean density of CD151 in control group was defined as 100%. Each experiment was performed at least in triplicate. *p<0.05 vs. control and GFP group. HUVECs, human umbilical vein endothelial cells; rAAV, recombinant adeno-associated virus; GFP, green fluorescent protein.

Figure 1. Expression of CD151 protein after transfection in HUVECs. (A) The gene sequences of CD151 and CD151-ARSA. The CD151 mutation (CD151-ARSA) changed the motif of YRSL245-248 to ARSA245-248. (B) HUVECs transfected with rAAV-GFP observed using an inverted fluorescence microscope 7 days after transfection (the same field). (C) Western blot analysis. (D) Quantitative analysis of CD151 protein expression. β-actin was used as an internal loading control. The mean density of CD151 in control group was defined as 100%. Each experiment was performed at least in triplicate. *p<0.05 vs. control and GFP group. HUVECs, human umbilical vein endothelial cells; rAAV, recombinant adeno-associated virus; GFP, green fluorescent protein.

Figure 2. Effects of CD151 and CD151-ARSA transfection on HUVECs. Cell Counting Kit-8 (CCK-8) assays were performed at 24 and 48 h after gene delivery. The CD151 group showed promoted proliferation ability. In the CD151-ARSA mutation group, the proliferation of HUVECs was decreased significantly at 24 or 48 h after delivery. Three independent experiments were carried out, and each experiment was in triplicate. *p<0.05 vs. control and GFP group. *p<0.05 vs. CD151 group. HUVECs, human umbilical vein endothelial cells; GFP, green fluorescent protein.

Effects of CD151 and CD151-ARSA transfection on the proliferation of HUVECs. To determine the proliferative
effects of CD151 and CD151-ARSA, we performed the CCK-8 assay at 12, 24 and 48 h after delivery. As shown in Fig. 2, the CD151 group showed increased proliferation ability, compared with the control group and the GFP group at 24 and 48 h after rAAV transfection. No significant difference was observed in the groups at 12 h after transfection (data not shown). However, the results showed that the CD151-induced proliferation of HUVECs was decreased significantly by the CD151-ARSA transfection, at 24 or 48 h after delivery (Fig. 2). These data suggest that the CD151-ARSA mutant was capable of impairing the cell proliferation ability of CD151.

**Effects of CD151 and CD151-ARSA transfection on the migration of HUVECs.** Regulation of cell motility is a prominent...
feature of CD151 (4,22). Cell migration was assessed by a cell wound-healing assay and observed at 0, 12, 24 and 48 h after CD151 transfection. As shown in Fig. 3A and B, the expression of CD151 significantly enhanced cell migration at 24 and 48 h, compared with the control and GFP groups. However, at 24 h after delivery, the migration ability of CD151-ARSA mutant cells was reduced compared with the CD151 group, and the most notable effect was at 48 h (Fig. 3).

Thus, these results suggest that CD151 transfection promotes wound healing while the CD151-ARSA mutant transfection delays wound healing.

**Effects of CD151 and CD151-ARSA transfection on the capillary network formation of HUVECs.** Upon plating on Matrigel basement membrane, HUVECs assembled into capillary network formation structures, which were observed at 12, 24 and 48 h after the rAAV transfection. As shown in Fig. 4A and B, CD151 transfection promoted the capillary network formation at 24 and 48 h after gene delivery, compared with the control and GFP groups (Fig. 4). By contrast, network formation was inhibited by transfection with the CD151-ARSA mutant at 24 and 48 h after gene delivery, compared with the CD151 group, and there was a marked difference at 48 h. These findings indicate that CD151-ARSA mutant delivery impairs the tube formation improved by CD151 gene transfer.

**Effects of CD151 and CD151-ARSA mutant on the PI3K/Akt and ERK signaling pathways.** A number of signaling pathways are involved in angiogenesis, such as Akt, eNOS, ERK and p38 MAPK (26,27). We analyzed the levels of PI3K, Akt, phospho-Akt and quantitative analysis. Inhibitor of PI3K (LY294002, 15 µM); inhibitor of MAPK (apigenin, 25 µM). HD group (HD-Fugene 6 transfection reagent) was used as a control group. Each experiment was performed at least in triplicate. *p<0.05 vs. control, GFP and HD groups. *p<0.05 vs. CD151 group. *p<0.05 vs. CD151-ARSA group (no signaling pathway inhibitor group). HUVECs, human umbilical vein endothelial cells; GFP, green fluorescent protein.

![Figure 5. Effects of CD151 and CD151-ARSA transfection on PI3K/Akt and ERK signaling pathways. Western blot analysis for PI3K, phosphorylated Akt, total Akt, phosphorylated ERK and total ERK. Inhibitors of MAPK (apigenin) and PI3K (LY294002) were applied to HUVECs following transfection with CD151 or CD151-ARSA.](image-url)
CD151-ARSA mutant resulted in reduced activation of PI3K/Akt and ERK signaling pathways (Fig. 5). In addition, MAPK inhibitor (apigenin) and PI3K inhibitor (LY294002) significantly reduced the activation of ERK and PI3K induced by CD151, respectively (Fig. 5).

As a whole, these results suggest that the gene transfer of CD151-ARSA mutant attenuates the activation of the PI3K/Akt and ERK signaling pathways.

**Discussion**

The present study was designed to investigate the molecular mechanisms that govern the effects of CD151 in angiogenesis. In this in vitro study, our results showed that the delivery of the CD151-ARSA mutant (YRSL→ARSA) into HUVECs decreased cell migration and capillary network formation on Matrigel, contrary to the effects of CD151 gene delivery. Furthermore, we demonstrated that mutation of the CD151 YRSL motif resulted in diminished activation of PI3K/Akt and ERK signaling pathways in HUVECs. These data provide evidence that the CD151 YRSL motif was indeed a key region of CD151 for regulating cell migration, capillary network formation and angiogenesis.

Multiple lines of evidence indicate that the YRSL sequence of the CD151 C-terminal cytoplasmic tail is important and it is thought to determine its intracellular trafficking and function (24). These motifs in the cytoplasmic domain of CD151 could be recognized by adapter protein (AP)-2 complex, a core component of clathrin endocytic machinery (28). It was found that the YRSL motif was required for CD151 endocytic processes, indicating that the YRSL sequence in the CD151 cytoplasmic domain determines its trafficking (22). When the YRSL→ARSA mutant human CD151 molecule was created and transfected into NIH3T3 mouse fibroblast cells, the vesicle trafficking was completely impaired and CD151-promoted NIH3T3 migration was diminished (22). Therefore, the YRSL sequence of CD151 is necessary for vesicle trafficking of CD151 and this sequence is also a key region for CD151-related cell migration.

Angiogenesis is a complex process involving extracellular matrix degradation, endothelial cell proliferation and migration, formation of tube structures and morphological differentiation (29-31). Although our previous data showed that CD151 was capable of promoting cell proliferation, cell migration and angiogenesis both in vivo and in vitro, the mechanisms remain to be elucidated. Based on above data, we hypothesized that the YRSL sequence of CD151 may also play an important role in the CD151-mediated process of angiogenesis. To test this hypothesis, in the present study we mutated the YRSL→ARSA motif in the human CD151 molecule and transfected it into HUVECs. As shown in the western blot analysis, CD151 and the CD151-ARSA mutant were both well expressed at the protein level, and the mutation did not influence the expression of CD151 protein. Furthermore, evidence indicates that the CD151-ARSA mutant transfection abrogates CD151-induced cell proliferation and migration in vitro. First, CD151-ARSA treatment decreased the HUVEC proliferation, in contrast to the promoting effects of CD151. Second, the directional motility of HUVECs (cell-wounding healing assay) showed that the CD151-ARSA mutant exhibited delayed cell motility, compared with CD151 transfectant. The YRSL motif was required for CD151 regulating cell migration, consistent with the results obtained from Liu et al (22).

Our previous work has shown that CD151 gene delivery promoted capillary formation in vitro (16). Although the specific region of CD151 QRD194-196 was emphasized (14,16), the YRSL motif may be another key region in CD151-induced angiogenesis. As was previously reported, endocytosis of adhesion molecules from cell-cell contacts may be regulated according to the YRSL motif (4,22,32). In particular, the capillary network formation was related to the maintenance of cell-cell contacts or junctions, which could be regulated by the lateral trafficking of cell adhesion molecules (4,21,22), and it was questioned whether CD151 may affect the capillary formation caused by the YRSL motif through vesicular trafficking. In the present study, it was shown that CD151-ARSA mutant transfection significantly disrupted the capillary network formation, while CD151 promoted the capillary network formation on Matrigel. The capillary network formation was inhibited by transfection with the CD151-ARSA mutant at 24 and 48 h after gene delivery. Notably, the difference became more significant over time, and a marked difference was observed at 48 h. Thus, it was observed that CD151 is capable of regulating the capillary network formation through the YRSL motif. Based on these data, it was accepted that the CD151 YRSL sequence is critical for CD151-mediated capillary network formation. Therefore, CD151 YRSL sequence-mediated vesicular trafficking may be another mechanism for CD151 regulation of angiogenesis in vitro.

The signaling mechanism of CD151 has been explored in the last decade (9,11,31). A growing list of signaling pathways that may be involved includes FAK, ERK and PI3K/Akt (11,13-15,31). Takeda et al showed that the adhesion-dependent activation of PKB/c-Akt and e-NOS was diminished in CD151-null mouse lung endothelial cells (11). In the present study, the CD151-ARSA mutant transfection resulted in diminished activation of PI3K/Akt and ERK signaling pathways, which was opposite to the findings with CD151 transfection. Thus, the activation of PI3K/Akt and ERK signaling pathways may be involved in CD151-mediated endocytosis trafficking.

In conclusion, our data demonstrated that the CD151-ARSA mutant abrogated cell proliferation, migration and capillary network formation in HUVECs. Furthermore, we observed the CD151-mediated activation of ERK and PI3K/Akt signaling pathways, but these effects were all impaired by CD151-ARSA gene delivery. Our observations emphasize that the specific YRSL region of CD151 is important for vesicle trafficking, which plays a key role in CD151-induced cell proliferation, migration and capillary network formation. Our study demonstrates for the first time the effect of CD151 YRSL in CD151-induced angiogenesis. Further studies should be performed and may provide indications for a better understanding of CD151 and CD151-ARSA via integrin-vesicle trafficking.

**Acknowledgements**

The project was supported by the grant no. 81000047 from the National Natural Science Foundation of China.
References

1. Fitter S, Tetaj TZ, Berndt MC and Ashman LK: Molecular cloning of cDNA encoding a novel platelet-endothelial cell tetraspan antigen, PETA-3. Blood 86: 1348-1355, 1995.

2. Hasegawa H, Utsunomiya Y, Kishimoto K, Yanagisawa K and Fujita S: SFA-1, a novel cellular gene induced by human T-cell leukemia virus type 1, is a member of the transmembrane 4 superfamily. J Virol 70: 3258-3263, 1996.

3. Sincock PM, Mayrhofer G and Ashman LK: Localization of the transmembrane 4 superfamily (TM4SF) member PETA-3 (CD151) in normal human tissues: comparison with CD9, CD63, and alpha5beta1 integrin. J Histochem Cytochem 45: 515-525, 1997.

4. Sincock PM, Fitter S, Parton RG, Berndt MC, Gamble JR and Ashman LK: PETA-3/CD151, a member of the transmembrane 4 superfamily, is localised to the plasma membrane and endocytic system of endothelial cells, associates with multiple integrins and modulates cell function. J Cell Sci 112: 833-844, 1999.

5. Yauch RL, Berditchevski F, Harler MB, Reichner J and Hemler ME: Highly stoichiometric, stable, and specific association of integrin alpha3beta1 with CD151 provides a major link to phosphatidylinositol 4-kinase, and may regulate cell migration. Mol Biol Cell 9: 2751-2765, 1998.

6. Zhang XA, Kazarov AR, Yang X, Bontrager AL, Stipp CS and Hemler ME: Function of the tetraspanin CD151-alpha6beta1 integrin complex during cellular morphogenesis. Mol Biol Cell 13: 1-11, 2002.

7. Hemler ME: Integrin associated proteins. Curr Opin Cell Biol 10: 578-585, 1998.

8. Serru V, Le Naour F, Billard M, et al.: Selective tetraspan-integrin complexes (CD81/alpha4beta1, CD151/alpha3beta1, CD151/alpha6beta1) under conditions disrupting tetraspan interactions. Biochim J 340: 103-111, 1999.

9. Zhang XA, Bontrager AL and Hemler ME: Transmembrane-4 superfamily proteins associate with activated protein kinase C (PKC) and link PKC to specific beta1 integrins. J Biol Chem 276: 25005-25013, 2001.

10. Wright MD, Geary SM, Fitter S, et al.: Characterization of mice lacking the tetraspanin superfamily member CD151. Mol Cell Biol 24: 5978-5988, 2004.

11. Takeda Y, Kazarov AR, Butterfield CE, et al.: Deletion of tetraspanin Cd151 results in decreased pathologic angiogenesis in vivo and in vitro. Blood 109: 1524-1532, 2007.

12. Lan RF, Liu ZX, Liu XC, Song YE and Wang DW: CD151 promotes neovascularization and improves blood perfusion in a rat hind-limb ischemia model. J Endovasc Ther 12: 469-478, 2005.

13. Zhao H, Liu Z, Liu X, et al.: CD151 gene delivery after myocardial infarction promotes functional neovascularization and activates FAK signaling. Mol Med 15: 307-315, 2009.

14. Liu WF, Zu ZH, Chai BL, et al.: Role of tetraspanin CD151-alpha3/alpha6 integrin complex: Implication in angiogenesis CD151-integrin complex in angiogenesis. Int J Biochem Cell Biol 43: 642-650, 2011.

15. Zhong ZZ and Liu ZX: Activation of the phosphatidylinositol 3-kinase/protein kinase Akt pathway mediates CD151-induced endothelial cell proliferation and cell migration. Int J Biochem Cell Biol 39: 340-348, 2007.

16. Zuo H, Liu Z, Yang J, et al.: Activation of the ERK signaling pathway is involved in CD151-induced angiogenic effects on the formation of CD151-integrin complexes. Acta Pharmacol Sin 31: 805-812, 2010.

17. Zuo H, Liu Z, Liu X, et al.: Assessment of myocardial blood perfusion improved by CD151 in pig myocardial infarction model. Acta Pharmacol Sin 30: 70-77, 2009.

18. Breitscher MS and Aguado-Velasco C: Membrane traffic during cell locomotion. Curr Opin Cell Biol 10: 537-541, 1998.

19. Kobayashi T, Vischer UM, Rossobelleti C, et al.: The tetraspanin CD63/lamp3 cycles between endocytic and secretory compartments in human endothelial cells. Mol Biol Cell 11: 1829-1843, 2000.

20. Dufield A, Kamsteeg EJ, Brown AN, Pangel P and Caplan MJ: The tetraspanin CD63 enhances the internalization of the H,K-ATPase beta-subunit. Proc Natl Acad Sci USA 100: 15560-15565, 2003.

21. Caswell PT, Vadrevu S and Norman JC: Integrins: masters and slaves of endocytic transport. Nat Rev Mol Cell Biol 10: 843-853, 2009.

22. Liu L, He B, Liu WM, Zhou D, Cox JV and Zhang XA: Tetraspanin CD151 promotes cell migration by regulating integrin trafficking. J Biol Chem 282: 31631-31642, 2007.

23. Ohno H, Stewart J, Fournier MC, et al.: Interaction of tyrosine-based sorting signals with clathrin-associated proteins. Science 269: 1872-1875, 1995.

24. Bonifacio JS and Traub LM: Signals for sorting of transmembrane proteins to endosomes and lysosomes. Annu Rev Biochem 72: 395-447, 2003.

25. Auricchio A, Hildinger M, O’Connor E, Gao GP and Wilson JM: Isolation of highly infectious and pure adeno-associated virus type 2 vectors with a single-step gravity-flow column. Hum Gene Ther 12: 71-76, 2001.

26. Berditchevski F and Odintsova E: Characterization of integrin-tetraspanin adhesion complexes: role of tetraspanins in integrin signaling. J Cell Biol 146: 477-492, 1999.

27. Fitter S, Sincock PM, Jolliffe CN and Ashman LK: Transmembrane 4 superfamily protein CD151 (PETA-3) associates with beta1 and alphaIIb beta3 integrins in haemopoietic cell lines and modulates cell-cell adhesion. Biochem J 338: 61-70, 1999.

28. Conner SD and Schmid SL: Regulated portals of entry into the cell. Nature 422: 37-44, 2003.

29. Carmeliet P: Mechanisms of angiogenesis and arteriogenesis. Nat Med 6: 389-395, 2000.

30. Rey S and Semenza GL: Hypoxia-inducible factor-1-dependent mechanisms of vascularization and vascular remodelling. J Endocrinol 1829-1843, 2000.

31. Ulrich F and Heisenberg CP: Trafficking and cell migration. Cardiovasc Res 86: 236-242, 2010.

32. Ulrich F and Heisenberg CP: Trafficking and cell migration. Traffic 10: 811-818, 2009.

33. Mellman I and Nelson WJ: Coordinated protein sorting, targeting and distribution in polarized cells. Nat Rev Mol Cell Biol 9: 835-845, 2008.

34. Conner SD and Schmid SL: Regulated portals of entry into the cell. Nature 422: 37-44, 2003.

35. Carmeliet P: Mechanisms of angiogenesis and arteriogenesis. Nat Med 6: 389-395, 2000.

36. Rey S and Semenza GL: Hypoxia-inducible factor-1-dependent mechanisms of vascularization and vascular remodelling. Cardiovasc Res 86: 236-242, 2010.

37. Ulrich F and Heisenberg CP: Trafficking and cell migration. Traffic 10: 811-818, 2009.

38. Mellman I and Nelson WJ: Coordinated protein sorting, targeting and distribution in polarized cells. Nat Rev Mol Cell Biol 9: 835-845, 2008.

39. Conner SD and Schmid SL: Regulated portals of entry into the cell. Nature 422: 37-44, 2003.

40. Carmeliet P: Mechanisms of angiogenesis and arteriogenesis. Nat Med 6: 389-395, 2000.

41. Rey S and Semenza GL: Hypoxia-inducible factor-1-dependent mechanisms of vascularization and vascular remodelling. Cardiovasc Res 86: 236-242, 2010.

42. Ulrich F and Heisenberg CP: Trafficking and cell migration. Traffic 10: 811-818, 2009.

43. Mellman I and Nelson WJ: Coordinated protein sorting, targeting and distribution in polarized cells. Nat Rev Mol Cell Biol 9: 835-845, 2008.