Perilipin+ embryonic preadipocytes actively proliferate along growing vasculatures for adipose expansion

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ABSTRACT
Despite the growing interest in adipose tissue as a therapeutic target of metabolic diseases, the identity of adipocyte precursor cells (preadipocytes) and the formation of adipose tissue during embryonic development are still poorly understood. Here, we clarified the identity and dynamic processes of preadipocytes in mouse white adipose tissue during embryogenesis through direct examination, lineage tracing and culture systems. Surprisingly, we found that lipid-lacking but perilipin+ or adiponectin+ proliferating preadipocytes started to emerge at embryonic day 16.5, and these cells underwent active proliferation until birth. Moreover, these preadipocytes resided as clusters and were distributed along growing adipose vasculatures. Importantly, the embryonic preadipocytes exhibited considerable coexpression of stem cell markers, such as CD24, CD29 and PDGFRα, and a small portion of preadipocytes were derived from PDGFRα+ mural cells, in contrast to the adult preadipocytes present in the stromal vascular fraction. Further analyses with in vitro and ex vivo culture systems revealed a stepwise but dynamic regulation of preadipocyte formation and differentiation during embryogenesis. To conclude, we unraveled the identity and characteristics of embryonic preadipocytes, which are crucial for the formation and expansion of adipose tissue during embryogenesis.

KEY WORDS: Adipogenesis, Adipose tissue, Angiogenesis, Preadipocytes

INTRODUCTION
The worldwide epidemic of obesity over the past half century has raised a surge of interest in the understanding of the origin, development, integrity and remodeling of adipose tissues (Rosen and MacDougald, 2006; Gesta et al., 2007; Cao, 2010; Wang et al., 2013). Two types of adipose tissues, which are different in structure and function, are found in mammals as white adipose tissue (WAT) and brown adipose tissue (BAT). WAT regulates energy storage and mobilization in the form of triacylglycerides, and has paracrine and endocrine functions via secretion of adipokines and growth factors, whereas BAT is specialized as an energy dissipater for heat during cold- and diet-induced thermogenesis. Recent reports (Wu et al., 2012; Rosenwald et al., 2013; Nedergaard and Cannon, 2014) reveal that WAT and BAT are interchangeable in certain pathophysiological situations, named ‘browning’ (WAT to BAT) and ‘whitening’ (BAT to WAT), indicating that adipose tissues are dynamic, versatile and communicative tissues. Nevertheless, the origin and development of adipose tissues are poorly understood.

Expansion of WAT during postnatal growth or obesogenic change in adults is primarily caused by appropriate or excessive filling of lipid into post-mitotic adipocytes, together with, to a lesser extent, an ongoing proliferation and differentiation of preadipocytes, which are predominantly localized adjacent to adipose vasculatures (Nishimura et al., 2007; Li et al., 2011; Tran et al., 2012). By contrast, formation of WAT during embryonic development is assumed by serial processes of commitment of putative adipoblasts, mainly from mesoderm, and proliferation and differentiation of adipoblasts into adipocyte precursor cells (preadipocytes) (Gesta et al., 2007; Tang et al., 2008). These processes have been supposed to appear along the growing adipose vasculature, which is termed the ‘adipose niche’ (Crandall et al., 1997; Cho et al., 2007; Birsoy et al., 2011; Han et al., 2011). Although there have been attempts to characterize the distinct cellular intermediates (adipoblasts and preadipocytes) between mesodermal cells and differentiated adipocytes, such studies have been hampered by the lack of specific cell surface markers to identify and prospectively isolate them in vivo. A recent genetic approach using a lineage-tracing mouse model, the ‘AdipoChaser mouse’, demonstrated that most adipocytes could be committed and differentiated from their putative progenitors before birth (Wang et al., 2013). However, lineage tracing does not reveal details about phase timing and developmental cues. Moreover, changes in phenotypes in the long-term bring into question the ability to reliably identify phenotypic fingerprints based on observations at a single time point. Therefore, considering that substantial changes in phenotypes could be generated at an early time point by genetic changes and manipulations, it is valuable to observe embryonic tissues directly.

In this study, through direct examination of the embryonic inguinal fat pad, we explored the dynamic processes of identification and development of preadipocytes in inguinal WAT during embryogenesis. Unexpectedly, we found that the preadipocytes exist as perilipin+ and adiponectin+ cells either as lipid-lacking cells or small-sized lipid-containing cells, which are distributed as clusters along growing adipose vasculatures and are also highly proliferative, particularly at the periphery of the adipose cluster.

RESULTS
Emergence of perilipin+ preadipocytes during embryogenesis
To explore the origin of preadipocytes in vivo, we attempted to identify the formation of WAT during murine embryonic development in the inguinal region, which is easily accessible

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compared with other regions, as well as being easily distinguishable by a developing lymph node at the center of a tripod-shaped vascular branch (Fig. 1A). As no specific markers for adipoblasts/preadipocytes are available, and there also is a possibility that a mixed population of adipoblasts/preadipocytes and perilipin 1+ matured adipocytes could exist in the developing adipose tissue, we traced prenatal adipogenesis by immunostaining of perilipin 1 (Plin1, hereafter called perilipin) in whole-mounted inguinal WAT (Fig. 1B; supplementary material Fig. S1A). At embryonic day (E) 14.5, neither apparent adipose tissue nor perilipin+ cells were observed (Fig. 1B). However, from E16.5, de novo lipid-lacking perilipin+ cells that were attached mainly along the CD31+ (Pecam1 – Mouse Genome Informatics) blood vessels started to appear (Fig. 1B). Remarkably, after two days, two different types of perilipin+ cells, either lipid-lacking or scant lipid droplet-containing cells that were stained with boron-dipyrromethene (BODIPY), were concomitantly increased and congregated into the shape of a cluster along the blood vessels (Fig. 1B,C). Moreover, the size of these clusters expanded profoundly without significant increases in cell sizes but with marked increase in the number of perilipin+ cells, which still had either no or small-sized lipid droplets, until postnatal day (P) 0 (Fig. 1C-F). These findings indicate that hyperplasia of perilipin+ cells was in progress in these clusters during the prenatal period. By contrast, lipid and other nutrients provided by nursing after birth (Rudolph et al., 2007) caused every perilipin+ cell to expand and change into lipid-filled perilipin+ adipocytes (Fig. 1C,D).

Crucial interactions between PPA and their supporting vascular network during adipogenesis

Because PPA were relatively abundant along the distinct vascular networks but rarely existed in avascular regions in the adipose clusters at E16.5 (supplementary material Fig. S2A), we speculated that vasculatures provide an external cue for fate determination and differentiation of putative adipoblasts into PPA. To test our hypothesis, we generated endothelial Vegfr2-depleted mice.
(VR2ΔEC) by intercrossing endothelial cell-specific VE-cadherin-Cre-ERT2 and VEGFR2+/fl mice, which enabled vascular disruption upon tamoxifen administration from E16.5. At E18.5, tamoxifen-treated VR2ΔEC embryos showed hemorrhages and edema, indicating that these mice underwent effective vascular disruption owing to the loss of VEGFR2 in growing blood vessels, including inguinal adipose vasculature (Fig. 2A-C). In the inguinal region of these mice, the development of clusters was no longer prominent, and the number of PPAs per cluster had reduced by 49% compared with control embryos (Fig. 2C,D). Accordingly, the gene expression of Vegfr2 and Pparg2 (Pparg – Mouse Genome Informatics) in the WAT of VR2ΔEC decreased by 77% and 62%, respectively (Fig. 2E,F). These findings demonstrate that endothelial-specific Vegfr2 depletion was effective enough to induce vascular disruption, leading to the interruption of in vivo generation of PPA during adipogenesis.

Conversely, we generated Col1a1-Cre/inducible COMP-Ang1-Tg (Ang1OE) mice that have connective tissue-driven overexpression of angiopoietin 1 (Ang1; Angpt1 – Mouse Genome Informatics) for a gain-of-function study on the developing adipose vasculatures. The gross appearance of E18.5 embryos of Ang1OE was reddish compared with that of control embryos, suggesting increased blood vessel density and vessel enlargement (Fig. 2G). In the inguinal region, Ang1OE embryos exhibited concomitantly expanded vascular network and clusters, and had a 2.1-fold increase in the number of PPAs per cluster (Fig. 2H-J). Moreover, the WAT of these embryos had 3.4-fold and 2.1-fold increases in the gene expression of Vegfr2 and Pparg2 compared with control embryos, respectively (Fig. 2K,L), indicating that adipose vasculature has a crucial relationship with in vivo adipogenesis during embryogenesis. Together, the embryonic distinct adipose vasculatures provide an external cue for the development of PPA and adipose tissue formation in a spatiotemporal manner.

**Temporal but distinct changes of proliferative ability in PPA**

To investigate whether active proliferation of PPA is an essential process for adipose growth during the prenatal period, we first stained the inguinal adipose tissues with a proliferation marker, PH3. In fact, a fair number of PH3+ cells that were noticeably matched with PPAs were observed from E16.5 to P0 (Fig. 3A; supplementary material Fig. S2B). Moreover, orthotopic-dissection image analysis revealed that most PH3+ PPAs were located at the periphery of the cluster at E18.5 (Fig. 3B). Second, using an alternative proliferation detection method, in vivo 5-ethynyl-2′-deoxyuridine (EdU) labeling, we confirmed that PPAs were actively proliferating at E18.5 (Fig. 3C; supplementary material Fig. S2C). Intriguingly, quantification revealed that the proportion of PH3+ cells in the PPA population in the cluster increased as the embryo got closer to birth (Fig. 3A,D). Furthermore, these findings were consistent with the observed changes in vascular network and clusters (Fig. 2).

**Fig. 2. Crucial interactions between PPAs and their supporting vascular network in adipogenesis.** (A-L) The supporting vascular network was disrupted in VR2ΔEC mice (A-F), whereas vascularity was expanded in Ang1OE mice (G-L). Unless otherwise denoted, n=4 per group. *P<0.05 versus control.

(A,B) Photographs showing gross features and inguinal WAT at E18.5. Scale bars: 1 mm. (C) Images of clusters in WAT stained for perilipin and CD31. Scale bars: 100 μm. (D) Comparison of the number of perilipin+ cells per cluster at E18.5. (E,F) Quantitative real-time PCR analyses of Vegfr2 and Pparg2 in WAT at E18.5. n=5 per group. (G,H) Photographs showing gross features and inguinal WAT of control and Ang1OE at E18.5. Arrows indicate a reddish skin color due to increased vascularity and vessel enlargement. Scale bars: 1 mm. (I) Images of clusters in WAT of control and Ang1OE at E18.5 stained for perilipin and CD31. Scale bars: 100 μm. (J) Comparison of the number of perilipin+ cells per cluster at E18.5. (K,L) Quantitative real-time PCR analyses of Vegfr2 and Pparg2 in WAT at E18.5. Error bars represent s.d.
consistent with the gene expression of another proliferation marker, Ki67 (Mki67 – Mouse Genome Informatics), in the inguinal WAT (Fig. 3E). However, the proportion of the proliferating cells decreased sharply when its cell size expanded, due to apparent lipid-fillings (Fig. 3A,D). To confirm this phenomenon in vitro, 3T3-L1 cells were cultured with adipogenic differentiation medium and observed at an early differentiation period. Consistently, perilipin+ PPARY+ PH3+ cells were detected at day 3 after adipogenic differentiation (supplementary material Fig. S2D), which was similar to the in vivo findings. To confirm this finding, we used a live-imaging device and were able to observe dividing cells containing scant lipid droplets that were stained with BODIPY in 3T3-L1 cells (supplementary material Fig. S2E). Furthermore, E18.5 WAT of VR2<sup>EC</sup> embryos showed a reduction in the number of PH3+ cells in the PPA population by 70% compared with control embryos (Fig. 3F,G), indicating that actively growing vasculature is required for proper development of WAT through promotion of PPA proliferation.

**Embryonic preadipocytes are different from adult preadipocytes**

Lipid-lacking PPAs and adult preadipocytes can be enriched as adipose stromal vascular fraction (SVF), a heterogeneous mixture of cells operationally defined by enzymatic dissociation of WAT followed by density separation from lipid-retaining adipocytes (Han et al., 2011). To evaluate the differences between PPAs and adult preadipocytes, we first performed a colony-forming unit fibroblast (CFU-F) assay in WAT from E18.5 and adult inguinal WAT. Both SVFs could produce fibroblast colonies, but the number of colonies from E18.5 WAT was higher than that from adult WAT, whereas the sizes of adult colonies were relatively larger than those of E18.5 (Fig. 4A-C). Moreover, flow cytometry analyses with surface markers of mesenchymal and adipose stem cells [CD24, CD29 (Itgb1), CD34, Sca-1 (Ly6a) and PDGFRα] on a lineage-negative population [CD31<sup>-</sup> CD45<sup>-</sup> (Ptprc Ter119<sup>-</sup> (Ly76)] revealed that CD24-, CD29-, CD34- and PDGFRα-expressing cells were highly enriched in the SVF of E18.5 WAT compared with the SVF of adult WAT (Fig. 4D,E). Our findings regarding the differences in preadipocyte characteristics between embryos and adults are in line with a recent report addressing two distinct adipose progenitor populations: developmental and adult (Jiang et al., 2014). As it was reported that the majority of adipose progenitors are derived from the PDGFRβ<sup>-</sup> mural cell compartment (Tang et al., 2008), we were curious about whether PPAs could share their origin and identity with the PDGFRβ<sup>+</sup> mural cell. To trace PDGFRβ<sup>+</sup> adipose progenitors, we intercrossed Pdgfrb-Cre-ERT2 and reporter tdTomato (supplementary material Fig. S3A,B) or mT/mG mice, and then administrated tamoxifen into a pregnant female at different time points (E10.5, E13.5 and E16.5) and analyzed inguinal WAT at E18.5 (Fig. 4F). We found that a subset of embryonic PDGFRβ<sup>+</sup> mural cells already had adipogenic potential. In contrast to the gradual increase in the number of GFP<sup>-</sup> pericytes or stromal cells during embryogenesis, the WAT of the embryos that were injected with tamoxifen at E13.5 had the highest number of GFP<sup>+</sup> cells in the perilipin<sup>+</sup> cell population at E18.5 (Fig. 4G-I), indicating that the conversion from PDGFRβ<sup>+</sup> cells to adipose-lineage cells either takes a few days or only occurs during specific periods. Despite this, and contrary to a previous study (Tang et al., 2008), the majority of embryonic preadipocytes inside the clusters were not derived from the PDGFRβ<sup>-</sup> mural cell compartment (Fig. 4G-I). These findings suggest that embryonic preadipocytes in WAT are derived from several subsets of mesenchymal cells.

**Stepwise but dynamic regulation of preadipocyte formation and differentiation during embryogenesis**

To gain insight into the detailed characteristics of PPAs, we harvested and cultured the SVF of inguinal WAT obtained at each prenatal period. Under adipogenic differentiation medium, SVFs
were evidently differentiated into Oil Red O+ lipid-filling cells at day 7. This adipose-lineage commitment started from E14.5, and robustly occurred after E16.5 (Fig. 5A,B). Quantitative real-time PCR analyses on the expression of key adipogenic molecules and markers in the SVF indicated that the gene expression levels of preadipocyte/adipocyte markers, such as Cebpb, Pparg2 and Cebpa, increased markedly, showing the highest peak at E18.5, and the expression of Plin1 elevated significantly after P0 (Fig. 5C). Consistent with the gene expression patterns, immunostaining for adipogenic transcription factors inside the cluster showed prominent PPARγ and C/EBPα protein expression after E16.5 (Fig. 5D,E). Interestingly, most PPARγ+ or C/EBPα+ cells already co-expressed perilipin (Fig. 5D,E), implying that this adipogenic transcriptional cascade actively occurs in the PPAs several days before birth. Moreover, the deposition of laminin-containing basement membrane was not distinctly found in PPAs at E16.5, whereas it was distinct on the margin of PPAs at P0 (Fig. 5F). To characterize PPAs further, we performed ex vivo culture of clusters derived from embryonic WAT. Plating the clusters induced cell proliferation around the cluster within a few days (supplementary material Fig. S4A). By Oil Red O staining, we noticed distinct cell populations inside and outside of the cluster. Because the cells inside the cluster were stained with Oil Red O without adipogenic differentiation medium, Oil Red O+ cells were regarded as nascent adipocytes that accumulated lipid during the culture (supplementary material Fig. S4A). However, as Oil Red O+ cells outside the cluster were present only under adipogenic differentiation medium (supplementary material Fig. S4A), these migrated Oil Red O+ cells outside the cluster were regarded as preadipocytes that were differentiated into adipocytes when cultured with adipogenic differentiation supplementation. Surprisingly, some of the migrated cells outside the cluster without adipogenic differentiation medium were stained with perilipin and PH3 (supplementary material Fig. S4B). Taken together, these findings revealed that the embryonic preadipocytes with proliferative and migrative capacity express perilipin, which is again contrary to the notion that perilipin is exclusively expressed by differentiated adipocytes.
Taken together, these findings indicate that there is a stepwise regulation of preadipocyte formation and differentiation during embryogenesis.

The adiponectin+ preadipocytes are almost identical to PPAs

To validate our findings, we evaluated the expression of adiponectin, which is a selective marker for adipocyte differentiation as well as a well-known adipokine, in PPAs of adiponectin-Cre/tdTomato (Ad-Tomato) mice. In Ad-Tomato embryos, tdTomato signal was highly detected in the inguinal WAT at E18.5, and PH3+ proliferating PPAs were frequently detected in the adiponectin+ PPA population at the inside of clusters (Fig. 6A). By immunostaining the cultured PPAs of Ad-Tomato embryos, we found a mixed population of fibroblast-like shaped adiponectin+ integrin α5high preadipocytes and round shaped adiponectin+ integrin α5low adipocytes, which were distinguishable by morphology (Fig. 6B). A subset of cultured adiponectin+ cells were perilipin+ and small lipid droplet-containing preadipocytes, and were also PPARγ+ and PH3+ proliferating preadipocytes (supplementary material Fig. S5A,B). Furthermore, like perilipin+ SVF cells (supplementary material Fig. S6, S7), more than 50% of adiponectin+ SVF cells also co-expressed CD24, CD29, CD34, Sca-1 and PDGFRα (Fig. 6C,D).

Next, we investigated the differences in the functional capacity between adiponectin+ preadipocytes and adiponectin− SVF cells. After digesting E18.5 WAT of Ad-Tomato embryos, we harvested and plated the SVF cells for 1 day to eliminate any existing hematopoietic cells (Gupta et al., 2012). Flow cytometry analysis showed two distinct populations of adiponectin+ cell and adiponectin− cells by the intensity of tdTomato signal; 10.3% of cultured SVF cells were adiponectin+ cells (Fig. 6E). We then sorted these two populations and plated and cultured them with adipogenic differentiation medium. The gene expression levels of tdTomato and adiponectin confirmed that sorted tdTomato+ SVF cells showed enrichment for adiponectin (Fig. 6H). As we expected, most of the sorted tdTomato+ SVF cells expressed strong tdTomato signal with a fibroblast-like cell shape in the undifferentiated state, suggesting that they are adiponectin+ preadipocytes (Fig. 6F). During adipogenic differentiation, tdTomato+ preadipocytes were robustly differentiated into Oil Red O+ adipocytes, to a greater extent than adiponectin− cells (Fig. 6G; supplementary material Fig. S5C,D).
The quantitative differences of adipocytes between the two populations were matched with the gene expression levels of Plin1 (Fig. 6H). Moreover, the gene expression levels of preadipocyte/adipocyte markers, such as Pparg2 and Cebpa, displayed markedly increased patterns from tdTomato+ embryonic SVF cells in both the undifferentiated and differentiated states (Fig. 6H). Thus, the adiponectin+ cells are almost identical to PPAs. Collectively, these data indicate the similarity between perilipin+ cells and adiponectin+ cells as embryonic preadipocytes.

**DISCUSSION**

In this study, we demonstrate that embryonic perilipin+ preadipocytes in WAT proliferate to form clusters and interact with growing adipose vasculature (which is at odds with the previously held belief that perilipin is expressed exclusively in differentiated adipocytes) and that they proliferate robustly within a few days during the prenatal period. In turn, generation of a massive quantity of PPAs during the prenatal period could be a prerequisite to building a certain size of WAT rapidly with fully lipid-filled, differentiated adipocytes by consumption of lipid-rich nutrients such as milk during the postnatal period.

Accumulating evidence demonstrates that adipose tissue expansion and remodeling is angiogenesis dependent during the postnatal period and adulthood (Rupnick et al., 2002; Cao, 2007; Cho et al., 2007; Sun et al., 2011). The closed structural relationship between adipocytes and vasculature facilitates constant and immediate responses to various conditions to maintain metabolic homeostasis. In addition to its basic roles, the adipose vasculature has important roles in providing hormones, growth factors, inflammation-related cytokines, and stem cells to adipose tissues (Nishimura et al., 2009; Christiaens and Lijnen, 2010; Cao, 2013). Our findings indicate that the role of the growing vasculature as an adipose niche applies also to the growth of PPAs as a cluster during embryogenesis.

Preadipocytes enter the cell cycle, go through one or two rounds of replication that is called mitotic clonal expansion, and proceed to terminal differentiation in vitro (Tang et al., 2003). However, little is known about whether this event is a prerequisite for adipogenic
Moreover, we discovered that a small number of embryonic (CD24, CD29, CD34 and PDGFRα) observed differences in the expression of stem cell markers after birth (Berry and Rodeheffer, 2013). Likewise, we also expression of CD24, one of the stem cell markers, in preadipocytes differentiated adipocyte markers resides in the embryonic WAT. Together, we can conclude that a subset of preadipocytes expressing differentiated adipocyte markers or adipocytes expressing proliferative markers in embryonic WAT.

To characterize the observed PPAs in more detail, we collected evidence to support our hypothesis. First, both cultured PPAs and adiponectinα-peripilinα preadipocytes evidently showed a typical fibroblast-like shape and robust adipogenic differentiation under adipogenic medium differentiation. Second, partially differentiated human preadipocytes have been shown to still be capable of cell division (Prins and O’Rahilly, 1997; Gregoire et al., 1998). These reports are in line with our findings that a subset of PPAs, without noticeable lipid droplets, also divided under adipogenic differentiation. Third, the differences in laminin deposition and integrin α5 expression in PPAs between E16.5 and P0 indicate the onset of prenatal adipogenesis, which is usually defined by extracellular matrix remodeling and characterized by the conversion from the fibronectin-rich stromal matrix to a laminin-rich basement membrane (Liu et al., 2005; Hoshiba et al., 2010). Together, we can conclude that a subset of preadipocytes expressing differentiated adipocyte markers resides in the embryonic WAT.

A recent report demonstrated that there is a decrease in the expression of CD24, one of the stem cell markers, in preadipocytes after birth (Berry and Rodeheffer, 2013). Likewise, we also observed differences in the expression of stem cell markers (CD24, CD29, CD34 and PDGFRα) between E18.5 and adult. Moreover, we discovered that a small number of embryonic preadipocytes was derived from the PDGFRαβ mural cell compartment by using Pdgfrb-Cre-ER T2 reporter mice, in contrast to a previous report that observed lacZ expression in P30 WAT under the control of Pdgfrb-Cre (Tang et al., 2008). Considering recent studies stating that region, sex, strain and age can affect the gene expression and physiological phenotypes of adipose tissue as well as the quantity of adipose progenitors (Yamamoto et al., 2010; Macotela et al., 2012; Chau et al., 2014), we also propose that the preadipocyte populations in WAT of embryos and adults are different in character, and even origin.

Like stem cell markers such as CD24, CD29 and CD34, perilipin has been used as marker of prenatal preadipocytes for this study. As perilipin knockout mice showed reduced body weight and resistance to diet-induced obesity (Tansey et al., 2001), which indicates alterations in metabolic homeostasis, elaborative studies on time- and tissue-specific ablation of perilipin by genetically modified mice will be required to understand the roles of perilipin in preadipocytes.

In conclusion, we demonstrate that the expression of perilipin or adiponectin is not confined to terminally differentiated adipocytes. Rather, embryonic preadipocytes expressing these markers proliferate in order to expand the adipose-lineage cell population. Although subsequent studies are required to explore the underlying mechanisms, we highlight the role of embryonic preadipocytes and the prenatal period in the control of in vivo adipogenesis, which we believe will subsequently open new directions to exploring the cause of obesity.

**MATERIALS AND METHODS**

**Animals**

Specific pathogen-free C57BL/6J, Adiponectin-Cre and tdTomato mice were purchased from Jackson Laboratory; Pdgfrb-Cre-ER T2 transgenic mice used in this study contained a BAC construct of Pdgfrb and the tamoxifen-inducible Cre recombinase (Feil et al., 1997; Stanczuk et al., 2015; R. H. Adams, unpublished). VEGFR2α- and PDGFRαβ (Albuquerque et al., 2009) mice were kindly provided by Professor Masanori Hirashima (Kobe University, Japan). VE-Cadherin-Cre-ER T2 (Wang et al., 2010), Coll1α-Cre (Liu et al., 2004), inducible COMP-Angl-Tg (Lee et al., 2013) and mT/mG (Muzumdar et al., 2007) mice were transferred and bred in our specific pathogen-free animal facilities at KAIST. Tamoxifen (Sigma-Aldrich) was dissolved in corn oil and administrated into the peritoneal cavity of pregnant females (3 mg/mouse) at the indicated time point.

Embryos in this study were generated by timed mating, and E0.5 was regarded as the day of prominent plug formation. Mice from E12.5 to 2-month-old adults were used for this study. All mice were bred in our specific pathogen-free animal facility and were fed normal chow diet (NCD) (PMI LabDiet) with ad libitum access to water. Animal care and experimental procedures were performed under the approval from the Institutional Animal Care and Use Committee (No. KA2013-37) of KAIST.

**Histological analysis**

Mice were anesthetized with a combination of ketamine (80 mg/kg) and xylazine (12 mg/kg) by intramuscular injection. All adipose tissues were from male adult mice, or from embryos or pups regardless of gender. Harvested tissues were fixed with 4% paraformaldehyde in PBS for 1 h, and whole-mounted or cryo-embedded followed by cryosectioning. After blocking with 5% goat or donkey serum (Jackson ImmunoResearch) in PBST (0.3% Triton X-100 in PBS for whole-mount method, 0.03% Triton X-100 in PBS for cryosections) for 1 h at room temperature, whole-mounted or sectioned tissue was incubated overnight with primary antibodies (supplementary material Table S1) at 4°C. After several washes with PBST, the samples were incubated with the following secondary antibodies (supplementary material Table S2) for 2 h at room temperature: Cy3-, Cy5- or FITC-conjugated goat anti-guinea pig antibody; Cy3- or Cy5-conjugated goat anti-hamster antibody; Cy3-, Cy5- or FITC-conjugated goat anti-rabbit antibody; Cy3- or FITC-conjugated goat anti-rat antibody; and Cy3- or FITC-conjugated donkey anti-goat antibody. Hoechst 33342 (Sigma-Aldrich) was used to detect nuclei, and boron-dipyrromethene (BODIPY, Invitrogen) was used to detect lipid droplets in adipocytes. For fluorescence labeling, prepared mice were injected intraperitoneally with 5-ethyl-2'-deoxyuridine (EdU; 2 nmol/mouse; Invitrogen) and sacrificed after the indicated interval. EdU was detected according to the manufacturer’s instructions. For cell staining, cells at the indicated time points were fixed with 4% paraformaldehyde in PBS for 1 h, blocked with 5% goat serum in PBST, and stained with antibodies (supplementary material Table S1). To label proliferating cells, 3T3-L1 cells were incubated with 10 μM EdU (Invitrogen) for 3 h and processed for EdU detection and staining.

**Morphometric analysis**

Confocal microscopes (ApoTome, LSM 510 and LSM 780, Carl Zeiss) equipped with argon and helium-neon lasers were used to visualize fluorescence images. Morphometric analyses were performed with Image J software (NIH) or Zeiss image software (ZEN 2012).

**In vitro culture of 3T3-L1 cell line**

The 3T3-L1 cell line was maintained in culture medium, which is a mixture of 10% calf serum and 100 μg/ml streptomycin in Dulbecco’s modified Eagle medium (DMEM). In order to induce adipogenic differentiation, 2-day-postconfluent cells were incubated with adipogenic differentiation medium.
[DMEM containing 10% fetal bovine serum, 5 μg/ml insulin (Sigma-Aldrich), 0.5 mM 3-isobutyl-1-methylxanthine (IBMX, Sigma-Aldrich) and 1 μM dexamethasone (Sigma-Aldrich)]. After 3 days, adipogenic differentiation medium was replaced with maintenance medium (DMEM containing 10% fetal bovine serum and 5 μg/ml insulin).

**In vitro culture of SVF and ex vivo culture of clusters**

Dissected inguinal WAT samples were chopped with scissors and incubated in digestion medium containing DMEM/F12 with 5 μg/ml insulin (Sigma-Aldrich), 100 U/ml hyaluronidase (Sigma-Aldrich), 10 ng/ml EGF (Sigma-Aldrich), 20 ng/ml cholera toxin (Sigma-Aldrich), 5% FBS, 500 ng/ml hydrocortisone (Sigma-Aldrich), 100 μg/ml streptomycin and 300 μM collagenase-1A (Sigma-Aldrich) for 1 h at 37°C. The resulting suspension was dissociated with 0.5 ml 0.25% trypsin/EDTA (Life Technologies) for 1.5 min at 37°C, and sequentially re-suspended in 0.1 mg/ml DNase for 5 min at 37°C. Erythrocytes were lysed with ACK Lysing Buffer (Life Technologies) for 5 min at 37°C and centrifuged at 470 g for 5 min. After several washes, the SVF pellet was re-suspended in culture medium. Two days after confluence was observed, SVF cells were cultured with adipogenic differentiation medium. After 3 days, adipogenic differentiation medium was changed to maintenance medium. For ex vivo culture of clusters, dissected adipose tissues derived from E18.5 embryos were incubated with HBSS containing 0.2% collagenase type II for 10 min at room temperature. Using sharp scissors under the microscope, clusters then were excised, trimmed, and plated in culture containing DMEM supplemented with 10% bovine serum.

**Colony-forming unit fibroblast (CFU-F) assay**

To quantify the proliferative activity of mesenchymal stem cells, we sorted lineage-negative SVF cells (CD31− CD45− Ter119−) from E18.5 and adult (2 months old) subcutaneous adipose tissue using FACSaria II (BD Biosciences) with the listed antibodies (supplementary material Table S1). Sorted cells (150, 300 or 450 per 100-mm plate) were seeded with culture medium. Two days after confluency was observed, SVF cells were cultured with adipogenic differentiation medium. After 3 days, adipogenic differentiation medium was changed to maintenance medium. For SVF culture of clusters, dissected adipose tissues derived from E18.5 embryos were incubated with HBSS containing 0.2% collagenase type II for 10 min at room temperature. Using sharp scissors under the microscope, clusters then were excised, trimmed, and plated in culture containing DMEM supplemented with 10% bovine serum.

**Live cell imaging**

3T3-L1 or SVF cells were cultured in 6-well plates with or without adipogenic differentiation. Using an incubator chamber (Live Cell Instrument) equipped with optimal environment settings of 5% CO₂ and 37°C, live cell imaging was performed and recorded at 5 or 10 min intervals by microscope (ApoTome, Carl Zeiss) equipped with argon and helium-neon lasers.

**Flow cytometry analysis**

SVF cells of E16.5, E18.5, P0, P2 and adult (2 months old) were harvested by digestion of subcutaneous adipose tissue. SVF was sequentially filtered through 70- and 40-μm filters. For intracellular staining, Cytofix/Cytoperm kit (BD Biosciences) was used according to the manufacturer’s instructions. Before incubation with antibodies, Fc receptors were blocked with CD16/CD32 antibody for 10 min to minimize non-specific binding. Cells were incubated with antibodies at 4°C for 30 min. Following antibody incubation, samples were washed, centrifuged at 300 g for 3 min and resuspended in PBS containing 5% calf serum (see supplementary material Table S1 for antibodies). The lineage-negative population was analyzed with FACSaria II (BD Biosciences). Data analysis was performed with FlowJo software (Tree Star).

**Oil Red O staining**

The cells in the wells were fixed with 4% paraformaldehyde for 40 min at room temperature. After being washed with PBS and 60% isopropanol, cells were incubated with filtered Oil Red O working solution for 50 min at room temperature. After staining, several washes with PBS and 60% isopropanol were performed to reduce non-specific staining. Images of stained cells were captured by a microscope equipped with a CCD camera (Carl Zeiss).

**X-gal staining**

Adipose tissues derived from VEGFR2+/lacZ mice were harvested, fixed with 4% paraformaldehyde in PBS for 1 h, and whole-mounted or cryo-embedded followed by cryosectioning. After several washes, whole or sliced tissues were incubated with X-gal working solution (Biosesang, Korea) at 37°C until galactosidase expression (indicated by blue color) was prominent. The staining was imaged using a microscope equipped with a CCD camera (Carl Zeiss).

**RNA extraction and quantitative real-time PCR**

Total RNA was extracted from adipose tissues with the RNeasy Plus Mini Kit (Qiagen) according to the manufacturer’s instructions. Then, the mRNA was reverse-transcribed into cDNA using the GoScript Reverse Transcription Kit (Promega). Quantitative real-time PCR was performed with the indicated primers using the Bio-RadTM CFX96 Real-Time PCR Detection System (Bio-Rad; see supplementary material Table S3 for primers). The PCR data were analyzed using Bio-Rad CFX Manager Software (Bio-Rad).

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