The significance of beige and brown fat in humans

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Abstract
Promotion of brown adipose tissue (BAT) activity or browning of white adipose tissue has shown great potential as anti-obesity strategy in numerous preclinical models. The discovery of active BAT in humans and the recent advances in the understanding of human BAT biology and function have significantly propelled this field of research. Pharmacological stimulation of energy expenditure to counteract obesity has always been an intriguing therapeutic concept; with the identification of the specific molecular pathways of brown fat function, this idea has now become as realistic as ever. Two distinct strategies are currently being pursued; one is the activation of bone fide BAT, the other is the induction of BAT-like cells or beige adipocytes within white fat depots, a process called browning. Recent evidence suggests that both phenomena can occur in humans. Cold-induced promotion of BAT activity is strongly associated with enhanced thermogenesis and energy expenditure in humans and has beneficial effects on fat mass and glucose metabolism. Despite these encouraging results, a number of issues deserve additional attention including the distinct characteristics of human vs rodent BAT, the heterogeneity of human BAT depots or the identification of the adipocyte precursors that can give rise to thermogenic cells in human adipose tissue. In addition, many pharmaceutical compounds are being tested for their ability to promote a thermogenic program in human adipocytes. This review summarizes the current knowledge about the various cellular and molecular aspects of human BAT as well as the relevance for energy metabolism including its therapeutic potential for obesity.

Introduction
There is now a large body of evidence – at least in rodents – that the actions of brown adipose tissue (BAT) or the browning of white adipose tissue (WAT) can protect against obesity and related complications (1, 2, 3). The molecular mechanisms that are responsible for the energy-dissipating qualities of brown fat have been studied in detail (4, 5). Uncoupling protein-1 (UCP-1) has been identified as the key factor controlling the thermogenic capacity of brown adipocytes (6). UCP-1 disrupts the electrochemical gradient across the mitochondrial membrane by allowing protons to reenter the mitochondrial matrix. Consequently, mitochondrial fatty acid oxidation is increased and chemical energy is wasted through heat production (thermogenesis) (4, 7). This process has long been believed to occur exclusively in brown adipocytes; however, in recent years, a large number of studies have reported about the emergence of UCP-1-positive cells in WAT with very similar properties as BAT cells (3, 8, 9). These cells have been termed ‘beige’ or ‘brite’ because of their intermediate function between brown and white adipocytes. A hallmark of beige adipocytes is their potential to take on a thermogenic phenotype in response...
to various stimuli such as cold, chemical compounds or genetic factors. Under basal conditions, beige cells express very similar molecular markers as classic white adipocytes (with low UCP-1 levels); however, when appropriately stimulated, they acquire a thermogenic signature with high UCP-1 expression and increased energy consumption similar to bone fide brown adipocytes (2, 3, 10, 11, 12, 13). Despite some mutual features, brown and beige adipocytes also have very distinct characteristics. For instance, they differ in their anatomical location; whereas beige cells emerge within WAT, particularly in the subcutaneous depots, classic brown adipocytes reside within bone fide BAT typically located in the interscapular region in rodents and human infants (3, 13, 14). Another difference between brown and beige adipocytes pertains to their developmental origin. Classical brown adipocytes originate from precursor cells in the embryonic mesoderm that express the transcription factor myogenic factor 5 (MYF5) and can give rise to either brown adipocytes or myocytes. The transcriptional co-regulator PR domain containing 16 (PRDM16) has been shown to control the switch between myogenic vs adipogenic differentiation in the MYF5-positive precursors (15, 16). In contrast, beige adipocytes are believed to come from a MYF5-negative lineage. However, the exact origin of beige adipocytes still remains a matter of great debate given that recent lineage tracing studies in rodents provide evidence for two fundamentally different concepts (3). One is the so-called precursor model because it implies that beige adipocytes are derived from a distinct progenitor cell population by de novo differentiation in response to distinct stimuli such as prolonged cold exposure or certain genetic modifications. The other concept proposes that mature white adipocytes can transdifferentiate into beige adipocytes when appropriately challenged and has therefore been termed transdifferentiation or more accurately interconversion model (8, 17, 18). Given the context in which these fate-mapping studies have been performed, it is possible that both developmental models are correct and that additional factors such as the environment (e.g. ambient temperature), the genetic background or fat depot specific differences may influence the decision how beige cells finally develop. Regardless of the precise developmental origin, the induction of thermogenically active beige adipocytes in WAT has proven to be effective in preventing or reducing obesity and diabetes in numerous preclinical models.

Current research is directed at discovering the various molecular cues that induce such a browning program in WAT. Besides chronic cold exposure, a number of transcription factors and co-regulators have been identified to control the browning process. Among those are peroxisome proliferator-activated receptor γ coactivator 1α (PGC-1α), PRDM16, bone morphogenetic proteins and many more (1). In addition, circulating factors such as thyroid hormones, bile acids, natriuretic peptides, retinoids, fibroblast growth factor 21 (FGF-21) and various cytokines can regulate a thermogenic program in WAT (14, 19, 20, 21, 22, 23). Notably, physical exercise has also been demonstrated to promote WAT browning in mice through a muscle-derived hormone named irisin (24). However, it is still a matter of investigation which of these pathways are conserved in humans and to what extent they contribute to thermogenesis and energy dissipation.

Detection of active human BAT

Despite previous observations in cancer patients undergoing positron emission tomography combined with X-ray computed tomography (PET/CT) showing significant uptake of radioactively labeled glucose 

$^{18}$F-fluorodeoxyglucose (FDG) in various cervical and thoracic fat depots, the metabolic relevance of this tissue has long been neglected. In fact, the FDG-positive fat depots have been viewed as impediment to diagnostic imaging because the signal could obscure tumor visualization or introduce false positive results (25, 26). Therefore, beta blockers have occasionally been used to repress the unwanted FDG uptake (27). However, the interest in these FDG-positive fat depots has risen significantly since 2009 when three prospective PET/CT studies coupled with tissue biopsies in human adults showed that these BAT depots contained UCP-1-expressing cells and could be activated by cold exposure (28, 29, 30). Notably, very early studies including one in Finnish outdoor workers had already indicated that the occurrence of BAT may be influenced by ambient temperatures (31). In line with these early observations are newer reports of seasonal variations in BAT activity showing negative correlations with average outdoor temperatures (32, 33).

In humans, BAT depots are typically located in the deep cervical, supraclavicular, parasternal and sometimes perirenal regions (28, 29, 30, 34 and Fig. 1). In some rare cases of pheochromocytoma, significant FDG uptake has also been detected in visceral adipose tissue reflecting a browning of this fat compartment due to the prolonged stimulation with endogenous catecholamines (35, 36).
However, after surgical removal of the tumor, FDG uptake was no longer present in the visceral fat suggesting reversal of the browning process (35).

Despite some limitations, FDG-PET/CT evolved as the gold standard for imaging human BAT, although a number of alternative approaches are currently being investigated. Particularly non-radioactive imaging techniques such as magnetic resonance imaging (MRI) have recently been pursued given that repeated PET/CT studies within the same individuals are limited due to the cumulative radiation exposure. Chemical-shift-encoded water-fat MRI allows to estimate the time-resolved fat fraction, which has been helpful to distinguish between WAT and BAT and also between active and inactive BAT (37, 38, 39). Other research is directed at testing alternative radiotracers beyond FDG (40, 41). Given that BAT is primarily utilizing fatty acids as fuel substrate, various lipid tracers such as $^{11}$C-acetate and $^{18}$F-fluoro-thiaheptadecanoic acid have been studied (42). However, FDG-PET/CT continues to be the most important platform for the detection and quantification of human BAT. In order to standardize the protocols for FDG-PET/CT imaging of human BAT and to ease comparisons between different studies, an expert panel assembled by the National Institute of Health (NIH) has established the first guidelines in 2014 (43). The ‘Brown Adipose Reporting Criteria in Imaging Studies’ (BARCIST) provide recommendations for experimental parameters and minimum requirements for acquisition, analysis and reporting of FDG-PET/CT BAT data in humans.

Beige or brown: does it really matter?

Ever since the discovery of cold-induced BAT in humans, researchers have tried to better understand the molecular characteristics of this adipose depot. Therefore, biopsy studies of FDG-positive neck fat depots have been performed to establish the genetic signature of human BAT. By comparing the results from human fat biopsies with those from mouse fat depots, researchers have tried to address the question whether human BAT resembles the molecular and functional aspects of bone fide brown or beige fat. Currently, there is evidence for both, which can in part be explained by the large heterogeneity of the human BAT samples studied. Depending upon the exact anatomical location, human BAT has been suggested to display either a classical brown or a beige signature or both (44, 45, 46, 47). Another important limitation besides the heterogeneous composition of human brown

Figure 1
Representative $^{18}$FDG-PET/CT scans before (A) and after (B) short-term cold exposure. FDG-positive BAT depots are visible at the typical cervical, clavicular and paravertebral regions.
Fat depots is the fact that the marker genes that have been used for genetic characterization of human BAT have previously been identified in mouse tissues (10, 45, 48). These gene sets may not fully reproduce the molecular landscape of human BAT, which makes it more challenging to unequivocally distinguish between beige and brown fat in humans. In this context, it is noteworthy that thermoneutral conditions occur at different ambient temperatures in mice and humans that is mice require slightly higher temperatures for thermoneutrality. Even though it is still a matter of debate which is the optimal housing temperature for mice to best mimic human thermal conditions, this needs to be considered when comparing molecular and functional aspects of human and murine brown/beige adipocytes (49, 50, 51). Regardless of those experimental caveats, human BAT is a metabolically highly active tissue with a respiratory capacity up to 50-fold greater than WAT and comparable to rodent BAT when normalized to the mitochondrial content (52).

Another important issue that has recently drawn a lot of interest is the capacity of human WAT to acquire BAT-like features. Turning the large amounts of white fat, as seen in obese subjects, into energy-dissipating beige fat represents an intriguing concept with great potential for therapeutic applications. Reports from pheochromocytoma patients with FDG-positive intra-abdominal WAT depots containing adipocytes with high UCP-1 expression suggest that browning of visceral WAT can occur in response to catecholamine excess (35, 36, 53, 54). More recently, a case study in a patient with papillary thyroid carcinoma found that TSH-suppressing treatment with levothyroxine (T4) induced BAT activity and browning of subcutaneous adipose tissue (56). The relevance of thyroid hormones for adipose browning in humans is also supported by findings that circulating free T4 concentrations significantly correlated with gene expression of classic brown fat markers in the subcutaneous WAT of 163 middle-aged obese men and women (57). Whether browning of WAT can also be induced under controlled pharmacological conditions and whether that results in increased energy wasting and body weight loss in humans is yet to be shown.

**Function of human BAT: relevance for energy metabolism**

Soon after the detection of cold-induced BAT in human adults in 2009, numerous follow-up studies confirmed that activated BAT contributed to non-shivering thermogenesis and oxidative metabolism in humans. Energy dissipation is significantly increased with BAT activation and was reported to account for up to 15% of resting energy expenditure in men (58). Chronic cold exposure (2 h daily for six weeks) resulted not only in enhanced BAT activity and cold-induced thermogenesis but also in a significant loss of body fat mass. The reduction in fat mass was greater in subjects with higher BAT activity and correlated with the degree of cold-induced thermogenesis (59). Both, glucose and fatty acid uptake contribute to increased BAT oxidative metabolism in humans (42, 60, 61, 62). Using hyperinsulinemic–euglycemic clamp studies, BAT activation was shown to improve whole-body glucose disposal and insulin sensitivity confirming its relevance for total energy metabolism (58, 61). However, rapidly elevated BAT radiodensity after short-term cold exposure led to the hypothesis that acute BAT thermogenesis is fueled predominantly by fatty acids hydrolyzed from intracellular triglycerides. This hypothesis has been tested in a recent study using 13C-acetate and 18F-FDG-PET/CT in combination with acute cold exposure and oral nicotinic acid, an inhibitor of intracellular triglyceride lipolysis (63). Not only did nicotinic acid administration suppress cold-induced BAT oxidative metabolism and glucose uptake in healthy men, but it also led to an increase in skeletal muscle shivering intensity. These results suggest that intracellular triglyceride lipolysis is likely essential for BAT thermogenesis and provide experimental evidence for a reciprocal role of non-shivering thermogenesis in BAT and shivering thermogenesis in skeletal muscle. In recent years, the importance of BAT as a major site for lipoprotein metabolism has been increasingly appreciated thanks largely to mouse models that demonstrated that cold-activated BAT can effectively clear plasma triglycerides predominantly through the actions of lipoprotein lipase and CD36 (64, 65, 66). Moreover, BAT activation mitigates hypercholesterolemia and atherosclerotic plaque formation (66, 67), suggesting a potential cardioprotective role, possibly independent of increased energy expenditure and weight loss. Whether this concept holds true for cardiovascular disease in humans is yet to be shown. Besides cardiovascular protection, another important question from a therapeutic standpoint is, whether active BAT can be recruited in obese and/or
Human brown adipose tissue transplantation in mice significantly improved glycemic conditions, WAT transplantation of BAT into diabetic or diet-induced obese mice from BAT transplantation studies in mice and rats. The recruitment of BAT during cold acclimation was not associated with any elevation in energy expenditure, which may be a consequence of the modest increase in BAT activity observed in this study. However, when comparing BAT-positive with BAT-negative subjects, there is a plethora of evidence for higher cold-induced thermogenesis, enhanced energy expenditure and improved whole-body glucose homeostasis in those with active BAT irrespective of the body mass (58, 59, 68, 69). A hyperinsulinemic–euglycemic clamp study in overweight men with type 2 diabetes found that 10-day cold acclimation significantly enhanced BAT activity and improved whole-body insulin sensitivity by 43%, which was mainly driven by increased insulin-stimulated glucose disposal (61). Interestingly, enhanced skeletal muscle glucose uptake due to elevated GLUT4 translocation seemed to account for majority of the glycemic effects seen in this study, raising the possibility that activated BAT may release endocrine factors that engage other metabolic tissues such as skeletal muscle. Recent evidence suggests that BAT is indeed a source of hormone production and that these BAT-derived factors may play a role in systemic energy metabolism.

**BAT as a secretory organ**

In addition to its function as a thermogenic organ, BAT has recently been identified as a tissue with high secretory capacity. Accumulating evidence from preclinical studies suggests that molecules released from BAT differ significantly from those released from WAT. The so-called BAT adipokines or BATokines can act in an autocrine, paracrine or endocrine fashion (71). Most of the autocrine and paracrine molecules such as nerve growth factor, fibroblastic growth factor 2 and vascular endothelial growth factor-A promote BAT growth, vascularization, innervation and blood flow, processes that are important for BAT recruitment during thermogenic stimulation (71). Strong evidence for BAT as an endocrine organ came from BAT transplantation studies in mice and rats. The transplantation of BAT into diabetic or diet-induced obese mice significantly improved glycemic conditions, WAT inflammation and systemic adipokine profiles (72, 73, 74). Secretion of insulin-like growth factor 1 has been proposed as one endocrine mechanism for the antidiabetic actions of transplanted BAT (72). Morphological and molecular studies from transplanted BAT revealed larger adipocyte size and decreased thermogenic gene expression in comparison to endogenous BAT (73). These findings suggested that the beneficial effects of BAT transplantation may not primarily be driven by the intrinsic thermogenic activity but by endocrine factors released from the transplant. Additional support for this concept came from a study showing that the favorable metabolic effects of BAT transplantation were lost when interleukin-6 (IL-6)-deficient mice were used as donors, suggesting IL-6 as another endocrine mediator of transplanted BAT (74). IL-6 has previously been identified to be secreted from activated brown adipocytes (75). The role of IL-6 as a classic pro-inflammatory cytokine has to be reconsidered in light of recent data that IL-6 signaling can promote insulin sensitivity in skeletal muscle and adipose tissue of mice (76, 77). The importance for BAT as secretory organ has also been highlighted by a recent report identifying BAT as a source for circulating exosomal miRNAs, which function as negative regulators of translation. Transplantation of BAT into mice lacking the miRNA-processing enzyme Dicer restored the levels of numerous circulating miRNAs that are associated with an improvement in glucose tolerance (78). Interestingly, serum concentrations of the exosomal miRNA miR-92 have been reported to correlate with BAT glucose uptake in human PET/CT studies, suggesting miRNAs as potential new biomarkers for BAT activity (79). In this context, a new class of lipids has been identified, that is released from tissues and acts locally or systemically on energy turnover, glucose metabolism and inflammatory pathways (80, 81). One of these so-called lipokines is the BAT-derived 12,13-dihydroxy-9Z-octadecenoic acid (12,13-diHOME), which is increased in response to cold exposure and promotes BAT activation in a feedforward mechanism (82). In search for new BAT biomarkers, targeted metabolomics recently showed that the lipid lysophosphatidylcholine-acyl C16:0 is strongly associated with BAT activity in humans as determined by PET/CT scans (83). These findings highlight the promising developments in the identification of novel non-invasive markers for the prediction of metabolically active BAT in humans.

Another circulating molecule that has lately received a lot of attention is fibroblast growth factor 21 (FGF21), which is mainly secreted by the liver but...
also by activated brown adipocytes (84). FGF21 acts in concert with the cofactor β-Klotho, which is required for the interaction with FGF receptors (85). FGF21 regulates energy metabolism through multiple pathways including enhanced browning of WAT and accelerated lipoprotein catabolism (23, 65). In addition, FGF21 alters food preferences through hypothalamic signals that is decreased desire for sweets and consequently reduces carbohydrate intake in mice (86, 87). As a result, FGF21 actions counteract diet-induced obesity, hyperglycemia and hyperlipidemia in various animal models. Notably, FGF21 is also expressed in human BAT and cold exposure increases circulating concentrations of FGF21 in humans, possibly as a reflection of enhanced BAT activation (88). In isolated human brown adipocytes, FGF21 and the muscle-derived hormone irisin have additive effects on norepinephrine-stimulated thermogenesis (89). Clinical trials are currently underway to test the efficacy and safety of FGF21 analogs in human metabolic disease. Initial data on long-acting FGF21 analogs in overweight/obese type 2 diabetic subjects demonstrated a significant decrease in body weight together with improved plasma lipid profiles and elevation of circulating adiponectin levels. However, blood glucose levels were not significantly reduced, whereas markers of bone turnover increased in a dose-dependent manner, which tempered the previous enthusiasm of FGF21-based treatment for metabolic disease (90, 91).

### Pharmacological activation of human BAT and outlook

Given the persuasive evidence from various preclinical and clinical studies that the promotion of BAT function has beneficial effects on total energy metabolism, pharmacologic activation of human BAT holds great promise as a therapeutic concept in metabolic disease. Among pharmacological substances, β3-agonists are well known to induce UCP-1 expression and thermogenesis in vivo in rodents and in vitro in isolated brown adipocytes (4, 92, 93, 94). Although initial attempts with β3-agonists have failed to increase BAT activity in humans (95), a recent study showed that mirabegron, a β3-agonist approved for the treatment of overactive bladder, activates BAT and elevates energy expenditure in humans similar to cold exposure (96). However, the potential cardiovascular side effects of β3-agonists may limit their widespread use in humans. Non-pungent capsaicin analogs (capsinoids) may represent an alternative therapeutic approach. Capsaicin enhances thermogenic gene expression and browning of WAT in mice through stimulation of transient receptor potential channels (97). Moreover, a six-week treatment with oral capsinoids increased cold-induced thermogenesis and energy expenditure in subjects with previously low or undetectable BAT activity. However, the effects were not sufficient to induce a loss of body weight or fat mass (59). In contrast, treatment with the antidiabetic agent liraglutide, which acts as an agonist at the glucagon-like peptide-1 (GLP-1) receptor results in significant weight loss in obese subjects (98). GLP-1-mediated weight loss is mainly driven by delayed gastric emptying and appetite suppression (99). However, a recent study found that energy expenditure is also increased in liraglutide-treated obese patients with type 2 diabetes (100). Interestingly, intracerebral liraglutide administration in mice stimulated BAT thermogenesis and browning of WAT through hypothalamic adenosine monophosphate-activated protein kinase action (100). Whether GLP-1 agonist mediated weight loss seen in humans also involves altered BAT activity is still a matter of investigation. The bile acid chenodeoxycholic acid (CDCA) on the other hand has already been demonstrated to increase BAT activity in mice and humans (101, 102). Ingestion of 15 mg/kg CDCA for two subsequent days significantly stimulated BAT FDG uptake in young healthy women and CDCA treatment in isolated human brown adipocytes enhanced mitochondrial respiration (102). In addition to the activation of the nuclear farnesoid X receptor, bile acids signal through the G protein-coupled receptor TGR5, which leads to thermogenic activation of brown adipocytes through the actions of intracellular thyroid hormones (103). Other intriguing drug targets are bone morphogenetic proteins (BMPs), in particular BMP7 and BMP8b, which are important for brown adipocyte development, the thermogenic response of BAT and the browning of WAT (104, 105, 106). Central actions of BMP8b also include increased sympathetic outflow to BAT, which resulted in weight loss in mice (105).

Although these early findings are very promising, more studies will be needed to assess the potency of these factors to recruit brown or beige adipocytes in humans. Current research is also directed at identifying the cell population in human fat depots that can undergo efficient thermogenic transformation in response to pharmacological stimuli. Even if new therapeutic interventions will achieve successful expansion and activation of BAT in humans, it remains to be seen whether
persistent weight loss and improved glucose metabolism will be the consequence. Given the tight control of energy balance under physiologic conditions, it is possible that increased appetite and food intake will compensate for the energy wasting induced by BAT activation. Hence, a combined intervention targeting both, adipose tissue thermogenesis and central appetite regulation, may represent the most effective therapeutic approach in the fight against obesity and related metabolic complications.

Declaration of interest
The author declares that there is no conflict of interest that could be perceived as prejudicing the impartiality of this review.

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