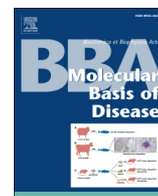




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## Review

Adipocyte lineages: Tracing back the origins of fat<sup>☆</sup>Joan Sanchez-Gurmaches, David A. Guertin<sup>\*</sup>

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## ABSTRACT

The obesity epidemic has intensified efforts to understand the mechanisms controlling adipose tissue development. Adipose tissue is generally classified as white adipose tissue (WAT), the major energy storing tissue, or brown adipose tissue (BAT), which mediates non-shivering thermogenesis. It is hypothesized that brite adipocytes (brown in white) may represent a third adipocyte class. The recent realization that brown fat exist in adult humans suggests increasing brown fat energy expenditure could be a therapeutic strategy to combat obesity. To understand adipose tissue development, several groups are tracing the origins of mature adipocytes back to their adult precursor and embryonic ancestors. From these studies emerged a model that brown adipocytes originate from a precursor shared with skeletal muscle that expresses Myf5-Cre, while all white adipocytes originate from a Myf5-negative precursors. While this provided a rational explanation to why BAT is more metabolically favorable than WAT, recent work indicates the situation is more complex because subsets of white adipocytes also arise from Myf5-Cre expressing precursors. Lineage tracing studies further suggest that the vasculature may provide a niche supporting both brown and white adipocyte progenitors; however, the identity of the adipocyte progenitor cell is under debate. Differences in origin between adipocytes could explain metabolic heterogeneity between depots and/or influence body fat patterning particularly in lipodystrophy disorders. Here, we discuss recent insights into adipose tissue origins highlighting lineage-tracing studies in mice, how variations in metabolism or signaling between lineages could affect body fat distribution, and the questions that remain unresolved. This article is part of a Special Issue entitled: Modulation of Adipose Tissue in Health and Disease.

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## 1. Introduction:

Obesity is a risk factor for many diseases including type 2 diabetes, cardiovascular disease, and cancer. Obesity occurs when energy intake exceeds energy expenditure causing adipose tissue to overgrow. However, body fat distribution patterns are highly variable between individuals and not all fat is equal with some depots having more favorable metabolic properties than others [1]. Some lipodystrophies also present as fat distribution disorders characterized by regional lipotrophy with other depots being spared or even expanding [2,3]. With a worldwide obesity epidemic well underway, the study of adipose tissue growth, distribution, and regulation is a major research focus. Yet the developmental origins of fat, the determinants of body fat distribution, and the signaling mechanisms that control fat growth remain poorly defined.

## 2. Adipose tissue is complex and heterogeneous

In mammals, fat is typically classified by morphological appearance as being either white adipose tissue (WAT) or brown adipose

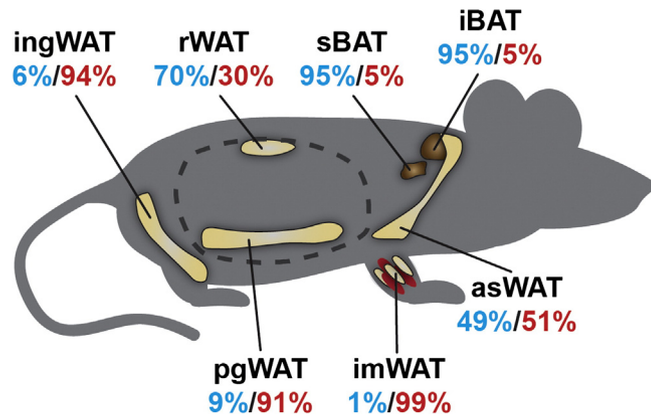
tissue (BAT). WAT (the primary site of energy storage) is mostly composed of adipocytes containing a large unilocular lipid droplet. WATs are found throughout the body; however, the distribution of mass between each depot varies in the population as a function of genetics, age, and for some depots, sensitivity to hormones and glucocorticoids. WAT location is often classified as being visceral (in the trunk cavity) or subcutaneous (below the skin). There is not a uniformly applied system for describing the anatomical location of each WAT depot and therefore depot names often vary between studies. According to [4], WAT depots in rodents include the anterior subcutaneous WATs (asWATs) including interscapular and axillary WAT (located in the scapular region), inguinal WAT (ingWAT; attached dorsally along the pelvis to the thigh of the hindlimb), perigonadal WAT (pgWAT; surrounding the uterus and ovaries in females and the epididymis and testes in males), retroperitoneal WAT (rWAT; located within the abdominal cavity along the dorsal wall of the abdomen behind the kidney but not attached to the kidney or mixed with perirenal brown fat), and mesenteric WAT (mWAT; lining the surface of the intestines) (Fig. 1). Although the primary function of WAT is energy storage, it also functions as an endocrine organ secreting hormones and cytokines such as leptin and adiponectin that regulate feeding and metabolism [5]. Epidemiological studies have found that the accumulation of visceral fat (determined by high waist-to-hip ratio) associates with metabolic disease (i.e. insulin resistance, type 2 diabetes, dyslipidemia,

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## Myf5<sup>+</sup> precursors Myf5<sup>neg</sup> precursors



**Fig. 1.** The Myf5 lineage contribution to the precursor pool in each fat depot varies with its anatomical location. The contribution of the Myf5 lineage to the adipocyte precursor cell compartment (defined as CD31<sup>-</sup>CD45<sup>-</sup>Terr119<sup>-</sup>CD29<sup>+</sup>CD34<sup>+</sup>Sca1<sup>+</sup> cells) was recently determined by lineage tracing with Myf5-Cre;R26R-YFP mice. More than 95% of the precursors in brown fat are labeled with Myf5-Cre. In the anterior subcutaneous WATs (including interscapular and axial WATs) nearly 50% of the precursors trace to Myf5<sup>+</sup> precursors, and in the rWAT (a visceral WAT), the Myf5 precursors give rise to approximately 70% of the adipocyte precursor cell pool. In contrast, the Myf5 lineage contributes very little to the adipocyte precursor pool in ingWAT and pgWAT, which are 90–95% Myf5<sup>neg</sup>. The contribution of the Myf5 lineage to the intramuscular adipogenic precursor pool (defined with slightly different cell surface markers) is very low. Dotted line indicates the abdominal cavity. References provided in the text.

hypertension, atherosclerosis, hepatic steatosis, and cancer) while the accumulation of subcutaneous WAT associates with improved insulin sensitivity and low risk for developing type 2 diabetes [6–14].

Brown adipocytes contain multiple smaller (multilocular) lipid droplets, are rich in mitochondria, and reside in depots that are highly innervated and vascularized. In rodents, BAT is located in discrete depots in interscapular (iBAT), sub-scapular (sBAT), and cervical (cBAT) regions of the upper anterior side of the trunk and neck (Fig. 1) [4]. BAT also grows around parts of the aorta and kidneys [15]. These depots are often called “classical” BAT to distinguish them from brown-adipocyte-like cells, called brite adipocytes, which reside within some WATs (discussed below). In contrast to energy storing white adipocytes, brown adipocytes are specialized to expend energy to generate heat in a process called adaptive thermogenesis [16]. BAT is stimulated by the sympathetic nervous system following exposure to cold temperature and regulates the acute non-shivering thermogenesis response as well as the adaptive cold acclimatization response following chronic cold exposure. Thermogenesis is mediated by uncoupling protein 1 (UCP1), which embeds in the inner mitochondrial membrane, and produces heat by dissipating the proton electrochemical gradient over the inter-mitochondrial membrane space without generating ATP [17]. BAT stores energy for thermogenesis as perilipin coated lipid droplets and glycogen granules [18,19] and upon stimulation rapidly increases glucose and FA uptake to replenish its supplies. The high glucose uptake capacity of BAT makes it readily detectable by <sup>18</sup>F-fluorodeoxyglucose positron emission tomography (FDG–PET) [17,20,21]. While long thought to be critical only in rodents and newborn humans, the recent realization that BAT functions in adult humans (made possible by its sensitivity to FDG–PET) [22–26] raises the possibility that therapeutically controlling BAT growth and/or energy expenditure could be a strategy to combat obesity [27–34].

Interest is also growing in a third potential class of adipocyte called a brite adipocyte (also known as a “beige”, “inducible brown”, or “recruitable brown” adipocyte) [19,21,35–41]. This mysterious type of adipocyte exists among classical white adipocytes and is morphologically indistinguishable from its neighboring white adipocytes in the basal or unstimulated state. However, upon stimulation by chronic cold exposure (or other mechanisms that mimic beta-adrenergic

stimulation) they become multilocular and begin expressing UCP1 [19,21,40,42]. The presence of brite adipocytes in the WAT of mice is not homogeneous. For example, many adipocytes in ingWAT or rWAT become multilocular and induce UCP1 following stimulation; however, only a few UCP1<sup>+</sup> brite adipocytes arise in pgWAT in the same mice [19,40]. Brite adipocyte content varies between mouse strains and correlates with overall strain sensitivity to high fat diet [43–47]. Whether brite adipocytes form by trans-differentiation of existing white adipocytes or arise from a unique preadipocyte lineage is under debate (discussed below) [19,40,41,48–51]. Perhaps the biggest unresolved issue pertaining to brite adipocytes is whether these cells actually contribute significantly to thermogenesis. For example, while UCP1 mRNA is induced in brite adipocytes several hundred-fold relative to white adipocytes (which barely expresses UCP1) [38,52,53], the total UCP1 expression is still an order of magnitude lower than that detected in brown adipose tissue [52]. In fact, the maximum thermogenic capacity of brite fat has been estimated at only ~10% of classical brown fat in mice. A debate is underway as to whether human brown fat is more similar to brite fat or to classical brown fat in mice. Early reports argue that what is currently being called human brown fat is more similar to murine brite fat than to classical murine brown fat [48,49]. However, recent studies that more extensively profile different layers of adipose tissue in newborns and adults reveals that humans have brown fat deposits—particularly in the neck—that have a classic brown fat signature [54,55]. Interestingly, there appears to be a gradient of fat cell types in the neck, with deep neck fat being classical BAT, intermediate cells possibly being more brite-like, and the most peripheral adipocytes being classical white adipocytes [55]. These findings are important considering the growing emphasis on developing therapeutic strategies to induce the “browning” of WAT in humans [35,56–59] because they suggest that the studies of classical brown fat in mice could also have important therapeutic implications in humans. The physiological significance and regulatory mechanisms of brown versus brite fat in humans clearly need to be determined.

Each individual fat depot is complex, composed not only of mature adipocytes but also of adipocyte precursor cells, fibroblasts, nerves, vascular cells, macrophages, and other cell types (collectively called the stromal vascular fraction or SVF). Adding to their complexity is the fact that adipose tissues are functionally heterogeneous. Although the sharpest functional divergence is clearly between energy storing WAT and energy-expending BAT, functional divergence is also evident between WATs best exemplified by the risk associated with excess visceral fat versus subcutaneous fat. However, such simple distinctions are oversimplifications because differences in lipogenic activity, cell dynamics, proliferative and differentiation capacity, and gene expression even between categorically similar WAT depots have been reported [60–67]. Heterogeneity likely exists even within a single WAT as suggested by studies showing that neighboring adipocytes can respond differently to genetic perturbations or drugs [68–72]. The existence of such heterogeneity within and between fat tissues suggests broad conclusions should not be based on a limited survey of select fat depots, and that each depot should be considered separately.

Does adipocyte tissue heterogeneity reflect different developmental origins of adipocytes, variances in the developmental cues a particular adipocyte may experience during differentiation, or extracellular influences on the mature adipocytes unique to the local cell or tissue environment? Answering these questions could lead to new anti-obesity therapies, improve the understanding of lipodystrophies, and increase the prospect of using adipocyte progenitor cells for cell-based therapeutics [17,73,74].

### 3. The origin of adipocytes

Central to understanding the complexities and heterogeneity of adipose tissue is to understand where the path to becoming an

**Table 1**  
Summary of recent lineage tracing studies of pre- and mature adipocytes.

|   | Genetic approach                                     | Promoter location   | Rationale   | Reported result   | References |
|---|--|---|---|---|------------|
| White adipocytes                                | <i>sox10-Cre;R26R-eYFP</i>                           | Transgene   | Sox10 expresses in pre- and migratory neural crest cells at all rostro-caudal levels  | Mature adipocytes in the salivary gland labeled positive; pgWAT and ingWAT labeled negative   | [103,139]  |
|   | <i>PPAR<math>\gamma</math>-tTA;TRE-Cre;R26R-LacZ</i> | Knock-in  | PPAR $\gamma$ is a nuclear receptor expressed in all tissues and is critical for adipogenesis   | ingWAT and rWAT positive; some positive SVF cells   | [82]       |
|   | <i>aP2-GFP</i>                                       | Transgene   | aP2 (FABP4) expresses in mature adipocytes and is a target of PPAR $\gamma$   | ingWAT positive   | [82]       |
|   | <i>aP2-Cre;R26R-LacZ</i>                             | Transgene   | Described above   | ingWAT and pgWAT positive   | [82]       |
|   | <i>PDGFR<math>\beta</math>-Cre;R26R-LacZ</i>         | Transgene   | PDGFR $\beta$ is expressed in vascular mural cells  | ingWAT and rWAT positive  | [82,140]   |
|   | <i>myf5-Cre;R26R-eYFP</i>                            | Knock-in  | Myf5 is a muscle differentiation transcription factor that is first expressed in the dermomyotome at E8.0 and expresses in adult satellite cells              | ingWAT and pgWAT negative   | [116,117]  |
|   | <i>wnt1-Cre;R26R-eYFP</i>                            | Transgene   | Restrictive marker of migrating neural crest cells  | Cephalic WAT positive; pgWAT and ingWAT negative. 70% of adipocyte precursors in cephalic WAT positive; all ingWAT adipocyte precursors negative  | [104,141]  |
|   | <i>LysM-Cre;R26R-LacZ</i>                            | Knock-in  | The lysozyme M is exclusively expressed in macrophages and granulocytes   | 5–20% of pgWAT adipocytes positive  | [110,111]  |
|   | <i>VE-cadherin-Cre;R26R-LacZ</i>                     | Knock-in  | VE-cadherin is expressed in endothelial cells, is involved in cell adhesion, and is essential for vascular system development                                 | ingWAT and pgWAT positive   | [96,98]    |
|   | <i>VE-cadherin-Cre;R26R-eGFP</i>                     | Knock-in  | Described above   | Subset of mature adipocytes in the pgWAT positive   | [96,98]    |
|   | <i>VE-cadherin-CreER<sup>T2</sup>;R26R-LacZ</i>      | Knock-in  | VE-cadherin-CreER <sup>T2</sup> is a tamoxifen-inducible version of the <i>VE-cadherin-Cre</i>  | Numerous adipocytes in the ingWAT, pgWAT and BAT positive   | [97,98]    |
|   | <i>Zfp423-GFP</i>                                    | Knock-in  | Zfp423 is a transcriptional regulator broadly expressed but essential for preadipocyte commitment   | ingWAT positive   | [95]       |
|   | <i>pax3-Cre;R26R-mTmG</i>                            | Knock-in  | Pax3 is a muscle differentiation transcription factor first expressed in dorsal neural crest and somites; it cooperates with Myf5 to drive muscle development | 50% of the adipogenic SVF cells from asWAT positive   | [123,124]  |
|   | <i>myf5-Cre;R26R-LacZ</i>                            | Knock-in  | Described above   | Mature adipocytes from asWAT and rWAT positive; few mature adipocytes from ingWAT and pgWAT positive; 50–60% of adipogenic SVF cells from asWAT and rWAT positive; <2% from the ingWAT and pgWAT positive | [83,117]   |
|   | <i>myf5-Cre;R26R-eYFP</i>                            | Knock-in  | Described above   | 50–70% of the adipocyte precursors from asWAT and rWAT positive; <10% of the adipocyte precursors from pgWAT and ingWAT positive  | [83,117]   |
| <i>vav1-Cre;R26R-mTmG</i>                       | Knock-in   | Vav1 is a proto-oncogene that expresses in the hematopoietic and lymphoid systems                                     | All adipocyte precursors and mature adipocytes from ingWAT, pgWAT, rWAT and mWAT negative   | [100,112]   |            |
| <i>tie2-Cre;R26R-mTmG</i>                       | Transgene  | Tie2 is an angiopoietin receptor specific to endothelial cells and some hematopoietic cells                           | All adipocyte precursors and mature adipocytes from ingWAT, pgWAT, rWAT and mWAT negative   | [100,142]   |            |
| <i>VE-cadherin-Cre;R26R-mTmG</i>                | Knock-in   | Described above   | All adipocyte precursors and mature adipocytes from ingWAT, pgWAT, rWAT and mWAT negative   | [96,100]  |            |
| <i>PDGFR<math>\alpha</math>-Cre;R26R-mTmG</i>   | Transgene  | PDGFR $\alpha$ is expressed in most mesenchymal cells can activate the MAPK, PI3K and PLC $\gamma$ signaling pathways | All adipocyte precursors and mature adipocytes from ingWAT, pgWAT, rWAT and mWAT positive   | [100,143]   |            |
| Brown adipocytes                                | <i>En1-Cre;R26R-LacZ</i>                             | Knock-in  | Engrailed 1 is a homeobox transcription factor that expresses in cells of the central dermomyotome  | BAT positive  | [114,144]  |
|   | <i>myf5-Cre;R26R-eYFP</i>                            | Knock-in  | Described above   | BAT positive  | [116,117]  |
|   | <i>pax7-CreER<sup>T2</sup>;R26R-LacZ</i>             | Knock-in  | Pax7 is expressed in medial and central cells of the dermomyotome and in adult satellite cells  | BAT positive  | [121,122]  |
|   | <i>myf5-Cre;R26R-LacZ</i>                            | Knock-in  | Described above   | BAT positive; >99% of adipogenic BAT SVF cells positive   | [83,117]   |
|   | <i>myf5-Cre;R26R-eYFP</i>                            | Knock-in  | Described above   | >95% of BAT adipocyte precursors positive   | [83,117]   |
| <i>VE-cadherin-Cre;R26R-LacZ</i>                | Knock-in   | Described above   | BAT positive  | [96,98]   |            |
| <i>VE-cadherin-CreER<sup>T2</sup>;R26R-LacZ</i> | Knock-in   | Described above   | Many brown adipocytes positive  | [97,98]   |            |
| Brite adipocytes                                | <i>myf5-Cre;R26R-eYFP</i>                            | Knock-in  | Described above   | Brite adipocytes in ingWAT negative   | [116,117]  |
|   | <i>myf5-Cre;R26R-LacZ</i>                            | Knock-in  | Described above   | CL316,243 induced brite adipocytes in asWAT and rWAT positive; brite adipocytes induced in ingWAT negative  | [83,117]   |
|   | <i>VE-cadherin-CreER<sup>T2</sup>;R26R-LacZ</i>      | Knock-in  | Described above   | Rosiglitazone induced bright adipocytes in ingWAT positive  | [97,98]    |
|   | <i>UCP1-CreER<sup>T2</sup>;R26R-tdRFP</i>            | Knock-in  | UCP1 is the most accepted functional marker of brown adipocytes   | Brite and white adipocytes interconvert in ingWAT depending on the temperature  | [51]       |

(continued on next page)

**Table 1** (continued)

|   | Genetic approach                     | Promoter location | Rationale       | Reported result | References |
|---|--------------------------------------|-------------------|-----------------|-----------------|------------|
| Adipocyte precursors in skeletal muscle | <i>myf5-Cre;R26R-eYFP</i>            | Knock-in          | Described above | Negative        | [84,117]   |
|   | <i>myf5-Cre;R26R-eYFP</i>            | Knock-in          | Described above | Negative        | [37,117]   |
|   | <i>myf5-Cre;</i><br><i>R26R-mTmG</i> | Knock-in          | Described above | Negative        | [117,124]  |
|   | <i>Pax3-Cre;</i><br><i>R26R-mTmG</i> | Knock-in          | Described above | Negative        | [123,124]  |
|   |                                      |                   |                 |                 |            |

R26R: Rosa 26 reporter.

tTA: tetracycline transactivator.

TRE: tetracycline response element.

SVF: stromal vascular fraction.

adipocyte begins. Moreover, strategies targeting nascent adipocytes could be useful in fighting obesity; however the identity of the adipocyte progenitor cells is just beginning to be unraveled. As obesity progresses, adipose tissues grow by hypertrophy (increasing the size of their adipocytes) and by hyperplasia (increasing the number of adipocytes) [75–79]. A yearly adipocyte turnover rate for lean and obese humans of 10% has been reported [79], but because mature adipocytes do not divide, a resident adipocyte precursor cell must exist.

Undifferentiated adipocyte progenitor cells were long assumed to reside in the stromal vascular fraction, but only recently have cell surface markers been used to distinguish adipocyte precursor cells from non-adipogenic stromal cells by fluorescence activated cell sorting (FACS) [37,80–82]. One study identifies a CD45<sup>−</sup>CD31<sup>−</sup>Ter119<sup>−</sup>(Lin<sup>−</sup>)CD29<sup>+</sup>Sca1<sup>+</sup>CD34<sup>+</sup>CD24<sup>+</sup> population that contains adipocyte progenitors capable of reconstituting a functional white adipose tissue when transplanted into lipodystrophic mice [80]. Similar cell surface markers have been used to isolate adipocyte precursors from brown fat and skeletal muscle [37,83,84]. Whether prospectively isolated adipocyte precursors originating from different fat depots are functionally similar is an open question. Moreover, whether a true adipocyte stem cell exists that can serially reconstitute an entire fat tissue including the same complement of stem and committed progenitor cells is unclear. Notably, depot-specific differences have been reported in adipocyte precursor cell surface marker expression levels, gene expression profiles, proliferation and differentiation capacity, and response to high fat diet [81,85–87]. Strain-dependent differences in adipocyte precursor cell number have also been observed [85]. It has been argued that using cell surface markers to isolate adipose precursors is problematic because these markers lack functional relevance and cannot be used to localize adipocytes in vivo [88]. While valid, alternative approaches rely on identifying adipocyte precursor cell functional marker genes and using the promoter to drive expression of a reporter to genetically mark lineages (to date, a truly specific adipocyte precursor cell functional marker remains elusive) [21,89,90]. Both approaches are in fact quite complimentary; however, in clinical practice the most practical method to isolate adipocyte precursor cells from human lipospirates is by cell sorting based on surface marker expression [73]. Protocols to prospectively isolate human adipocyte precursor cells are currently being optimized.

Lineage tracing on the other hand is the only technique that can conclusively indicate the developmental origin of an adipocyte or reveal which tissue-resident progenitors actually become mature adipocytes, and therefore these studies can only be performed in mice. Many groups are performing lineage-tracing analyses using transgenic and Cre-Lox technology [Table 1]. Hypothesizing that peroxisome proliferator-activated receptor  $\gamma$  (PPAR $\gamma$ ) expression is an early marker of adipocyte commitment, one study marked PPAR $\gamma$ -expressing stromal vascular fraction cells with GFP (using a PPAR $\gamma$ -tetracycline transactivator (tTA) knock-in mouse together with the tetracycline response element (TRE)-H2B-GFP reporter system) [82]. This labeled an adipocyte precursor population that expresses cell surface markers that are similar to the prospectively isolated adipocyte precursors described above suggesting that early precursors express PPAR $\gamma$  [80]. Interestingly, these GFP marked

precursors appear to commit to the adipocyte lineage in embryonic development or shortly after birth, and localize in vivo to the adipose tissue mural cell compartment of the vasculature [82]. Using platelet-derived growth factor receptor  $\beta$  (PDGFR $\beta$ ) expression (which marks mural cells in many tissues) to isolate and fractionate stromal vascular fraction cells confirmed that the PDGFR $\beta$ <sup>+</sup> mural cells in adipose tissue, but not in other tissues like the kidneys, skeletal muscle, and cardiac muscle, are enriched for adipogenic precursors. In fact, when PDGFR $\beta$ -Cre is combined with the Rosa 26 reporter (R26R)-LacZ reporter (which permanently labels PDGFR $\beta$ -Cre expressing cells and their descendants) rWAT and ingWAT are homogeneously marked [82]. The PDGF receptors (PDGFR $\alpha$  and  $\beta$ ) are platelet-derived growth factor sensing receptor tyrosine kinases that activate several down-stream signaling pathways including the MAPK, PI3K and PLC $\gamma$  pathways [91]. In contrast, PPAR $\gamma$ -GFP lineage tracing only labels a subset of mural cells, indicating that while all the adipocytes appear to arise from mural cells, only a subset of the mural cells are adipogenic [82]. What determines the identity of the adipogenic population is not known. These observations are consistent with previous ultra-structural findings suggesting the adipose vasculature may function as the adipocyte progenitor cell niche [92–94].

The theory that adipocytes arise from a vascular bed has recently been expanded to include the hypothesis that adipocytes actually originate from a subset of endothelial cells. For example, transgenic mice driving GFP expression from the promoter of the zinc finger protein 423 (*zfp423*), a transcription factor involved early in preadipocyte commitment, not only labels mural cells, but also some endothelial cells in ingWAT and iBAT as determined by co-expression with the endothelial cell marker CD31 and PDGFR $\beta$  [88,95]. Interestingly, the *zfp423*-GFP marked cells are different from those marked with PPAR $\gamma$ -GFP, and exhibit depot-specific differences in the expression of surface markers [80,82,95]. Thus, *zfp423* could be a functional marker of white and brown preadipocytes early in commitment. However, the specificity of *zfp423* to preadipocytes is unclear because it is also expressed in skeletal muscle and brain tissues at comparable levels [95]. A related study also arrives at an endothelial-cell-of-origin model using the endothelial cell specific VE-cadherin-Cre knock-in allele with the R26R-LacZ reporter, which reportedly marks adipocytes in pgWAT, ingWAT and iBAT [96–98]. However, another study found that conditionally deleting PPAR $\gamma$  with the endothelial specific receptor tyrosine kinase-Cre (Tie2-Cre) did not affect PPAR $\gamma$  expression in fat or adipose tissue mass [99]. Moreover, when the same constitutive VE-cadherin-Cre driver or Tie2-Cre is used in combination with the R26R-mTmG reporter (in which Cre turns off expression of membrane-targeted Tomato fluorescent protein and induces expression of membrane-targeted GFP) no adipocytes expressed GFP indicating that VE-cadherin or Tie2 was not expressed in their lineage [100]. These contradictory findings might suggest that only subsets of adipocytes are of endothelial cell origin. This needs further investigation.

Accurate lineage tracing can be challenging in adipocytes and this warrants a brief discussion of certain considerations. For starters, in any lineage tracing study drivers expressed from an endogenous locus/promoter are preferred because they more likely reflect the

physiological expression pattern. One factor to consider specifically when tracing mature adipocytes is their paucity of cytoplasm. This poses a particular problem when analyzing tissue sections for the expression of cytoplasmic reporters, such as LacZ or YFP/GFP. The mTmG reporter overcomes this caveat because the fluorescent reporters are membrane targeted and therefore concentrate the reporter at the cell membrane surrounding the predominantly lipid-filled interior of the adipocyte. It is also wise to employ careful co-staining with known markers to distinguish between signals emanating from the miniscule membrane/cytoplasmic region of the adipocyte from signals originating in the abutting stromal vascular fraction cells, especially in highly vascularized depots.

While progress is being made in identifying the white adipocyte progenitors residing in mature adipose tissues, much less is understood about the embryonic origins of these progenitors. Previously all adipocytes were thought to arise from a common mesodermal precursor [17], but in recent years lineage tracing studies appear to have challenged this assumption and the emerging view—perhaps not surprisingly—is that white adipocyte ancestry is complex. An early clue suggesting white adipocytes might have different ancestries comes from the pathology of certain partial lipodystrophies. For example patients with familial partial lipodystrophy type 2 (FPLD2; OMIM #151660 — also called Dunnigan syndrome) suffer from the gradual atrophy of adipose tissue in the limbs and parts of the trunk while adipose tissue in the face, neck, and intra-abdominal areas are spared and often increase in mass. Similarly, in congenital infiltrating lipomatosis of the face (CIL-F) excessive adipose tissue only infiltrates the facial soft tissue forming a lipomatous tumor that causes severe disfigurement [101,102]. Such regional characteristics suggest a distinct adipogenic route in the cephalic region. Consistent with this hypothesis, lineage-tracing studies in mice indicate that Sox10-Cre, which expresses in pre-migratory and migratory cells of neural crest, mark adipocytes in the head but not in the ingWAT and pgWAT [103]. A related lineage tracing study using the Wnt1-Cre, which is induced during neural crest formation, finds that craniofacial progenitors and adipocytes but not subcutaneous or visceral adipocytes trace to a neural crest cell of origin [104]. This study also finds that the neural crest adipogenic lineage is the predominant contributor to a pool of adipogenic progenitors residing in both the cephalic fat and muscle. Curiously, the number of neural crest-derived progenitors declines with age and are replaced with an unidentified lineage in what has been called a lineage-succession model [104]. Interestingly, these two distinct adipogenic lineages have similar metabolic properties indicating that their divergent origin does not impart any unique functional feature that distinguishes them. Bones and skeletal muscles in the head are known to originate from the neural crest [105]. Thus, one possibility is that some adipocytes could share a common origin with their neighboring organs and this might reflect an important developmental relationship.

It has been reported that some adipogenic cells originate in the bone marrow, and that after bone marrow transplants in rodents, hematopoietic cells become adipocytes in iBAT, asWAT, omental WAT, and mWAT [106–108]. Moreover, it has also been reported that a substantial number of adipocytes in the pgWAT are labeled with lysozyme 2-Cre in *LysM-Cre;R26R-LacZ* mice in which Cre expression marks the myeloid lineage [109–111]. However, this too is up for debate because Vav1 oncogene-Cre (*Vav1-Cre*), which marks hematopoietic lineages [112] fails to label any adipocytes or adipocyte progenitors [100].

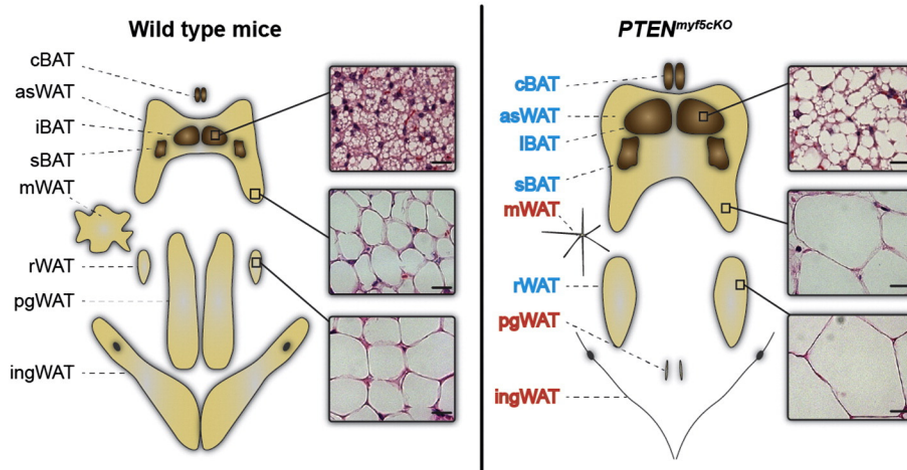
#### 4. The Myf5 lineage contribution to fat

While the vision of white adipocyte origins remains blurred, what seemed to be increasing in clarity was that brown adipocytes arise from a distinct mesenchymal precursor cell shared with skeletal muscle, but not with any white adipocyte. Evidence suggesting that BAT and WAT could have distinct developmental origins first came

from a lineage tracing analysis of central dermomyotome cells expressing the transcription factor *Engrailed1* (a pattern formation related Homeobox-family member) with an *En1-CreER* knock-in allele [113]. Using the R26R-LacZ reporter this study revealed that iBAT but not ingWAT or pgWAT shares a common origin with skeletal muscle and dermis [114] [Radhika Atit, personal communication]. Using microarrays to compare the gene expression profiles of preadipocytes during *in vitro* differentiation, a subsequent study determined that brown preadipocytes isolated from a mix of iBAT, sBAT and cBAT but not white preadipocytes from pgWAT have a myogenic-like transcriptional signature (including expression of myogenic factor 5 (*Myf5*) and myogenic differentiation antigen (*MyoD*)) [115]. This model seemed all-but-confirmed by a report that traced the *Myf5* lineage in fat [116]. In this study, the *Myf5-Cre* knock-in allele [117] was used to drive YFP expression from the R26R-YFP reporter. Using an anti-GFP antibody to detect YFP by immunofluorescence, it was shown that regionally localized iBAT and skeletal muscle, but not interscapular WAT expresses YFP [116]. YFP was also not detected in the brite adipocytes induced to form within ingWAT. This *Myf5-Cre* labeling pattern was confirmed by examining YFP mRNA expression. Other related work found that when *Myf5-Cre* drives expression of the R26R-diphtheria toxin A-chain construct, the *Myf5*-lineage is ablated and this greatly reduces the classical BAT depots [118,119] [Thomas Braun, personal communication]. Finally, lineage tracing performed with Pax7 (paired box7)-Cre or Pax3 (paired box3)-Cre knock-in alleles (Pax7 and Pax3 are transcription factors that cooperate with *Myf5* to drive myogenesis) also label BAT [120–124]. The notion that BAT and skeletal muscle share a common origin distinct from WAT is logical because of the metabolic similarities between BAT and muscle (such as high mitochondrial content and increased energy expenditure) and thus from these studies emerged a model arguing that brown adipocytes can be delineated from white (and brite) adipocytes by their ancestral expression of *Myf5* [17,89].

The simplicity and logic behind the *Myf5* model led to its widespread acceptance. Therefore, the recent finding that deleting *phosphatase and tensin homolog (PTEN)* with *Myf5-Cre* affects brown and white adipose tissue was quite surprising [83]. *PTEN* is a tumor suppressor and negative regulator of the insulin-PI3K-AKT-mTOR growth pathway [125]. In a genetic study aimed at understanding how this pathway regulates the growth of mesodermal derived tissues, the same *Myf5-Cre* knock-in allele used previously to trace the origin of brown adipocytes was used to conditionally delete *PTEN*. The *myf5-cre;PTEN<sup>fl/fl</sup>* conditional knockout (*PTEN<sup>myf5-CKO</sup>*) mice have severely overgrown interscapular, subscapular, and cervical BATs that are *PTEN*-deficient, consistent with brown adipocytes originating from a *Myf5*-positive (*Myf5<sup>+</sup>*) lineage. However, in contradiction to the model that all white adipocytes originate from a *Myf5*-negative (*Myf5<sup>neg</sup>*) precursor, *PTEN<sup>myf5-CKO</sup>* mice also have hypertrophic anterior subcutaneous WATs and retroperitoneal WAT that are also *PTEN* deficient [83]. Even more surprising is that pgWAT and mesenteric WAT as well as posterior subcutaneous WATs including ingWAT are completely absent [83] (Fig. 2). This total body fat redistribution phenotype led to a comprehensive reevaluation of the *Myf5<sup>+</sup>* lineage contribution to fat.

Originally, the *Myf5* lineage had only been examined in a few adipose tissue depots. To revisit the *Myf5<sup>+</sup>* lineage contribution to adipose tissue, the *Myf5-Cre* knock-in allele [117] was again combined with R26R-YFP reporter and separately with the R26R-LacZ reporter, and lineage tracing was performed on a broad selection of depots [83]. Here, *Myf5-Cre* was found to drive YFP mRNA expression not only in classical brown fat depots (iBAT, sBAT and cBAT) as expected, but also in anterior subcutaneous white fats (interscapular and axillary) and in rWAT. In contrast, very little YFP mRNA expression was detected in white adipocytes located in the ingWAT or pgWAT. These results were confirmed using *Myf5-Cre;R26R-LacZ* mice indicating that the *Myf5*-lineage contribution to fat is broader than previously thought. As mentioned above, *En1-CreER* had previously been shown to label dorsal dermis in addition to skeletal muscle



**Fig. 2.** Deleting *PTEN* with *Myf5*-Cre models partial lipodystrophy. Deleting *PTEN* with *Myf5*-Cre causes lipohypertrophy of the fat depots largely derived from *Myf5*<sup>+</sup> adipocyte precursors (labeled in blue) and lipotrophy of the fats that are largely *Myf5*<sup>neg</sup> (labeled in red). The overgrown fats are bigger because they contain more cells due to expansion of the *Myf5*<sup>+</sup> adipocyte precursor cell population (not shown) and bigger cells as a result of increased lipid content (see histology insets). The absent fats are reduced to fibrotic tissue. This body fat distribution phenotype resembles the pathology of a rare human lipodystrophy syndrome called Madelung's disease.

and BAT; in fact, *Myf5*-cre also labels dorsal dermis [126] (J Sanchez-Gurmaches & DA Guertin) and reportedly (using a different *Myf5*-Cre knock-in allele) a portion of the ribs and cervical vertebrae [127]. Thus, it is perhaps not surprising that *Myf5*-Cre also labels white adipocytes in the same general anatomical location.

Interestingly, when stromal vascular fraction cells are prepared from various WAT depots isolated from *Myf5*-Cre;R26R-LacZ mice and differentiated *in vitro*, about half of the adipogenic cells from the asWATs or rWAT trace with *Myf5*-Cre compared to less-than 1% in ingWAT and pgWAT. This mixed lineage contribution is also detectable in the adipocyte precursor pool. Using *Myf5*-Cre;R26R-YFP and a FACS-based cell sorting strategy based on [80], it was found that most of the CD31<sup>-</sup>CD45<sup>-</sup>Ter119<sup>-</sup>CD29<sup>+</sup>Sca1<sup>+</sup>CD34<sup>+</sup> cells isolated from BAT, nearly half of the adipocyte precursors isolated from the asWATs, and three-quarters of the adipocyte precursors isolated from rWAT, trace with *Myf5*-Cre. In contrast, less than 10% of the adipocyte precursors isolated from ingWAT and pgWAT are marked (Fig. 1). Gene expression analysis indicates that the *Myf5*-Cre tracing white adipocyte precursors do not express brown preadipocyte markers [83]. Consistent with these data, a recent study finds that the Pax3-Cre knock-in allele also labels adipogenic precursors isolated from the stromal vascular fraction of asWAT [124]. Interestingly, the Pax3-Cre driver also labels adipocyte precursor cells with a pattern similar to the *Myf5*-Cre driver (J Sanchez-Gurmaches and DA Guertin, unpublished). Why previous work failed to uncover the *Myf5*-lineage contribution to white fat could reflect the limited survey of fat depots, technical limitations, or strain differences.

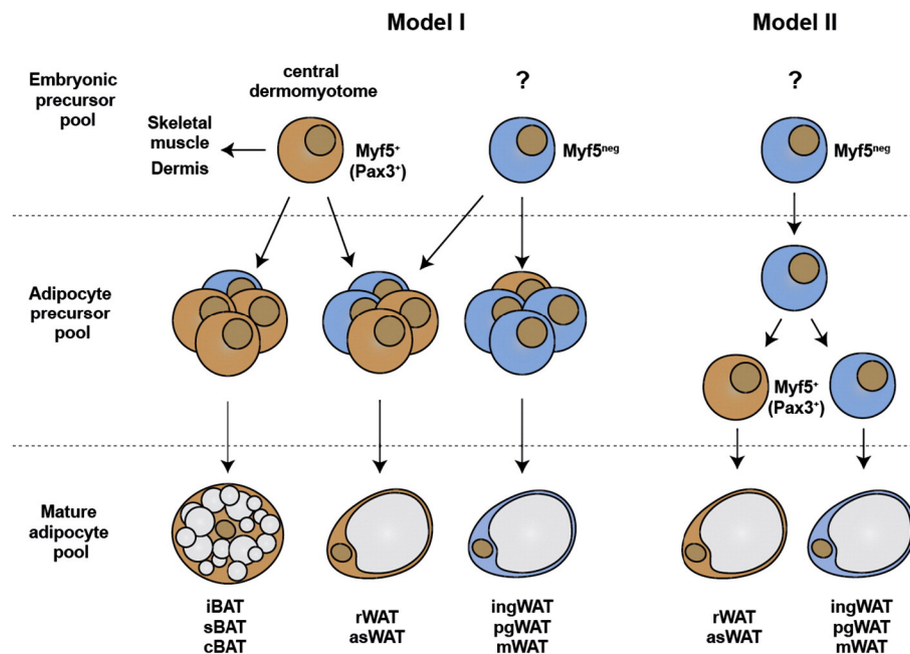
## 5. A revised *Myf5* lineage model and its implications

These new insights suggest a revised model of the *Myf5*-Cre lineage contributions to adipose tissue. We propose two possibilities: In Model (I) the *Myf5*<sup>+</sup> (Pax3<sup>+</sup>) adipocyte precursors could originate in the central dermomyotome and mix with *Myf5*<sup>neg</sup> (Pax3<sup>neg</sup>) lineage(s) in the adipocyte precursor cell compartment of each depot (Fig. 3). In BAT, most of the adipocyte precursors are *Myf5*<sup>+</sup>, in asWATs and rWAT the mix is more evenly distributed, while in other WATs such as ingWAT and pgWAT, a *Myf5*<sup>neg</sup> lineage(s) predominates (Fig. 1). Based on this model, one might predict that the mature adipocyte pool in each depot would reflect this heterogeneity. Interestingly however, *Myf5*-Cre lineage tracing of mature adipocytes with the R26R-LacZ reporter reveals a large fraction of the mature adipocytes in BAT, asWAT, and rWAT are LacZ positive, while only a few (<1%) mature adipocytes in ingWAT and pgWAT are positive [83]. Thus, despite the heterogeneity in the

adipocyte precursor cell compartment, it appears that the *Myf5*<sup>+</sup> lineage may selectively differentiate in the BAT, asWATs, and rWAT, while *Myf5*<sup>neg</sup> lineages selectively give rise to most adipocytes in the ingWAT and pgWAT. Importantly, exactly what cell types are contained within the adipocyte precursor pools as identified by cell surface markers, and how selective expansion of certain lineages might be determined is not known. Notably, the true adipocyte progenitor is likely contained within a subset of the adipocyte progenitor cell pool and it cannot yet be ruled out that *Myf5*<sup>neg</sup> precursors give rise to *Myf5*<sup>+</sup> (Pax3<sup>+</sup>) precursors within that pool. Accordingly in Model (II), *Myf5*-Cre (and Pax3-Cre) expression could be induced during determination or adipocyte differentiation in some depots (Fig. 3). Thus, it will be important to determine exactly when *Myf5*-Cre and Pax3-Cre express in the adipocyte lineages (e.g. with an inducible Cre), how many mature adipocytes in each depot arise from these lineages, and whether *Myf5*/*Pax3* have functional roles in these adipocytes [128]. This requires further investigation.

Previously, it made sense that the mitochondria rich and energy expending brown fat might share a common precursor with skeletal muscle, but what is the significance of arising from a *Myf5*<sup>+</sup> (Pax3<sup>+</sup>) versus a *Myf5*<sup>neg</sup> (Pax3<sup>neg</sup>) precursor for a white adipocyte? One possibility is that progenitors and/or mature adipocytes arising from *Myf5*<sup>+</sup> precursors could have a metabolic advantage (or disadvantage). Metabolic variations between distinct lineages could affect depot-specific fat growth and thus body fat patterning. Another possibility is that endocrine function could vary between lineages. Alternatively, there may be no functional differences between the adipocyte precursor cells arising from *Myf5* positive or negative precursors. Precedence for such functional convergence exists. For example, it was recently shown that dermal precursors in the trunk arise from *Myf5*<sup>+</sup> precursors while dermal precursors in the face arise from *Myf5*<sup>neg</sup> (*Wnt1*<sup>+</sup>) precursors, yet despite having different developmental origins, the mature dermal cells appear functionally similar based on marker profile, proliferation capacity, and ability to self-renew and differentiate [126]. Similarly, adipogenic precursors that arise from the neural crest appear functionally similar to mesoderm-derived adipogenic precursors [103,104]. Thus, being of a different developmental origin does not automatically imply a functional difference.

While it remains to be seen whether natural variations in metabolism between white adipocytes of different origins contributes to body fat patterning, is it possible that a mutation could arise that targets a specific adipocyte lineage. Such a mutation could alter the growth, metabolism, or insulin sensitivity of that lineage, and it seems reasonable to speculate that such a scenario could be the basis of certain lipodystrophy disorders. Lipodystrophy can be inherited or more



**Fig. 3.** Where is Myf5-Cre expressed during adipocyte development? At least two models could explain the origins of adipocytes that trace to Myf5-Cre<sup>+</sup> precursors. (I) In the “common origin model”, Myf5<sup>+</sup> (and possibly Pax3<sup>+</sup>) adipocyte precursors originate in the central dermomyotome during embryonic development and give rise to adipocyte progenitor cells residing in the stromal vascular fraction of each depot. Both Myf5<sup>+</sup> and Myf5<sup>neg</sup> lineages contribute to the adipocyte precursor cell pool (Lin<sup>−</sup>CD29<sup>+</sup>CD34<sup>+</sup>Sca1<sup>+</sup>) to varying degrees in each depot; however in BATs, asWATs, and rWAT the Myf5<sup>+</sup> lineage precursors appear to be preferentially selected to become adipocytes while in the ingWAT, pgWAT, and mWAT the Myf5<sup>neg</sup> lineage precursors selectively differentiate. (II) Alternatively, in the “selective induction model” Myf5<sup>neg</sup> precursors residing in the stromal vascular fraction of asWAT and rWAT (and perhaps even in BAT) induce Myf5-Cre (and possibly Pax3) expression upon differentiation giving rise to Myf5<sup>+</sup> precursors, while in the other WATs Myf5-Cre is not induced. Whether Myf5 protein has a functional role in these adipocyte lineages is unknown.

commonly acquired, and can affect all body fat (i.e. generalized lipodystrophy) or only specific regions (i.e. partial lipodystrophy). In many cases of partial lipodystrophy the spared regions increase in mass for example in Dunnigan-type familial partial lipodystrophy (described above) and Barraquer–Simons partial lipodystrophy (in which lipodystrophy begins in the head and neck and spreads to the chest, abdomen and upper extremities, while the adipose tissues in the lower abdomen, gluteal region, and lower extremities expand) [129]. Even in congenital generalized lipodystrophy or Berardinelli–Seip syndrome, which is characterized by the near-complete absence of adipose tissue, some fat depots are spared including those in the head, palms, and sole of the foot [2]. These regional variations in fat pathology have contributed to the hypothesis that adipocytes have distinct developmental origins.

Consistent with the idea that lineage-restricted mutations could cause lipodystrophy is the discovery that *PTEN* loss in the Myf5 lineage causes severe body fat redistribution [83]. In *PTEN*<sup>Myf5<sup>CKO</sup> mice, body fat is redistributed such that Myf5<sup>+</sup> adipocytes expand (causing lipohypertrophy of BAT, asWATs, and rWAT) at the expense of Myf5<sup>neg</sup> adipocytes (i.e. inguinal, perigonadal, and mesenteric WAT), which are reduced to fibrotic tissue (Fig. 2). The hypertrophic fats have more cells due to expansion of the adipogenic precursors, and bigger cells caused by increased lipid content. *PTEN*<sup>Myf5<sup>CKO</sup> mice also exhibit skeletal muscle atrophy, which appears to be caused by a systemic effect because *PTEN* expression is maintained at normal levels in most skeletal muscles by a post-transcriptional mechanism. Losing *PTEN* hyperactivates the major intracellular effectors of insulin including PI3K, AKT, and mTOR, which likely alters the metabolic balance in the Myf5 adipocyte lineage to favor energy uptake and storage. Interestingly, the *PTEN*<sup>Myf5<sup>CKO</sup> mouse phenotypically resembles a rare partial lipodystrophy syndrome called multiple symmetric lipomatosis or Madelung’s disease [3]. Patients with Madelung’s disease suffer from mild to severe adipose tissue expansion in the head and neck with extension down the back (type I) and/or the shoulders, upper arms and abdomen (type II). The *PTEN*<sup>Myf5<sup>CKO</sup> phenotype also resembles the</sup></sup></sup></sup>

buffalo hump syndrome often seen in HIV patients on aggressive anti-retroviral therapy, which is currently the most common lipodystrophy disorder in the United States [130]. Thus, the *PTEN*<sup>Myf5<sup>CKO</sup> mouse is a novel model of partial lipodystrophy and its phenotype supports the idea that mutations targeting metabolism or the metabolic regulatory circuits in a particular adipocyte lineage could be the pathological basis of certain lipodystrophies.</sup>

## 6. Where do brite adipocytes come from?

In response to cold or  $\beta$ -adrenergic agonists, brite adipocytes appear within white fat and can comprise a significant fraction of the total adipocyte population depending upon the depot. While ingWAT and to a lesser extent rWAT are predisposed to browning based on their rapid response to acute cold challenge, other depots like pgWAT are more resistant and only show some browning after chronic cold challenge [4,21,41,52]. Two models have been proposed to explain the origins of brite adipocytes. One posits that brite adipocytes arise by transdifferentiation of existing mature white adipocytes; the second argues that brite adipocytes arise de novo from a distinct precursor population [41,48,50,131]. These models may not be mutually exclusive and depot-specific differences may exist (discussed below).

The brite adipocytes that arise in the largely Myf5<sup>neg</sup> lineage fats (e.g. pgWAT and ingWAT) are reportedly Myf5<sup>neg</sup> while the brite adipocytes arising in Myf5<sup>+</sup> lineage fats (e.g. asWATs and rWAT) are Myf5<sup>+</sup> [83,116]. This is an important point because it is argued that all brite adipocytes are Myf5<sup>neg</sup>, which is true in ingWAT but not in rWAT or asWATs. Thus, brite adipocytes from different depots likely have different origins. Factors like the proximity to noradrenergic parenchymal nerve fibers, to the vasculature, or exposure to other micro-environmental conditions could determine which cells become brite [19,40,41]. Recently, several preadipocyte cell lines were sub-cloned from immortalized stromal cultures derived from ingWAT, and only some of these lines responded like a brite preadipocyte (called beige in this report) would be predicted too following stimulation (i.e. induce

UCP1) [48]. These could be brite preadipocytes and although they were found to express distinct surface markers, lineage tracing was not performed, thus it is unclear if they represent a unique brite-adipocyte lineage. Brite/beige adipocytes from other white fat tissues were not examined [48].

Interestingly, another study finds that when mice are treated with a low dose of the  $\beta_3$ -agonist CL316,243 that promotes proliferation but avoids inflammation, the brite adipocytes appearing in the pgWAT arise from a resident progenitor cell while the brite adipocytes induced in the ingWAT appear to arise by transdifferentiation [41,50]. This suggests that the mechanism by which brite adipocytes arise varies with the depot and the duration of stimulation. Perhaps immediate responding brite adipocytes exist in ingWAT but not in pgWAT, which would explain why brite adipocytes only appear after chronic cold exposure in pgWAT, but are often readily detectable in ingWAT even at ambient temperature. The emerging nascent brite cell population in pgWAT traces to a platelet-derived growth factor receptor  $\alpha$  (PDGFR $\alpha$ ) positive precursor residing in the stromal vascular fraction (labeled with an inducible PDGFR $\alpha$ -CreER<sup>T2</sup> allele). This precursor population lacks expression of the cell surface markers CD24, PDGFR $\beta$ , the endothelial marker IB4, and is negative for smooth muscle actin and PPAR $\gamma$ , indicating it may be distinct from other precursors described above [50,80,82]. Originally, it was reported that the fate of these PDGFR $\alpha$  positive precursors is not restricted to brite fat because high-fat diet feeding instructs them to become white adipocytes [50]. A more recent lineage tracing study using a constitutive PDGFR $\alpha$ -Cre driver together with the R26R-mTmG reporter argues that all mature adipocytes in all WAT depots are in fact derivatives of PDGFR $\alpha$ <sup>+</sup> precursors [100]. Mature adipocytes do not express PDGFR $\alpha$  [50] indicating these adipocytes are branded with PDGFR $\alpha$ -Cre as precursors. Whether PDGFR $\alpha$  has a functional role in the adipocyte precursor is unclear. However, PDGFR stimulation with ligand inhibits adipogenesis in  $\beta_3$ T<sub>3</sub>-L1 cells [132] and expressing an activated mutant allele of PDGFR $\beta$  in pericytes or mesenchymal progenitor cells blocks white adipocyte differentiation [133], suggesting down-regulating PDGFR signaling might be essential for adipocyte differentiation. More information regarding PDGFR signaling can be found in [91].

A strong argument in favor of brite-to-white interconversion in the ingWAT was recently reported [51]. In this study, researchers used lineage tracing to mark the cold-induced brite adipocytes in the ingWAT (with a UCP1-CreER driver) then followed their fate upon returning the mice to warmer temperature. In response to warming (or “whitening”) the brite adipocyte characteristics disappeared but importantly the adipocytes that were labeled during cold-induced “brightening” retained their lineage mark and now expressed “white” characteristics. This is in contrast to classical brown adipocytes, which retain their brown characteristics even when unstimulated, and argues that brite adipocytes are not dedifferentiated or eliminated when no longer needed. Moreover, these same white adipocytes can be reconverted to brite adipocytes upon subsequent cold challenge. Thus, in the ingWAT interconversion between the white and brite state appears to be responsible for much of the cold-induced changes in adipocyte morphology. Future studies are needed to determine whether depot, age, or strain specific differences exist in the interconversion rate and whether any lineages have inherent potential to become brite when stimulated.

## 7. Ectopic adipocytes in skeletal muscle

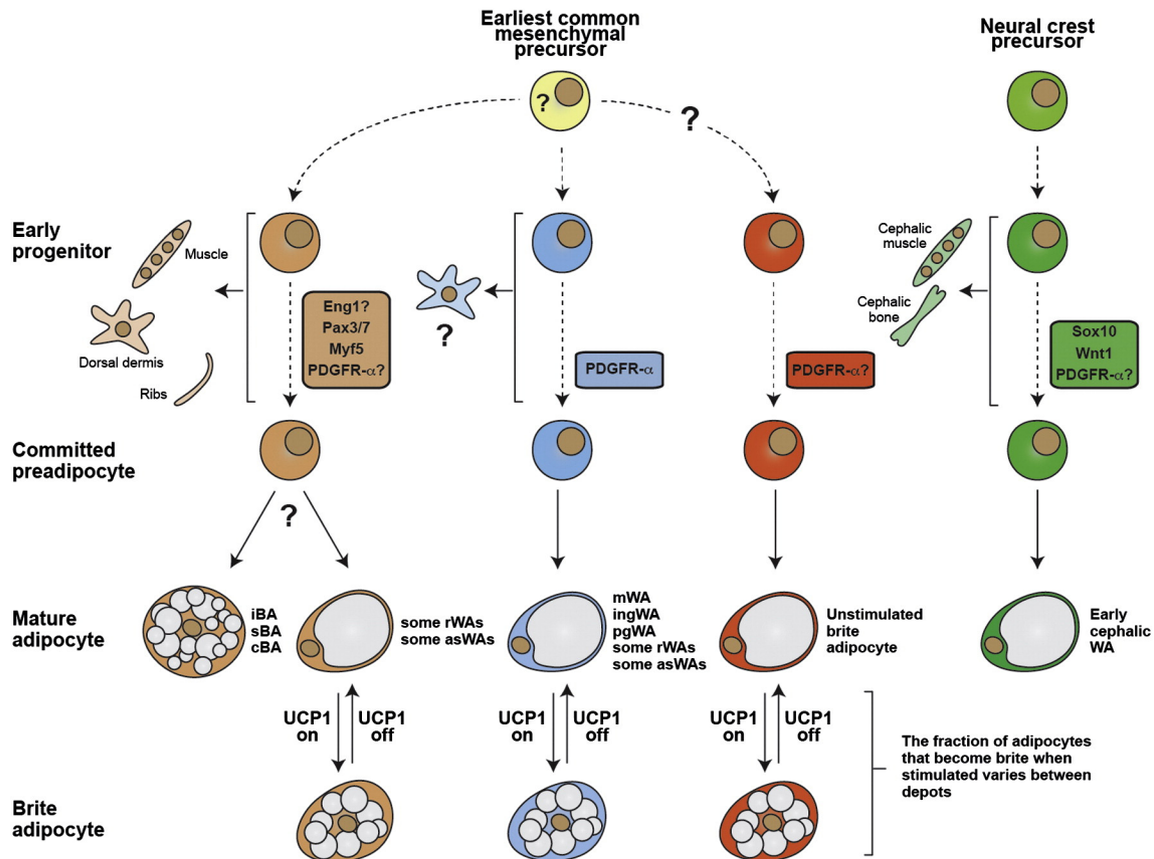
Ectopic fat accumulation in muscle is a pathological feature of several conditions including myopathies like Duchenne muscular dystrophy (OMIM #310200), obesity, and type 2-diabetes [134,135]. Studies using FACS technology to prospectively isolate adipogenic precursor cells from rodent limb muscles are also beginning to expose the identity of muscle resident adipogenic precursors [37,84,136]. While the cell surface marker profiles differ slightly between the different protocols (e.g. CD45<sup>-</sup>CD31<sup>-</sup>CD34<sup>+</sup>Sca1<sup>-</sup> $\alpha$ 7-integrin<sup>-</sup> in [84]

and CD31<sup>-</sup>CD45<sup>-</sup>SM/C-2.6<sup>-</sup> in [136]), almost all of the adipogenic precursors identified in hind limb skeletal muscles express PDGFR $\alpha$  [84,136]. These PDGFR $\alpha$ <sup>+</sup> precursors are of mesenchymal origin and reside in the interstitial space between muscle fibers; however, they are not marked with Myf5-Cre or Pax3-Cre by lineage tracing and are distinct from satellite cells based on their localization [84,124,136]. Muscle-resident adipogenic cells can also be isolated using a CD45<sup>-</sup>Mac1<sup>-</sup>Sca1<sup>+</sup> protocol [37]. In vitro, these cells can be induced to express high levels of UCP1 when stimulated with bone morphogenetic protein 7 (BMP7) suggesting they may function like inducible brown adipocytes. In fact, a histological examination of the thigh muscle revealed that brown adipocytes, defined as UCP1-positive multilocular cells, are interspersed in the muscle of mice [137]. It is interesting that muscle-associated adipogenic precursors originate from Myf5<sup>neg</sup>/Pax3<sup>neg</sup> lineages because this suggests that the adipogenic cells in muscle may have migrated from a different developmental location. It will be important to analyze the Myf5<sup>+/</sup>Pax3<sup>+</sup> lineage contribution to the muscle adipogenic precursor populations across several muscles to determine if the Myf5<sup>+</sup> precursor contribution varies with anatomical location.

## 8. What can we learn from our ancestors?

The ability to safely store large amounts of energy in adipocytes was a critical adaptation in mammalian evolution. But as the energetic cost of obtaining calories has plummeted compared to evolutionary times, the remarkable energy storing capacity of adipose tissue is now being revealed, as are its vices. White fat depots are dispersed throughout the body, and while a typical healthy individual probably has around 80,000 cal stored in fat (equivalent to over one month of meals), the distribution of that fat between different depots is highly variable in the population. These variations are exaggerated among obese individuals. In general white adipose tissue biology is not well understood, but perhaps one of the most clinically relevant mysteries is why expansion of some depots (e.g. visceral white fat) correlates with high risk of metabolic disease, while expansion of other depots (e.g. subcutaneous white fat) is benign or protective. Although one reason is likely that visceral depots drain directly into the liver causing damage, there may be other underlying functional differences related to origin. Like white fat, the amount of brown fat also varies between individuals and its presence negatively correlates with body mass index [22,28,32,33]. The existence of brown fat cells in adult humans may open doors to novel anti-obesity therapies that could function by increasing brown fat energy expenditure; however, the study of human brown fat is in its infancy.

Could lineage-tracing studies explain some of the variability between fat depots? Previously it was thought that all adipocytes descended from a common mesodermal precursor, but recent studies have argued that this assumption is incorrect. A current trending model is that brown, white, and brite adipocytes have distinct origins; however, evidence is mounting that this view is too simplistic (Fig. 4). Although all adipocytes have a common precursor if traced far enough back, moving forward it will be important to determine when unique adipocyte lineage marks are established and whether early adipocyte ancestors convey unique properties to their descendant adipocytes that make them functionally different from adipocytes that arise from other lineages. Adipocyte lineages could vary in their developmental gene expression pattern, which could affect adipocyte metabolism, the regulatory circuits that control metabolism or endocrine function, or the ability of resident progenitor cells to respond to growth cues. How might this affect body fat patterning? One possibility is that variations in metabolism or insulin sensitivity between lineages could affect depot growth rate. Partial lipodystrophies in which body fat is redistributed may be extreme examples of this. Notably, the regional absence of fat tissue associated with many lipodystrophies may often be caused by intrinsic loss of adipocyte



**Fig. 4.** A model based on recent lineage tracing studies that explains some of the complexity of adipocyte ancestry. A trending idea is that morphologically brown (multilocular), white (unilocular), and brite (interconvertible) adipocytes develop along unique lineages. However, recent studies in mice using Cre-Lox technology to irreversibly label cellular lineages argues that it is likely not this simple. An alternative view is that morphologically white adipocytes arise from multiple distinct lineages. One of these lineages is also marked by ancestral Myf5 expression and gives rise to morphologically brown adipocytes, a subset of morphologically white adipocytes (e.g. in the asWAT and rWAT), skeletal muscle cells, dermis, and ribs. Some of the Myf5-positive lineage derived white adipocytes become brite adipocytes when stimulated (e.g. Myf5-positive lineage white adipocytes in rWAT and asWAT). Other markers such as Engrailed-1 might also express early in this lineage. Whether the Myf5 lineage derived brown and white adipocytes arise from the same precursor is still unknown. Most white adipocytes in ingWAT and pgWAT develop from a Myf5-negative lineage (or possibly multiple lineages). Some of these Myf5-negative lineage white adipocytes can also become brite when stimulated (e.g. the Myf5-negative lineage white adipocytes that undergo interconversion in the ingWAT). Some brite adipocytes might also originate from a lineage distinct from the other white adipocytes although a Cre driver unique to a brite adipocyte lineage has not been found. Although white/brite adipocyte lineages are just beginning to be defined, PDGFR $\alpha$  appears to express in the ingWAT, pgWAT, rWAT, and mesenteric WAT lineages. Importantly, when each lineage mark actually expresses, when the lineages diverge during development, and the earliest common mesenchymal progenitor cell remain unclear and this is indicated by dotted lines and question marks. Another distinct white adipocyte lineage arises from Sox10 and Wnt1-expressing neural crest precursors in the cephalic region of young mice (notably, an unidentified lineage begins replacing neural crest derived cephalic adipocytes with age). These adipocytes label positive for PDGFR- $\alpha^+$  by antibody staining and therefore likely would trace with PDGFR-Cre, although this has not been shown. In sum, the emerging lineage tracing data presents a complex picture of adipocyte origins. iBA, sBA, and cBA = interscapular, subscapular, and cervical brown adipocyte; asWA, rWA, pgWA, and ingWA = anterior subcutaneous, retroperitoneal, perigonadal, and inguinal white adipocyte. For more detailed information see [50,51,83,100,103,114,116,127].

function. However, the *PTEN<sup>myf5cKO</sup>* model suggests the opposite scenario is also possible, that is a subset of regionally overactive fat tissues can prohibit the formation of fat depots in distal regions (Fig. 2). To what extent adipocytes vary in their origin, and whether origin influences function, remains to be seen. One thing that is clear is that different fat depots are heterogeneous, and thus it seems prudent at this point to view each depot as a unique organ when experimenting.

Is there an advantage to having adipocytes that arise from different developmental origins? One possibility is that having multiple origins might ensure that fat tissue still grows if one lineage becomes compromised (for example by a spontaneous mutation). Partial lipodystrophy syndromes again are examples of this in which regional lipodystrophy is often compensated for by lipohypertrophy of other depots. A second possibility is that some adipose tissue depots may grow in coordination with adjacent lean tissues, perhaps from a common precursor, to provide a locally available fuel reserve or other support service. This suggests a fate switching mechanisms could exist in a common precursor. Such a model has been proposed for brown adipose tissue and skeletal muscle development [116]. Adipose tissues of different origins could have other specialized functions, such as

hormone production, responding to diet, or fueling exercise. In mice, adipose tissues appear at different times during embryonic and post-natal growth. Perhaps early growing fat depots arise as part of a developmental program, and these are metabolically more favorable depots. In contrast, the growth of less favorable fat tissues could arise from different precursors later perhaps in close correlation with caloric load. In mice for example, WATs that arise early such as subcutaneous WATs tend to be more metabolically favorable, while pgWAT arises later, is less vascularized, less innervated, and more static in response to cold challenge [41,138].

What is the potential impact of understanding adipocyte origins for human health? As coping with the consequences of extreme adipose tissue growth is now a major burden on the health care system, therapeutic methods to restore normal energy balance are in great demand because exercise alone is no longer enough. Determining where adipocytes originate is an essential step towards elucidating the instructional program that functionally controls adipocyte development and function, and elucidating these pathways may reveal new mechanisms to block fat growth or increase the energy expenditure. On the flip side, adipose tissue is useful for reconstructive

surgery and so knowing how to grow fat also has clinical value. Adipose tissue is a source of adult stem/progenitor cells that are obtainable by minimally invasive procedures, and there is clearly no shortage of them. Understanding the origins of adipose stem cells could be valuable in optimizing cell-based therapies. While the mouse is invaluable for lineage-tracing studies because of its genetic tractability, there are many differences between rodents and humans with respect to fat distribution. Protocols to isolate human adipocyte progenitor cells are under development and it will be interesting and important to determine which functional markers are conserved.

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