Brown adipose tissue (BAT) is a specialized organ responsible for thermogenesis, a process required for maintaining body temperature. BAT is regulated by the sympathetic nervous system (SNS), which activates lipolysis and mitochondrial uncoupling in brown adipocytes. For many years, BAT was considered to be important only in small mammals and newborn humans, but recent data have shown that BAT is also functional in adult humans. On the basis of this evidence, extensive research has been focused on BAT function, where new molecules, such as irisin and bone morphogenetic proteins, particularly BMP7 and BMP8B, as well as novel central factors and new regulatory mechanisms, such as orexins and the canonical ventromedial nucleus of the hypothalamus (VMH) AMP-activated protein kinase (AMPK) – SNS – BAT axis, have been discovered and emerged as potential drug targets to combat obesity. In this review we provide an overview of the complex central regulation of BAT and how different neuronal cell populations co-ordinately work to maintain energy homeostasis.

Key words: AMPK, β-adrenoreceptors, brown adipose tissue (BAT), hypothalamus, obesity, orexins, sympathetic nervous system (SNS), thermogenesis, thyroid hormone, uncoupling protein 1 (UCP1)

Brown adipose tissue is a thermogenic machine

Brown adipose tissue (BAT) is a special class of adipose tissue responsible for energy and dissipation, being the most important organ for non-shivering thermogenesis (NST) (1–10). Morphologically, BAT is clearly distinguishable from white adipose tissue (WAT). In particular, brown adipocytes show polygonal shape with multilocular lipid droplets, many large mitochondria, and are richly innervated by sympathetic nerve efferent fibers (2,5,8,11,12), which ensure central control of thermogenesis. WAT, on the other hand, is the primary fat storage organ in the body and consists of unilocular adipocytes with a large lipid droplet for storage of excess calories when energy intake exceeds energy expenditure. Moreover, BAT is highly vascularized to allow the dissipation of generated heat (13), whereas WAT has evolutionarily enabled animals to survive for longer periods without meals, storing energy, mainly as triglycerides, and releasing fatty acids during fasting periods. Despite some initial controversies it is now accepted that brown adipocytes have a developmental origin that largely differs from that of the white adipocytes, as shown by the finding that even before differentiation brown and white fat cells express different genes (14–16).

Recently, evidence for a second type of brown adipocyte has been reported in both rodents and humans. It was found that brown fat cells may appear after thermogenic stimuli at anatomical sites corresponding to WAT. These beige or brite (brown in white) adipocytes derive from precursor cells different from those in classical BAT and are closer to the white adipocyte’s cell lineage (17–21). Despite the fact that the cell origin and anatomical placement are different from BAT, this new lineage has all the morphological and molecular characteristics of classical brown adipocytes present in BAT depots and acts as true thermogenic brown adipocytes (15,22,23). However, the thermogenic
capacity of beige/brite cells in vivo is much lower compared to brown adipocytes (23–26).

**Thermogenic processes in brown adipocytes**

Since the 1970s, BAT has been recognized as the main NST organ. Thermogenesis is facilitated via uncoupling of mitochondrial respiration from ATP production, mediated by uncoupling protein 1 (UCP1), also called thermogenin, which is located in the inner mitochondrial membrane (2,5,12,27). Using electron transport from the oxidation of fatty acids as substrate, UCP1 allows the free movement of protons back across the mitochondrial membrane, avoiding ATP synthesis and instead dissipating the excess of energy as heat (Figure 1) (2,5,28–31). The thermogenic potency of this mechanism is considerable, and fully activated BAT in humans has been appraised to contribute to 5% of the basal metabolic rate (32). By scaling up energetic values obtained in rodents, this effect is theoretically estimated to have the potential to increase daily energy expenditure by up to 20% (33).

BAT function is controlled by both the central and the peripheral nervous system. The sympathetic nervous system (SNS) is essential to activate BAT thermogenesis. An increase in the firing rate of the sympathetic nerves subserving BAT leads to norepinephrine release at the nerve terminal and activation of the β-adrenergic receptors (β-ARs, which are G protein coupled receptors) expressed in the brown adipocytes, mainly the β3 subtype (β3-AR). Upon receptor stimulation, associated protein Gs activates adenylate cyclase (AC), increasing cAMP, which in turn activates protein kinase A (PKA), inducing thermogenesis and downstream activation of p38 mitogen-activated protein kinase (MAPK) (2,34). PKA has both acute and chronic effects on BAT. The acute response of PKA increases lipolysis leading to elevated cytosolic free fatty acid (FFA) level. This process occurs by activation of adipose triglyceride lipase (ATGL), hormone-sensitive lipase (HSL; the activated form being pHSL), and monoacylglycerol lipase (MGL), the three of them sequentially hydrolyzing triglycerides to release FFAs. Carnitine palmitoyltransferase 1a (CPT1a) introduces FFA-CoA into the mitochondria, where FA oxidation leads to the formation of NADH and FADH, which are then further oxidized in the electron transport chain (Figure 1) (2,5,35–38). Purine nucleotides such as GDP and ADPN are bonded to UCP1 in resting conditions, inhibiting it; however, FFAs, which directly activate UCP1, displace the inhibitory nucleotides, increasing UCP1 activity within seconds (2). In addition to acute effects, prolonged stimulation of BAT for hours and days will result in increased amounts of UCP1 protein, mitochondrial biogenesis, and both hyperplasia and hypertrophy of BAT (1,2,5,39). Thus, although the total energy balance depends on many factors (food intake, energy expenditure, BAT thermogenesis, etc.), stimulation of BAT can have an impact on long-term energy balance and hence body weight. BAT not only increases energy expenditure, but also leads to plasma triglyceride clearance (40) and improved glucose homeostasis (41,42) through its ability in uptake and disposal of large quantities of lipid and glucose from the circulation for use as substrates for thermogenesis (2,40,43). On this basis, brown fat has emerged as a target to combat obesity and related metabolic disorders (6,8,33). This concept is of relevance because it is known that increased BAT function is a physiological counter-regulatory mechanism in high-fat diet (HFD)-induced obesity, eliciting recruitment of BAT, including increased UCP1 expression levels (44). However, using BAT as a

![Figure 1. Brown adipose tissue uncoupling and thermogenesis.](image-url)

In brown adipose tissue (BAT), sympathetic stimulation releases norepinephrine (NE) activating β3-adrenergoreceptors (β3-AR) in brown adipocytes coupled to G-proteins, which activates adenylate cyclase (AC), turning AMP into cAMP that in turn activates protein kinase A (PKA). PKA induces lipolysis activating adipose triglyceride lipase (ATGL), hormone-sensitive lipase (HSL; the active isoform being pHSL), and monoacylglycerol lipase (MGL) which hydrolyzes triglycerides, diacylglycerol, and monoacylglycerol, respectively, releasing free fatty acids (FFAs). FFAs are then imported into the mitochondria through carnitine palmitoyltransferase 1a (CPT1a), where they are oxidized (via β-oxidation and further citric acid cycle), leading to the formation of NADH and FADH, which are then oxidized by the electron transport chain. This results in pumping protons out of the mitochondrial matrix and the creation of a proton-motive force that drives the protons back into the mitochondrial matrix through uncoupling protein 1 (UCP1). The energy stored in the proton-motive force is then released, starting mitochondrial heat production. Furthermore, PKA activation increases UCP1 expression via mitogen-activated protein kinase (MAPK) and extracellular signal-regulated kinases (ERK)1/2 pathway responsible for gene transcription, cell growth, cell differentiation and protein synthesis.
target organ for drugs should be approached with care, since recent evidence has demonstrated that increased BAT activity may have harmful side effects. A recent report has demonstrated that atherosclerotic plaque growth was accelerated by cold-induced BAT thermogenesis, indirectly caused by the associated increase in lipolysis. Of note, UCP1 knockout mice were protected from that effect, suggesting that activation of UCP1 function may account for low-temperature-associated cardiovascular risks (45).

**Novel findings about BAT in adult humans**

For a long time, BAT was considered as a tissue relevant only to rodents, hibernating mammals, and newborn humans (1–3,5). However, current evidence demonstrates that BAT is also present in adult humans. Using different approaches, including positron emission tomography (PET) studies alongside 18fluorodeoxyglucose (18FDG) uptake, BAT is found in defined, but dispersed, areas in the body of adult humans, distributed mainly in the cervical, supraclavicular, perirenal, intercostal, and periaortic regions (3,12,46–48). Despite the initial enthusiasm following the identification of BAT in adult humans, two recent publications have reported some unexpected results regarding BAT in people. Indeed, human adipose tissue biopsies from the supraclavicular area have unexpectedly found that the expression profile of some biomarkers was closer to that of beige/brite cells than of that brown cells (18,19), suggesting that human BAT may be composed of mainly beige/brite adipocytes. The process in which precursor cells placed in WAT become beige/brite cells instead of white adipocytes is called ‘browning’, and will be explained in the next section.

**Beige/brite adipose tissue: a new thermogenic organ?**

As mentioned above, there is evidence that beige/brite adipocytes exist in rodents and humans (17–22,49). According to that concept, browning consists of beige/brite adipocytes emerging in white fat depots by cold stimulation. These new cells come from a completely different lineage than the myogenic (Myf5+) lineage, as classical brown adipocytes do (15), but they still exhibit the same properties as BAT with respect to UCP1-mediated thermogenesis in response to cold. In fact, specific molecular markers of beige fat cells, rather than those of the classical brown fat cells, are enriched in the human UCP1-positive tissues, being brown-like adipocytes located in WAT depots but exhibiting the properties of classic brown adipocytes upon cold stimulation (19,22). Recent data demonstrate that both cold exposure and β3-adrenoreceptor agonists induce the production of beige/brite cells (20,23,50–52). Based on the above findings, activation of thermogenesis through stimulation of beige/brite adipocytes in WAT might be a new way to increase energy expenditure. Thus, these beige/brite adipocytes represent a new cell type with therapeutic potential against obesity and related metabolic disease (6,8,53).

**Molecular determinants of browning**

PPARγ, PGC1α, PRDM16, and RIP140

The first indispensable factor for browning is peroxisome proliferator-activated receptor γ (PPARγ), an essential transcription factor for differentiation and survival of both brown and white adipocytes (22,54–57). PPARγ activation also promotes the formation of brown adipocyte-like cells, inducing the expression of UCP1, proliferator-activated receptor-gamma coactivator 1 alpha (PGC1α), and other genes related to mitochondrial biogenesis, as well as those implicated in the repression of WAT genes (57). Recently, it has been demonstrated that PPARγ induction of brown fat genes in subcutaneous WAT is mediated by activation of PRDM16 (PR domain containing 16) (57), a process that is mediated by deacetylation of PPARγ by sirtuin 1 (SIRT1) (58). PRDM16 is the second determinant factor to promote brown adipogenesis in WAT (15,59,60). The adipogenic actions of PPARγ are not restricted to BAT. It is well known that PPARγ is critically required for white adipogenesis, PPARγ2 (instead of PPARγ1) being the more potent adipogenic isoform in vivo (61–65). The specific mechanism by which PPARγ activation drives brown or white adipocytes depends on specific transcription factors. In fact, mesenchymal stem cells give rise to precursor cells of bone, muscle, and fat under appropriate conditions. PPARγ+ CD24 + white adipocyte precursor cells reside in mural cell compartments of the adipose vasculature. White adipocyte differentiation is driven by the transcription factors PPARγ and C/EBPs (CCAAT-enhancer-binding proteins), specially C/EBPα, giving rise to triglyceride-storing WAT. This proliferating WAT precursor cell population constantly reconstitutes WAT depots throughout adult life and can react to increased demand for energy storage with increased differentiation. Brown fat cells share precursors (Myf5+) with muscle cells but not with white adipocytes. Induction of PRDM16 expression in Myf5+ cells directs them to develop into brown fat cells (15,66). In the absence of PRDM16, these precursor cells will develop into muscle cells under the influence of the transcription factors myogenin and MyoD (myogenic differentiation protein) (15,66,67).

PGC1α is increased by cold exposure (68), and controls mitochondrial biogenesis and nuclear respiratory factors, making it indispensable for proper thermogenesis but not for BAT differentiation (56,68,69). PRDM16, similarly to PPARγ, is involved in the regulation of both BAT and WAT genes (70–72). PRDM16 is able to interact with PGC1α and C/EBPs (C-terminal-binding proteins) to activate brown fat genes and suppress white fat genes, respectively (73). Genetic overexpression of PRDM16 increases energy expenditure and improves glucose tolerance in HFD mice through augmented brown-like adipocytes in WAT, while deletion of PRDM16 in vivo has been shown to decrease UCP1 expression (74). Stimulation of BAT-specific genes by PRDM16 requires association with PGC1α and PGC1β (66), which are the third activators of browning as well as coactivators of PPARγ (18,57).

RIP140 (receptor-interacting protein 140) is a nuclear receptor co-regulator highly expressed in metabolic and reproductive tissues (75). Its overexpression inhibits genes involved in mitochondrial biogenesis and oxidative metabolism (76–78). By contrast, deletion of RIP140 increases brown-like adipocytes in WAT (76). RIP140-null mice have a 70% reduction in total body fat mass and have higher oxygen consumption when fed a HFD; these mice are resistant to obesity and have elevated expression of UCP1 in WAT (76,79). In brown adipocytes, RIP140 suppresses UCP1 by antagonizing the binding of PPARα/PGC1α to the UCP1 promoter (80,81). Consistently, its overexpression inhibits genes involved in mitochondrial biogenesis and oxidative metabolism (76–78).

**Irisin: role of exercise**

In 2012, Bostrom and colleagues identified irisin, a muscle tissue-secreted peptide, proposing a possible mechanism by which exercise increases BAT activity (82). During exercise, PGC1α is induced in the muscle of rodents and humans (83,84). As a consequence, fibronectin type III domain containing (FNDC5) is released and further cleaved to irisin. This active form of the protein can act on different tissues, among them BAT, where it increases
its thermogenic activity by increasing UCP1 expression (82,85). Irisin also stimulates browning by increasing the expression of UCP1 and other BAT-genes in subcutaneous WAT through mitogen-activated protein kinase p38 MAP kinase and ERK MAP kinase signaling (85), while genes characteristic of WAT are down-regulated (84). More recently, one report showed that the actions of irisin might be of clinical value since both irisin and fibroblast growth factor (FGF21; see below) are cold-induced endocrine activators of BAT activity in humans (86). However, there are also several studies that have shown controversial results, and some authors failed to document an effect by contraction in circulating irisin levels in humans or an effect on beige/brite differentiation of human preadipocytes (87,88). Therefore, the potential beneficial metabolic actions of irisin during exercise are still under debate, and some issues await clarification. These include the full characterization of the different tissues expressing irisin (89) and, more importantly, the different proteolytic mechanisms involved in its post-translational processing and the generation of putative secreted molecules (90). Thus, it seems obvious that further studies are needed to elucidate, in depth, this field.

**Transient receptor potential V4 channel**

The transient receptor potential (TRP) family of cation channels mediates sensation across a broad physiological range of temperatures. Several TRP members have been described, each of them sensing a particular temperature range. The TRPM8 channel is activated by modest cooling (environmental temperatures < 27°C) (91,92), and TRPM8-deficient mice exhibit a reduced ability to tolerate innocuous cold temperatures (93–95). TRPA1 is activated by colder temperatures (environmental temperatures < 17°C) than TRPM8 (96–98), while TRPV1 is activated by a noxious range of heat (> 43°C) (99,100). TRPV3 and TRPV4 are warm-sensitive channels activated by innocuous warm temperatures, with environmental temperature thresholds of 33–39°C and 25–34°C, respectively (101–105). TRPV4 was first identified as an osmolality sensor (10,106,107), and it is highly expressed in adipose tissue (106), where it acts as a negative regulator of PGC1α, UCP1, and cellular respiration and therefore as an inhibitor of the thermogenic program in WAT (108). Additionally, it potently controls the expression of multiple proinflammatory genes involved in the development of insulin resistance (108). This evidence is also supported by functional data of genetic ablation or pharmacological inhibition of TRPV4, both of which modulate thermogenic and proinflammatory pathways in adipose tissue (108). Furthermore, genetic ablation of TRPV4 induces elevated thermogenesis and protects from diet-induced obesity, adipose inflammation, and from insulin resistance (10,108), suggesting that inhibition of TRPV4 could be a target for treating obesity and type 2 diabetes by inducing a thermogenic program in white adipocytes. Whatever the case, due to its alleged role in regulating osmotic pressure in the kidney, some possible secondary effects should be considered when regarding TRPV4 as a drug target (109,110).

**Natriuretic peptides**

The cardiac natriuretic peptides (NPs), atrial NP (ANP) and ventricular NP (BNP), are potent vasodilators secreted by heart muscle cells in response to high blood volume. A thermogenic effect of NPs, in both mice and human adipocytes, was recently demonstrated (111). Thus, when NPs bind to their receptors (NPR) on brown adipocytes, this activates p38 MAPK for induction of the expression of Upf1, Pgc1α, and other genes implicated in thermogenesis. Chronic administration of BNP also increases browning in white fat and enhanced oxygen consumption and energy expenditure. Accordingly, in mice exposed to cold, in which BAT activity is increased, plasma levels of NPs were elevated (111). The physiological significance of these data is intriguing (9,10), but it would be reasonable to think that BAT thermogenesis would demand higher cardiac activity, and NPs would establish a logic communication between heart and BAT. Further studies are required for a better understanding of this heart–BAT axis.

**Retinoids**

Retinoids, such as vitamin A (retinol), are involved in multiple essential biological functions including fuel metabolism by controlling adipogenesis and energy homeostasis (112–114). Retinoid acid (a metabolite of retinol) is a direct activator of sympathetic activity and an inducer of UCP1 expression in BAT and WAT. In fact, its chronic administration induces browning in white adipocytes (9,10,115,116).

**Prostaglandins**

Prostaglandins (PG) are a group of lipid compounds involved mainly in inflammation, fever induction, and pain. Cyclooxygenase-2 (COX-2) is a key enzyme in PG formation. COX-2 has been described as downstream effector of β-adrenergic signaling in WAT, being required for the induction of brown adipocytes in WAT depots, increasing energy expenditure, and protecting against HFD-induced obesity (10,117). Genetic and pharmacological approaches have shown that COX-2 and prostaglandin E2 (PGE2) are involved in UCP1 expression in white, but not in brown adipocytes, suggesting that the browning process in WAT is COX-dependent (118). Current evidence has also demonstrated that lipocalin prostaglandin D synthase (L-PGDS) plays also a role in fuel utilization by BAT (119). In fact, under cold-acclimated conditions, mice lacking L-PGDS show elevated reliance on carbohydrate to provide fuel for thermogenesis and increased expression of genes regulating glycolysis and de novo lipogenesis in BAT. These transcriptional differences are associated with increased lipid content in BAT and a BAT lipid composition enriched with de novo synthesized lipids. Actually, when L-PGDS knockout mice showed unchanged levels of PGD2 in BAT, this suggests that L-PGDS may act as a lipocalin (lipid binding protein) rather than as a prostaglandin synthase. Moreover, L-PGDS knockout mice exhibit increased expression of genes involved in thermogenesis and increased norepinephrine-stimulated glucose uptake to BAT (119).

**Peripheral regulation of BAT**

**Thyroid hormones**

Thyroid hormones (THs) are produced in the thyroid gland in the form of thyroxine (T4) which is an inactive prohormone with low biological activity. T4 is transformed into the bioactive hormone triiodothyronine (T3) by catalytic action of enzymes iodothyronine deodinases (D1 and D2) expressed in different tissues (120–124). THs play a key role in energy metabolism acting on peripheral tissues, such as liver, pancreas, skeletal muscle, WAT, and BAT (124). Thereby, animal models and patients with hyperthyroidism, characterized by high circulating levels of TH, show weight loss despite increased food intake, due to very high caloric demand (124–130). By contrast, hypothyroidism is associated with decreased metabolic rate, with increased body weight despite reduced food intake (124–126,131–134).

BAT is a direct target of THs and displays high expression of TH receptors (TR). In fact, THs work synergistic with norepinephrine and are required to generate a full thermogenic response (124,130,135). The TRα subtype in BAT ensures maintenance of normal adrenergic thermogenic response, while the TRβ1 sub-
type increases Ucp1 gene expression (136,137). The functional relevance of THs on BAT is supported by animal data (138–142). Genetic and pharmacological approaches have confirmed the role of the different TR isoforms in BAT thermogenesis, showing that TR deficiency leads to hypothermia and cold intolerance associated with reduced BAT thermogenesis due to sympathetically impaired response, although morphology, recruitment, and development of BAT are unaltered (138–142). Furthermore, D2 is well recognized as an important enzyme in BAT physiology for TH-induced thermogenesis (143–146), and studies on genetic models with ablation of D2 expression suggest that D2-catalyzed T4 to T3 conversion is required to support BAT adrenergic responsiveness to generate heat in response to cold exposure (147–149). This evidence has been partially confirmed in humans, where chronic exposure with TH increased both brown adipose tissue activity and volume, as well as the expression of TH-responsive genes, such as Ucp1, leading to improvement in glucose control (150).

**Leptin**

The importance of leptin in the regulation of thermogenesis is well established. Leptin is a circulating hormone/cytokine which is released mainly by subcutaneous white adipose tissue, and has effects in both the central nervous system (CNS) and in peripheral tissues (151–155). Centrally, leptin controls food intake, energy expenditure, fat distribution, insulin sensitivity, FFAs oxidation, and lipolysis in the periphery. It is known that leptin increases SNS activity to BAT (156), and that ob/ob (leptin-deficient) mice, db/db (leptin receptor-deficient) mice, and fa/fa (leptin receptor-deficient) rats have atrophied BAT, with little UCP1 expression, leading to reduced BAT activity and impaired thermogenesis (157,158). The thermogenic effect of leptin is mediated by the melanocortin system, more specifically through stimulation of α-melanocyte-stimulating hormone (α-MSH) release, which then activates sympathetic outflow to BAT. Thus, leptin binding to its receptor, LEPR-B, leads to an increase of proopiomelanocortin (POMC) and a decrease of neuropeptide Y (NPY)/agouti-related protein (AgRP) levels. The overall consequence is triggering of SNS activation, which leads to an increase of UCP1 transcription and thermogenesis in BAT (159).

**Insulin**

Insulin is secreted in pancreatic β-cells in response to circulating glucose increase, acting as an anabolic hormone in peripheral tissues. BAT is one of the most insulin-responsive tissues with respect to stimulation of glucose uptake (2,160). Physiological conditions in which plasma insulin levels are elevated (refeeding) show increased glucose uptake into BAT (2,161,162), whereas states with low insulin levels (starvation or fasting) demonstrate reduced glucose uptake (2,162,163). In rodents, it was also found that insulin can promote thermogenesis through its action in the brain, triggering sympathetic activation to BAT (164). Interestingly, insulin infusion increased glucose-uptake in BAT to levels similar to that in skeletal muscle, which could be associated to the high expression of GLUT4 in BAT, although it was not accompanied by concomitant thermogenesis unlike cold exposure (165). Phosphatidylinositol-3-kinase (PI3K), a major kinase mediator of insulin and insulin-like-growth factors, is counteracted by phosphatase and tensin homolog (PTEN). Recent data showed that PTEN positively regulates a BAT-selective thermogenic program by blocking the PI3K pathway (166). Importantly, pharmacological PI3K inhibitors increase BAT thermogenesis and whole-body energy expenditure (166). However, the systemic significance of brown adipocytes for insulin-induced glucose clearance was demonstrated using transgenic mice lacking the insulin receptor in BAT (BATIRKO) in which the weight of BAT is decreased and enzymes of fatty acid synthesis are decreased (without change in the number of brown adipocytes), suggesting that insulin action is necessary for lipid accretion in BAT (41). In parallel, these mice develop an insulin-secretion defect, providing evidence that BAT plays a role in the regulation of insulin secretion (41).

**Adiponectin**

Adiponectin (ADPN; also called adipocyte complement-related protein (Acrp30), apM1, or adipoQ) is a 244-amino-acid protein secreted from adipose tissue (167–170), placenta (171), and cardiomyocytes (172), among other tissues. ADPN increases fatty acid combustion and energy consumption via PPARα activation, leading to decreased triglyceride content mainly in the liver and skeletal muscle through activation of AMP-activated protein kinase (AMPK; see below) activity (173). Adiponectin also increases hypothalamic AMPK activity via its receptor AdipoR1 to stimulate feeding (174–176). Recent evidence has shown that ADP reduces thermogenesis by inhibiting BAT function. ADPN treatment reduces PKA signaling (177) and UCP1 expression in BAT (45,174,178). In keeping with this, genetic ablation of the Adpn gene increases body temperature, increases Ucp1 expression in BAT, and more brown-like structure in inguinal fat compared to wild-type mice (179). Reconstitution of ADPN in these animal models blunted β-adrenergic receptor agonist-induced thermogenesis of interscapular BAT (179). Taken together, these data suggest that the suppressive effect of ADPN on BAT thermogenesis could be associated to an increase in hypothalamic AMPK activity (see below), an idea that needs to be confirmed.

**Resistin**

Resistin (RSTN) is an adipokine, originally identified in adipose tissue (180) but expressed in a variety of tissues, including the hypothalamus (181). Recently it has been demonstrated that resistin activates the ERK1/2 signaling pathway in the hypothalamus, involved in diverse cellular functions including cell growth, proliferation, and neuronal activity, which, in turn, reduces BAT thermogenesis (182,183). This evidence, alongside RSTN’s anorectic role and its effect on hypothalamic AMPK (184,185), points towards an extensive role of this adipokine in the modulation of BAT function.

**Adrenal steroids**

Glucocorticoids (such as cortisol and corticosterone) and mineralocorticoids (aldosterone) are steroid hormones secreted in the adrenal cortex and involved in metabolism, cell growth, and regulating ion and water transport. Cortisol acts by binding to the glucocorticoid receptor (GR), while aldosterone acts through the mineralocorticoid receptor (MR). The presence of GR on brown adipocytes was described many years ago (186); its activation in BAT reduces UCP1 expression (187,188). MR is also expressed in brown fat (189). Subsequently, studies in cells show that aldosterone inhibited the expression and function of UCP1 in brown adipocytes (187), while aldosterone treatment induced a significant increase of triglyceride’s accumulation in BAT and increased expression of adipogenic genes (lipoprotein lipase-Lpl-, Pparγ) (190). These data suggest that GR and MR signaling in BAT appears to act as a pivotal signal favoring lipid storage at the expense of heat production, hence promoting in brown adipocytes a specific function of white adipocytes.

**Fibroblast growth factor**

FGF21 is a metabolic regulator involved in the control of glucose homeostasis, insulin sensitivity, and ketogenesis, mainly
produced in the liver from where it is released into the blood (191–194). There is evidence that FGF21 is also involved in BAT-induced thermogenesis (195). Current data have provided a mechanistic explanation, showing that induction of cAMP in brown adipocytes induces FGF21 release from BAT in response to noradrenergic stimulation (196). FGF21 expression is under the control of PPARα and FFAs availability, and has also been reported to promote thermogenic activity, increasing energy expenditure, and UCP1 expression in vivo as well as in vitro in WAT (197). In fact, mice deficient in FGF21 display an impaired ability to adapt to chronic cold exposure, with diminished browning of WAT (197). Adipose-derived FGF21 increases the expression of UCP1 and other thermogenic genes in fat tissues, regulating this process, at least in part, by enhancing adipose tissue PGC1α protein levels and also cyclic adenosine 5’-monophosphate response element-binding protein (CREB) and downstream genes associated with oxidative metabolism (195,198,199).

Recent findings have shown that FGF21 also exerts central actions on energy homeostasis. In HFD-fed rats, FGF21 intracerebroventricular (ICV) injection increases feeding, elevates energy expenditure, and improves insulin sensitivity, although with no changes in body weight (200). On the other hand, genetic models of β-Klotho (transmembrane protein that acts with FGF receptors) ablation in the suprachiasmatic nucleus of the hypothalamus (SCN) and the dorsal vagal complex (DVC) simulate features of physiological starvation response, such as increased glucocorticoid levels, suppressed physical activity, and altered circadian behavior (201). Overall, this evidence has led to the proposal of FGF21 as a novel appealing therapeutic reagent for type 2 diabetes mellitus and obesity. However, its clinical development has been challenging due to its poor pharmacokinetics (202).

Central regulation of thermogenesis: the anatomical issue

Normothermia in a cold environment is maintained in part through metabolic activation of BAT in order to generate heat, besides skin vasoconstriction to avoid heat loss. Both processes are regulated by SNS and accompanied by adrenergic cardiac stimulation, to improve distribution of heat generated in thermogenic tissues (203–206). Furthermore, somatic motor nerves promote the generation of heat by skeletal muscle through shivering (207). In addition to these changes, cold responses engage activation of the thyroid and adrenal axes (208,209), All of these changes are co-ordinated at the central level, and in this section we will describe the main central circuits regulating BAT thermogenesis (5), summarized in Figures 2 and 3.

The preoptic area: the temperature central sensor

For many years the preoptic area (POA), also called anterior hypothalamic nucleus, has been considered the only region in the brain responsible for the control of thermoregulation, where cold-sensitive neurons are located (210–214). The POA also receives input from thermosensitive areas elsewhere in the body. Cold and heat receptors in the abdominal viscera send temperature information through splanchnic and vagus nerves to the CNS, where it is integrated into the final response (211,215). Temperature changes could also be sensed by the spinal cord, which contains thermoreceptors detecting cold environment (216). It is well known that electric stimulation of POA activates BAT (217,218). The POA also contains heat-sensitive neurons whose tonic discharge is reduced by skin cooling and whose thermosensitivty to preoptic temperature is increased when the skin is cooled (219). As a result, skin cooling or direct cooling of POA neurons induces sympathetic thermogenesis in BAT, as well as shivering thermogenesis (212,220). Within the POA cold appears to signal mainly to the median part (MnPO), where glutamatergic stimulation with N-methyl-D-aspartate (NMDA) induces physiological responses mimicking a cold-defensive response (221). On the contrary, stimulation of the medial POA (MPO) or lateral POA (LPO) does not recapitulate that effect (221). Furthermore, the POA projects to other central nuclei, modulating thermogenesis. Indeed, destruction of the ventromedial nucleus of the hypothalamus (VMH; see below) abolishes the ability of external cooling signaling to POA to stimulate BAT (222,223), suggesting that POA activation of BAT is mediated by the VMH.

Febrile responses are also mediated by the POA. Fever is a physiological condition that requires an increase of body temperature in response to endogenous pyrogens released, for example, during infection. This provides an optimal hyperthermic environment for mounting host defenses against invading bacteria or viruses, while reducing pathogen viability (213). Prostaglandins (PG), which are involved in mediating the febrile response, are synthesized in the brain vasculature and in peripheral tissues in response to immune signals. Besides previously mentioned effects on BAT thermogenesis and browning, there are studies using lipocalin prostaglandin D synthase (L-PGDS; which synthesizes D-series PG) demonstrating that L-PGDS mRNA in BAT is strongly and positively correlated with activation of BAT metabolism (119,224). Prostaglandins E2 (PGE2) also stimulate POA, activating BAT thermogenesis in a CAMP-dependent manner (225,226). In addition, the inhibitor of phosphodiesterase (the enzyme degrading cAMP), blunts fever induced by intra-POA PGE2 (227). Thereby POA is known as a ‘fever or thermoregulatory center’. The POA population with EP3 subtype of PGE receptor (EP3) is mainly GABAergic and projects to neurons at the dorsomedial nucleus of the hypothalamus (DMH) and the rostral raphe pallidus nucleus (rRPa), antagonizing GABA receptors and inducing fever by activating BAT thermogenesis (228). Fibers connecting POA to the DMH and rRPa will be explained below.

The ventromedial nucleus of the hypothalamus: the integrator center of peripheral signals

The VMH was the first identified as a hypothalamic site involved in thermoregulation. It was reported that electrical stimulation of that nucleus increased interscapular BAT temperature, and that this effect was abolished by β-adrenergic blockade (217,229–233). Also, GABA agonists applied into the VMH abolish the prostaglandin E2-induced BAT activation via a sympathetic efferent mechanism (225,234). On the other hand, glutamate injection into the VMH was shown to activate BAT (233–236), an effect promoted also by hydroxybutyrate (237), norepinephrine, serotonin, and tryptophan (238). Anatomical data have also demonstrated a link between the VMH and BAT. First, there is evidence that the VMH can be trans-synaptically infected from BAT with pseudorabies virus. For instance, 6 days after BAT inoculation with a transneuronal viral tract tracer, Bartha’s K strain of the pseudorabies virus, the infection penetrates into the VMH (239), an effect considered at some point controversial (240). Second, there are projections from the VMH to autonomic centers (2,127,241,242). In addition, there are several brainstem areas, such as the raphe pallidus (RPa) and inferior olive (IO), two nuclei which have been functionally linked to the regulation of BAT thermogenesis (240,243–246), where VMH neurons have been postulated to relay to modulate SNS activity (2,127,241). However, the role of the VMH in the regulation of BAT activity remains controversial due to the lack of
information about the precise neuronal pathways that connect the VMH to BAT (247).

More recently, genetic evidence has also supported the role of the VMH in the modulation of BAT thermogenesis. For example, VMH-specific knockout of steroidogenic factor-1 (SF-1) shows lower energy expenditure and expression of UCP1 in BAT (248,249). On the other hand, silencing estrogen receptor α (ERα) in the VMH increases body weight gain, visceral adiposity, hyperphagia, hyperglycemia, and reduces thermogenesis in BAT, leading to metabolic syndrome (250). Moreover, mice bearing conditional ERα knockout in SF1 neurons exhibit reduced plasma levels of norepinephrine and thermogenesis in BAT, with decreased UCP1, PPARγ, PGC1α, and β3-AR expression (251). These data support the idea that the VMH is a key modulator of BAT. We have extensively studied the mechanisms within the VMH that regulate BAT activity. Our studies point towards AMPK in the VMH as a key negative regulator of sympathetically activated BAT thermogenesis, integrating peripheral signals, such
Central regulation of BAT

As thyroid hormone, estradiol, and BMP8b, as well as drugs such as nicotine (127,241,252,253).

The dorsomedial nucleus of the hypothalamus

The DMH plays key roles in feeding and metabolic regulation associated with ingestive behavior (247,254,255), which is closely related to thermoregulation. After many years of controversial data, there is evidence that activation of neurons in the DMH elicits an increase in body temperature through a classic thermoregulatory mechanism (228,247,256). Excitatory amino acid receptors in the DMH mediate prostaglandin-evoked thermogenesis in BAT (257). In keeping with this, disinhibition of DMH neurons increases BAT sympathetic activation and thermogenesis (228,256). The cellular basis of this mechanism is not totally clear, but GABAergic neurons in the POA appear to provide a key source of the tonic inhibitory input to sympathetic nerve fibers in the DMH (228,258 – 261). However, neurons in the DMH do not project directly to BAT sympathetic fibers, but they rather send monosynaptic projections to the rRPa that have been involved in mediating the effects of DMH neurons on BAT thermogenesis. Thus, disinhibition of neurons in the DMH activates glutamate receptors in the rRPa, triggering BAT sympathetic

Figure 3. Interactions of hypothalamic nuclei regulating BAT thermogenesis. All complex peripheral signals are integrated at the central nervous system regulating the final output to brown adipose tissue (BAT). The main brain region to integrate them is the hypothalamus where different nuclei play key roles in the thermogenic response. Other brain sites, such as the preoptic area (POA) and rostral raphe pallidus (rRPa), also are involved in this network regulation. Therefore, cold exposure induces neuronal activation of POA, which projects to other neurons in the ventromedial nucleus of hypothalamus (VMH), leading to activation of sympathetic fibers to BAT. There are several lines of evidence showing that this mechanism involves AMP-activated protein kinase (AMPK), which can be regulated by glucagon-like-peptide 1 (GLP-1), bone morphogenetic protein 8B (BMP8B), thyroid hormones (THs), or estradiol (E2), all of which induce sympathetic activity to BAT by decreasing AMPK activity. Some evidence supports that ghrelin may well be involved in activation of BAT thermogenesis; however, definitive proof is lacking. The dorsomedial nucleus of hypothalamus (DMH) is highly involved in BAT regulation. The DMH projects fibers to the rRPa inducing BAT activity by increasing sympathetic tone. The paraventricular nucleus of hypothalamus (PVH) is also involved in the modulation of BAT thermogenesis by projecting inhibitory fibers to the rRPa. The lateral hypothalamic area (LHA) contains orexin (OX-A and OX-B)-expressing neurons which project to the VMH and the rRPs, activating BAT thermogenesis. The arcuate nucleus of hypothalamus (ARC) is characterized by the presence of an orexigenic neural population (coexpressing neuropeptide Y (NPY) and agouti-related protein (AgRP)) and an anorexigenic neuron population (coexpressing proopiomelanocortin (POMC) and cocaine- and amphetamine-regulated transcript which are inversely regulated by leptin. Leptin induces POMC expression and releasing of α-melanocyte-stimulating hormone (α-MSH), which binds to melanocortin-4 receptor (MC4R) within PVH, activating sympathetic fibers to BAT. Leptin also inhibits NPY and AgRP which inhibits sympathetic neurons to BAT trough PVH fibers. 3V = third ventricle; β3-AR = β3-adrenergic receptor; ERα = estrogen receptor alpha; GABA-R = GABA receptor; BMP8B = bone morphogenetic protein 8B receptor; GLP1-R = GLP-1 receptor; Glu = glutamate; LEPR-B = leptin receptor isoform B; SNS = sympathetic nervous system; TR = thyroid hormone receptor; PGE2 = prostaglandin E2; EP3 = prostaglandin E receptor type 3.
activation and thermogenesis (262). Functional data support this concept; in fact, inhibition of neurons in the DMH or blockade of local glutamate receptors in the DMH reverses febrile and cold-evoked excitations of SNS and subsequent stimulations of BAT thermogenesis (256,262,263). Other data suggest that the IO may also have a role in the SNS control of BAT and that this nucleus may also be involved in functional interactions between the motor and thermoregulatory systems, through the DMH in the same way as the rRPa (5,245). Overall, the reported evidence so far suggests that, in a normothermic environment, GABAergic neurons in the POA tonically inhibit the DMH. However, in a cold environment, signals from somatosensory nerves in the skin activate GABAergic neurons suppressing the tonic firing of the GABAergic neurons in the POA, leading to disinhibition of DMH neurons, which in turn stimulate neurons in the rRPa controlling BAT sympathetic tone and the IO, and therefore stimulating thermogenesis.

Recent findings raise the possibility of alternative mechanisms through which the DMH activates BAT thermogenesis. This is based on the finding that there are leptin receptors on the DMH which are involved in the stimulation of the rRPa and consequent sympathetic activation of BAT (264,265). Finally, there is evidence suggesting that knockdown of NPY specifically in the DMH promotes both the development of brown adipocytes in WAT and also BAT activity, increasing thermogenesis and energy expenditure (266). Furthermore, this manipulation prevents HFD-induced obesity and improves glucose homeostasis (266). Leptin is probably involved in this process by activating NPY in the DMH (267), although more studies are needed to elucidate this mechanism. Finally, the hypothalamic factor modulating the hypothalamic–pituitary–adrenal (HPA) axis, namely, corticotrophin-releasing hormone (CRH), increases BAT thermogenesis through the DMH and mRPa (268).

The paraventricular nucleus of the hypothalamus (PVH) plays an important role in energy homeostasis; however, its possible involvement in the regulation of thermogenesis and energy expenditure in BAT has been controversial. Because PVH neurons that project to the spinal sympathetic preganglionic cell column are activated during fever (269), and lesions of the PVH attenuate fever (270,272), it was initially thought that neurons in the PVH are activated during fever (269), and lesions of the PVH attenuate fever (270,272), it was initially thought that neurons in the PVH played a role in the excitation on BAT thermogenesis. This is again a controversial issue. Acute administration of the pseudorabies virus (239,273,274) or the H129 strain of the herpes simplex virus-1 (HSV-1) (275) demonstrates that the PVH contains a large number of melanocortin-4 receptor (MC4R)-expressing neurons that project to BAT. Thus, although PVH involvement in the regulation of BAT function remains unclear, existing and emerging data suggest that the PVH exerts a negative regulation on BAT thermogenesis.

The arcuate nucleus of the hypothalamus (ARC)
The ARC is considered as a 'master control center of feeding', containing neural populations inducing orexigenic actions (NPY/AgRP) and also neural populations inducing anorexigenic actions (α-MSH, the product of POMC, as well as CART) (255,297–302). However, there is evidence that ARC also regulates BAT thermogenesis. In this sense, it has been demonstrated that the ARC orexigenic population inhibits thermogenesis and that partial loss of AgRP/NPY neurons leads to a lean, hypophagic phenotype, also characterized by sympathetic activation of BAT (303,304).

Similarly to the PVH, the major implication of the ARC in thermogenesis regulation is linked to the central melanocortin system. Thus, loss of function of the key players of this system, such as POMC and MC4R, has been reported to cause massive obesity in rodents and humans (305,306). In this regard, MC4R deficiency has been shown to blunt the ability of leptin to increase UCP1 expression in BAT and WAT (307). Given that the ARC is the major site for regulation of physiological processes by leptin, the possible role of this nucleus in the integration of leptin actions on thermogenesis is also of interest. Upon activation by leptin, POMC neurons in ARC release the catabolic fragment α-MSH from nerve terminals to increase UCP1 expression in BAT (307). Recent data have demonstrated that leptin receptors in the ARC are necessary for leptin-induced increases in BAT sympathetic discharge; in keeping with this, deletion of leptin signaling in the ARC abolishes sympathetic activation to BAT (308).

The chemical signature of the ARC populations regulating BAT is controversial, and it seems that more than one neuronal type is taking part in this control. Mice lacking synaptic GABA release from RIP-Cre neurons in the ARC have reduced energy expenditure, become obese, and are extremely sensitive to HFD-induced obesity (309). In the same model, leptin's ability to stimulate thermogenesis, but not to reduce feeding, is markedly attenuated. Acute, selective activation of arcuate GABAergic RIP-Cre neurons, which monosynaptically innervate BAT neurons, rapidly stimulates brown fat and increases energy expenditure but does not affect feeding. Importantly, this response...
is dependent upon GABA release from RIP-Cre neurons. This evidence demonstrates that GABAergic RIP-Cre neurons in the ARC selectively drive energy expenditure, contribute to leptin's stimulatory effect on thermogenesis, and protect against diet-induced obesity (309).

Since it is known that thermogenesis can be induced by diet, it could be hypothesized that activation of leptin receptors in the ARC could be involved in diet-induced BAT activity. Although leptin levels are higher in obese subjects due to increased WAT mass, the leptin resistance in these patients likely avoids the capacity of leptin to activate GABAergic RIP-Cre neurons in the ARC, and thereby does not activate thermogenesis through PVH connections. On the other hand, mice lacking RIP-Cre neurons are extremely sensitive to diet-induced obesity due to a defect in diet-induced thermogenesis (290). However, other mechanisms involved in diet-induced BAT activation still remain unclear. Recent data using genetic mouse models of conditional creatine kinase knockout mouse also indicate that NPY reduction in the ARC is coupled to increased UCP1 expression in BAT, resulting in activation of thermogenesis, and suggesting that NPY in the ARC negatively regulates BAT function (310). The same authors recently reported the implication of Rho-kinase 1 (ROCK1) from ARC neuronal population in the regulation of BAT thermogenesis. ROCK regulates leptin action and body weight homeostasis, and genetic deficiency of ROCK1, in POMC and AgRP neurons, promotes obesity and decreased leptin sensitivity (311), also decreasing the effect of leptin on the activation of the thermogenic program. Finally, mice with an AgRP neuron-specific deletion of vesicular GABA transporter are lean, resistant to obesity, and show increased BAT activity (312).

Others central factors influencing BAT thermogenesis: novel targets to combat obesity?

The VMH AMPK–SNS–BAT axis: a canonical regulator of energy balance

Besides the classical neuropeptide-based theory for both feeding and energy expenditure control, recent data demonstrate that basic cellular sensors, such as AMPK and the mechanistic target of rapamycin (mTOR), play a major role in the modulation of energy balance (300,313–317). At hypothalamic level, the AMPK pathway is a canonical route regulating energy homeostasis, by integrating peripheral signals, such as hormones and metabolites with hypothalamic networks. AMPK is a cellular gauge that is activated in conditions of low energy, increasing energy production and reducing energy-wasting (300,313,314,316). AMPK is an important regulator of fatty acid biosynthesis, suppressing de novo lipogenesis by inactivation of acetyl-CoA carboxylase (ACC) by phosphorylating it (300,314). AMPK also regulates neuropeptide expression in the ARC (318–320).

Current evidence from our group has linked hypothalamic AMPK with BAT thermogenesis through modulation of the SNS. Although the first evidence linking hypothalamic AMPK with energy balance was mainly focused on feeding (300,318–320), it has become evident that AMPK also modulates the other side of the balance, namely energy expenditure. Null mice of protein tyrosine phosphatase 1B (PTP1B, which acts as an inhibitor of both insulin and leptin receptor signaling) show decreased body weight and are resistant to diet-induced obesity, associated with decreased hypothalamic AMPKα2, activity and increased BAT thermogenesis (321). This led to the speculation that hypothalamic AMPK may control BAT thermogenesis through the SNS, although it was not shown by those authors. This relationship was further explored by our group, and we demonstrated that decreased AMPK activity, specifically within the VMH, activates BAT thermogenesis through increased SNS outflow (127,241,252,308). Notably, this pathway, which was initially described for central effects of thyroid hormones on energy balance (127), is also shared by leptin (322), BMP8B (bone morphogenetic protein 8B) (241), estradiol (E2) (M.L., unpublished data), glucagon like peptide-1 (GLP-1) (R.N., unpublished data), and nicotine (252), which have led us to propose the VMH AMPK–SNS–BAT axis as a canonical mechanism modulating energy homeostasis (124), with possible implications in the treatment of obesity. This was demonstrated by using constitutive genetic activation of AMPK in the VMH which blocked the actions of the factors described above (127,241,252,322). Additional studies are ongoing to test whether other hypothalamic sensors, such as uncoupling protein 2 (UCP2) and mTOR (M.L., unpublished observations), are also involved in the modulation of BAT.

Orexins

Orexins (OX-A and OX-B), also named hypocretins, are produced in neurons located in the lateral hypothalamic area (LHA), the DMH, and perifornical hypothalamus (323–325). It is known that activation of the LHA promotes BAT function (326). In keeping with this, orexins have important roles in energy balance and feeding (323–325,327), and compelling evidence derived from genetic mouse models has suggested that orexins promote energy expenditure (328,329) through modulation of locomotor activity and BAT thermogenesis (330–332). In fact, recent data suggest that orexins are required for BAT development, differentiation, and function (330). Moreover, lack of orexin's action compromises energy balance, as demonstrated in orexin knockout mice, which are prone to diet-induced obesity, when compared with normal mice (330). In vitro data pointed to a direct effect of orexin on differentiation of new brown adipocytes (330,333). Morphological data have also identified a role for orexin neurons in the regulation of BAT thermogenesis via the SNS, involving direct projection from the LHA to the rRPa, that activates its efferents to premotor sympathetic neurons via a catecholaminergic mechanism (273,334–337). Consistent with this, central injection of OX-A or OX-B, activation of orexin neurons in the LHA, or specific administration of OX-A in the VMH or the rRPs increased BAT thermogenic activity (326,335–338). This effect can be reproduced by administration into the diagonal band of Broca (339). Of note, the thermogenic effect of OX-A displays a very precise anatomical pattern, since its administration into the PVH does not impact BAT (283). The relevance of the thermogenic effect of orexin remains inconclusive but it has been speculated that it may be acting as an adaptive response to stress (340).

A protective role in aging-decreased thermogenic capacity was also recently suggested for orexins (341). The aging process causes an increase in percent body fat, but the mechanism remains unclear. Current data from rodent experiments have shown that aging is associated with defective differentiation of BAT, alongside morphologic abnormalities and thermogenic dysfunction in rodents (341) and humans (47,342,343). In fact, interscapular brown tissue in aged mice is progressively populated by adipocytes, bearing white morphologic characteristics (341). Aged mice also fail to mobilize intracellular fuel reserves from brown adipocytes and exhibit deficiency in homeothermy. Of note, those effects are reversed by OX-A administration, while they are accelerated in mice in which OX-producing neurons were ablated (341). Further work will be required to address whether the orexin...
system may be targeted for intervention to reverse age-associated increase in fat mass. In this regard, aging is also associated with important changes such as leptin resistance, blunting normal central and peripheral functions of leptin (344,345). Thus, restoration of leptin sensitivity during aging might be another possibility to increase BAT function and reduce the amount of fat. Also, ablation of the ghrelin receptor (growth hormone secretagogue receptor, GHS-R) in old mice increases energy expenditure and resting metabolic rate and prevents age-associated decline in UCP1 expression (346). These results suggest that the ghrelin/GHS-R system might also be of clinical relevance. Finally, reduced nuclear SIRT1 activity initiates age-related mitochondrial decline (347,348), so it is also tempting to speculate that the reactivation of SIRT1 in old age might restore BAT activity.

Bone morphogenetic proteins (BMPs)

BMPs belong to the transforming growth factor superfamily and have been recently discovered to be key actors in the differentiation and development of BAT (241,349). Recently, it has been discovered that BMP8B is expressed in BAT and also in the hypothalamus. Notably, central administration of BMP8B induces thermogenesis, and increased core temperature, leading to weight loss. In the VMH, this action of BMP8B is AMPK-dependent, and increases sympathetic outflow to BAT, without altering feeding behavior (241). This evidence, alongside our data showing that nicotine stimulates thermogenesis through the same mechanism (252,253), indicates that this pathway may be a useful target for obesity treatment. Likewise, systemic administration of BMP7 reduces body weight gain at least in part due to induction of BAT-mediated energy expenditure (349). Stimulation of β3-adrenoreceptors and BMP7 synergistically increase UCP1 expression in WAT, suggesting that BMP7 may increase the sensitivity of adipocytes to sympathetic tone, like BMP8B (350). Whether BMP7 actions are mediated by the VMH AMPK–SNS–BAT axis remains unclear, but it is a possibility worth investigating. In this sense, BMP receptors are present in different hypothalamic regions involved in metabolic control (241). Thus, it seems that a cross-talk between BMP signals and SNS activation on peripheral and central levels may be a critical way to modulate global thermogenic capacity.

Besides its central action, BMP8B appears to be produced by mature brown adipocytes and its expression regulated by temperature and other metabolic cues, suggesting that it is under the control of central temperature sensing (241). By contrast, BMP7 is produced by the stromal vascular cells within the adipose tissue and may serve as a niche factor to drive the tissue-resident progenitors to differentiate into the brown lineage (350).

Glucagon-like peptide 1

Another recently reported factor that influences central regulation of BAT thermogenesis is GLP-1, which when directly administrated into the brain reduces body weight by increasing BAT thermogenesis, a process correlated with increased activity of sympathetic fibers innervating BAT (351). This finding suggests that increased BAT thermogenesis may be an additional mechanism in which pharmacological stimulation of GLP-1R controls energy balance (351,352). However, the exact brain regions and pathways through which GLP-1 stimulates BAT activity are unreported, although evidence from our group indicates that this effect is mediated by modulation of AMPK in the VMH (R.N., unpublished data). Taking into account that long-acting GLP-1 agonists are now at the forefront in the clinical management of type 2 diabetes and have been shown also to decrease BMI, studies addressing and uncovering their effects on energy expenditure are eagerly awaited.

Amylin

Amylin is a pancreatic hormone that was recently shown to be involved in central regulation of thermogenesis (353–357). Its ICV administration reduces body weight and food intake and increases body temperature by inducing sympathetic activity to BAT. These effects are blocked by pharmacological antagonism of amylin receptors, supporting their implication in BAT thermogenesis activation (357). In line with these data, neuronal overexpression of the amylin receptor subunit, RAMP1 (receptor-activity modifying protein 1) leads to higher energy expenditure, as indicated by increased oxygen consumption, elevated body temperature, and higher sympathetic tone subserving BAT. Consistent with this, the nestin/hRAMP1 transgenic mice had elevated BAT mRNA levels of PGC1α, UCP1, and UCP3, which can all be reversed by chronic blockade of sympathetic signaling (356). These data have a potential clinical impact, since amylin and its analogs have recently emerged as potential antiobesity drugs. In particular, the demonstration that amylin can restore or enhance leptin sensitivity in obese patients has raised interest in the CNS-mediated physiological effects of amylin and its role in the pathogenesis of obesity (356–359). Pramlintide is the clinically available amylin analogue currently approved for the treatment of patients with type 1 diabetes mellitus or type 2 diabetes mellitus who have not achieved optimal glycemic control after insulin therapy. Used as an adjunct to lifestyle changes, this drug decreased food intake and body weight in obese individuals followed for 12 months (360). In another clinical study conducted in obese or overweight subjects, the combined treatment with leptin and amylin for 4 weeks caused more weight loss than in the individual treatment groups (361). Therefore, these findings support amylin as a promising antiobesity target, particularly for use in combinatorial therapy, a strategy that is now arising to maximize weight reduction (359,362). Whatever the case, an exact knowledge of the molecular underpinnings modulating its actions will be essential to refine the design of further antiobesity therapies. Further work will be required to assess the exact brain populations and molecular mechanisms involved in the control of BAT activity by amylin. Investigating the role of AMPK-expressing neurons in the VMH is an interesting strategy.

Summary and conclusions

BAT is a key organ in the maintenance of temperature through dissipation of heat by mitochondrial uncoupling. Although the ‘classical’ central control of BAT function was mainly based on the modulation of sympathetic outflow, the molecular details of that interaction have remained elusive. Recent developments have demonstrated that ‘traditional’ hormones, such as leptin (308), thyroid hormones (127), estradiol (250,251) (M.L., unpublished observations), and GLP-1 (352) (R.N., unpublished observations), as well as ‘new’ peripheral molecules, such as amylin (356,357,359), nesfatin-1 (363), cannabinoids (364,365), irisin (84), FGF21 (196), BMP7 (349), and BMP8B (241), modulate BAT function by acting on several hypothalamic nuclei. Thus, AMPK in the VMH (127,241,252,322), NPY neurons in the DMH (266), GABAergic neurons in the ARC (309), and orexin neurons in the LHA (328–331), among others, play a major and canonical role in the modulation of BAT thermogenesis. However, although the recent developments in this field have
generated a lot of excitement, some doubt still exists with regard to the possibility of targeting BAT thermogenesis as a mechanism to treat obesity (5,6,8,33,53). Novel techniques, such as optogenetics, will clarify with more detail the existing mechanisms controlling BAT activity. Additionally, it is likely that new circuits implicated in thermogenesis will be also identified. The use of state-of-the-art methods will give us valuable and accurate information regarding the molecular targets to stimulate thermogenesis. Although the existence of cold-sensitive neurons has been documented long ago, the recent characterization of temperature-sensitive receptors open up new avenues to uncover the neural mechanisms involved in adaptation to changes in environment. Additional work will be necessary to demonstrate if our current hope in mechanisms controlling thermogenesis will be of valuable clinical therapeutic application in the future.

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