Thyroid hormones in the regulation of brown adipose tissue thermogenesis

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Abstract

A normal thyroid status is crucial for body temperature homeostasis, as thyroid hormone regulates both heat loss and conservation as well as heat production in the thermogenic tissues. Brown adipose tissue (BAT) is the major site of non-shivering thermogenesis and an important target of thyroid hormone action. Thyroid hormone not only regulates the tissue’s sensitivity to sympathetic stimulation by norepinephrine but also the expression of uncoupling protein 1, the key driver of BAT thermogenesis. Vice versa, sympathetic stimulation of BAT triggers the expression of deiodinase type II, an enzyme that enhances local thyroid hormone availability and signaling. This review summarizes the current knowledge on how thyroid hormone controls BAT thermogenesis, aiming to dissect the direct actions of the hormone in BAT and its indirect actions via the CNS, browning of white adipose tissue or heat loss over body surfaces. Of particular relevance is the apparent dose dependency of the observed effects, as we find that minor or moderate changes in thyroid hormone levels often have different effects as compared to high pharmacological doses. Moreover, we conclude that the more recent findings require a reevaluation of older studies, as key aspects such as heat loss or central BAT activation may not have received the necessary attention during the interpretation of these early findings. Finally, we provide a list of what we believe are the most relevant questions in the field that to date are still enigmatic and require further studies.

Introduction

The maintenance of body temperature according to a central set point is crucial for several physiological processes and largely regulated by balancing heat production as well as heat dissipation. Particularly the latter process is of great relevance in humans, which possess a large surface area that unlike in rodents is only sparsely covered with fur or hair. Therefore, vasodilation is an important response to dissipate excess heat, that is, during physical exercise, and vasoconstriction often occurs to conserve heat, that is, during cold exposure. Simultaneously, heat production processes are regulated accordingly, most prominently the induction of shivering in cold environments. However, also other means can significantly contribute to maintain body temperature including behavioral changes such as moving less in the heat or cuddling in the cold (1).

In the past decade, a special research focus has been put on the mechanisms underlying heat generation, as these processes could theoretically be used to fight obesity and metabolic disorders by burning excess calories. In humans, skeletal muscle is the primary thermogenic tissue, capable of providing rapid heat by shivering. However, in small mammals and newborn humans, the specialized brown adipose tissue (BAT) is the main thermogenic organ, providing a more long-lasting and sustainable source of converting stored energy into heat via non-shivering thermogenesis (2). Thus, BAT constitutes a potential...
anti-obesity drug target, especially as it was recently also identified in adult humans (3).

BAT thermogenesis is mainly mediated by the uncoupling protein 1 (UCP1) (4, 5), a transmembrane protein with six α-helical domains, which possesses a single substrate binding site a proton binding site, and several hydrophobic residues (5, 6, 7). The highest amount of this protein is located in the inner mitochondrial membrane of brown adipocytes, where, upon activation by free fatty acids, it drives the uncoupling of oxidative phosphorylation from ATP production by operating as proton carrier (8). The energy produced by the oxidation of substrates is thus released as heat giving BAT its thermogenic features. Under normal conditions, BAT is activated by the release of norepinephrine (NE) from the nerve terminals of the sympathetic nervous system (SNS). In response to NE-binding, β-adrenergic receptors facilitate second messenger signaling including the increase of intracellular cAMP concentrations and subsequently the stimulation of protein kinase A activity (Fig. 1A). While in the activation of rodent thermogenesis, the β3 adrenergic receptor plays the predominant role with a possible potentiating effect of α1 adrenergic receptors, recent findings suggested that in humans the β2 isof orm might be more relevant (9, 10). The downstream targets of the adrenergic receptors in both species, however, include for example, the hormone-sensitive lipase, which provides free fatty acids after breaking down triacylglycerols, and CREB, which activates gene expression (11).

An important regulator of BAT thermogenesis is thyroid hormone (TH), which is well-known for its effects on body temperature. Hyperthyroid patients, for instance, display an elevated body temperature and are sensitive to heat, while hypothyroid patients exhibit problems to maintain their body temperature and often feel cold (12). This review aims to summarize the current knowledge on how TH interacts with the different pathways important for maintaining body temperature with a special focus on BAT function.

Impaired thyroid hormone action interferes with correct BAT function

It is long known from early studies in thyroidectomized rats that TH is required for proper BAT function (13). On the molecular level, this comes down to two major effects, namely regulating Ucp1 transcription and controlling the sensitivity of BAT to NE (14). The latter is of particular importance, as NE increases type II deiodinase (DIO2) activity leading to an augmented intracellular conversion of the biologically less active thyroxin (T4) to the receptor-active form 3,3′,5-triiodothyronine (T3) by removing an iodine from the outer ring, thus constituting an important feed forward stimulation within the tissue. This complex synergistic interplay between the direct actions of TH in BAT, the local generation of T3 and the central stimulation of BAT by NE lies at the heart of understanding TH action in BAT thermogenesis (15).

Regulation of Ucp1 gene expression

The regulation of UCP1 content in BAT mitochondria by TH has been established long ago by studies showing lower UCP1 in rats made hypothyroid through thyroidectomy (16, 17, 18). Most remarkably, Ucp1 expression in these animals could not even be upregulated when the rats were exposed to cold, which is usually a very strong stimulus for their thermogenic response (17, 19, 20). This clearly demonstrates that TH signaling is absolutely required for Ucp1 induction and cannot be compensated by increased sympathetic stimulation of BAT. Vice versa, TH treatment by itself is also not sufficient to induce Ucp1 mRNA or protein at 23°C or 10°C ambient temperature as compared to untreated controls (21, 22). Interestingly, however, T3 treatment was found to induce Ucp1 mRNA and UCP1 protein in mice kept at thermoneutrality – a condition with minimal NE stimulation of BAT thermogenesis (22), indicating that there may be conditions under which the hormone alone can trigger Ucp1 expression. Nevertheless, due to the lack of sympathetic input in mice at 30°C, the baseline levels of Ucp1 are extremely low to begin with, and T3 treatment did not restore the UCP1 levels in those animals to the amounts present at room temperature.

Taken together, these findings underline that the individual responses of Ucp1 to NE or T3 alone are rather small and that only the combined synergistic actions of both factors can efficiently induce Ucp1 expression.

Local T3 production by DIO2 in BAT

Molecular studies revealed that Ucp1 gene expression induced by T3 occurs via TH-response elements (TRE) located in the 5′-flanking region of Ucp1 (23). For a maximal response, the saturation of thyroid hormone receptors (TRs) with T3 seems to be required, which is physiologically achieved by a bulk conversion of T4 to T3 within the BAT by DIO2 (16, 17, 19). Therefore, the intracellular production of receptor-active T3 is a key event in the thermogenic response of BAT. Upon NE-stimulation, the downstream cAMP signaling...
cascades strongly increase DIO2 activity leading to a three- to four-fold elevation of T3 concentration resulting in saturation of the nuclear TRs and the full induction of BAT thermogenesis (24). This was first observed in studies with thyroidectomized rats, which showed a normal cold response including an appropriate upregulation of Ucp1 after the application of T4, but not T3, suggesting that the local conversion of T4 by DIO2 is required for saturation of TRs (16, 19, 25). This was later confirmed in Dio2 knockout (D2KO) mice, which, despite normal T3 and even elevated T4 serum levels, exhibit an impaired BAT function and their ability to properly react to cold exposure is reduced and results in hypothermia (25, 26). Additionally, D2KO mice display impaired lipolysis and lipogenesis, which additionally affects UCP1-mediated respiratory uncoupling and thermogenesis, as activation of this protein greatly depends on the availability of free fatty acids (25, 27, 28). In line with these in vivo experiments, isolated brown adipocytes treated with the DIO2 inhibitor iopanoic acid showed diminished induction of Ucp1 gene expression (29, 30).

**Interactions of NE and TH signaling in BAT**

A key regulator of BAT thermogenesis is sympathetic signaling, which mediates the central regulation of body temperature by releasing NE from the nerve terminals. This NE release is crucially required for the TH effects on BAT, as it provides for example, the intracellular DIO2 upregulation needed for full activation of TRs. Consequently, TH alone cannot trigger Ucp1 upregulation in hypothroid rats in the absence of sympathetic stimulation (17). On the molecular level, it is assumed that the increase of intracellular CAMP levels upon NE-binding is required to active nuclear CREB (Fig. 1A), which in turn allows TREs to drive the upregulation of Ucp1 (31).
Vice versa, a thermogenic response to NE cannot be sufficiently induced in thyroidectomized rats even if sufficient UCP1 protein is available, suggesting that TH also affects other mechanisms potentially including intracellular UCP1 translocation or activation by free fatty acids (32). Likewise, it was observed in cultured rat brown adipocytes that the response to different NE stimulations depends on the availability of TH (15). These findings clearly suggest that TH somehow also modulates the thermogenic response of BAT to NE, presumably by affecting the adrenergic signaling pathway downstream of the respective receptors. It is, however, still controversial to date in how far TH also directly affects NE-turnover in BAT upstream of the receptor. Some studies did not find any change in hyper- or hypothyroid rats (33), while others showed increased NE during hypothyroid conditions (34). This would be in line with the reasoning that the tissue aims to partly compensate for missing circulating TH by upregulating DIO2 for increased local conversion of T4 to T3 (35, 36).

Divergent pathways for NE sensitivity and Ucp1 expression

Interestingly, thyroidectomized rats rescued with twice the physiological replacement dose of T3 showed a normal thermal response to NE infusion, despite the fact that their UCP1 protein content had not been normalized (32). In contrast, thyroidectomized rats treated with physiological T4 replacement doses had normalized adrenergic responsiveness of BAT and normalized Ucp1 mRNA content (32, 37). While the duration of the treatment might be a crucial factor contributing to these divergent effects, the findings still suggest that serum T3 might primarily affect NE responsiveness, while the locally produced T3 by DIO2 might be more effectively and more rapidly regulating Ucp1 expression.

Unfortunately, to date little is known on the precise mechanisms that mediate TH’s effects on NE responsiveness. While some early studies showed that TH stimulates the expression of β-adrenergic receptors in BAT (34), more recent studies found the opposite, namely reduced expression of Adrb3 at 23°C (22). As this effect was also observed at 10°C and 30°C, temperatures with higher and lower sympathetic input to BAT, it seems to be a direct T3 effect independent of NE input. In line with these findings, an increased expression of Adrb3 was observed in mice and rats with hypothyroidism (34, 38). It is tempting to speculate that these changes in Adrb3 serve as a local physiological feedback mechanism to reduce BAT sensitivity to NE in hyperthyroidism, where the animals exhibit excess heat production due to an elevated metabolic rate (22, 39), or to increase NE action in hypothyroidism, a period in which the animals might need additional facultative thermogenesis.

Thyroid hormone receptors in BAT

Two main types of nuclear TRs mediate the different functions of TH, TRα1 and TRβ. Both receptors act as ligand-dependent transcription factors but display a tissue-specific distribution across the body. Although TRα1 and TRβ are never fully mutually exclusive, TRα1 is mostly located in the heart and in the brain, while TRβ isoforms play a major role in the hypothalamus-pituitary-thyroid axis, the liver, and inner ear development (40, 41). However, whether the individual contributions of each receptor isofrom in various tissues and organs rely primarily on their expression levels or if there are isofrom specific regulatory mechanisms is not yet fully elucidated. Brown adipocytes accommodate both TRα1 and TRβ nuclear receptors (42). However, it was only in the late 90s, when the TRβ-selective analog, GC-1, opened the door to identify isofrom specific functions in BAT, as the compound binds to TRβ equally well as T3 but has a ten-fold reduced affinity to TRα1. This effect was used in a study from 2001, revealing that diet-induced hypothyroid rats treated with GC-1 restored Ucp1 expression in brown adipocytes but failed to normalize the thermal response of BAT to NE (43). This TRβ isofrom dependence of Ucp1 expression in BAT was later confirmed in TRβ knockout animals (43, 44). However, more recent studies did not find altered Ucp1 expression levels and UCP1 protein in TRβ knockout mice in comparison to wildtype animals (22), suggesting that TRα1 might be able to partially compensate the loss of TRβ under specific conditions. The importance of TRα1 signaling for the adrenergic sensitivity of BAT, however, was supported by findings in several different animal models. Mice lacking all TRα gene products have a blunted response to BAT stimulation by NE (45), and TRα1 mutant mice kept at 30°C showed no response to NE at all (46); however, for the latter study it needs to be kept in mind that the overall response to NE is generally reduced at 30°C, even in wildtypes.

Effects of increased TH action on BAT

Studies on the regulation of UCP1 and BAT thermogenesis by TH were made over an extremely broad range of treatment conditions (Fig. 1B), which makes the comparison of the results quite complicated. Moreover, most experiments were...
conducted in animal models that reflected the transition from severe hypothyroidism to hyperthyroidism, which, despite being a solid model system, has little relevance in humans. Therefore, researchers also used model systems, which resembled the development of hyperthyroidism from an euthyroid starting point. These studies revealed that, in contrast to hypothyroid animals indicating an enhanced sympathetic tone in BAT, hyperthyroid animals display a reduced adrenergic responsiveness in BAT (47, 48, 49, 50). As this correlation acts in parallel with the requirement of TH for the NE-mediated induction of UCP1, the overall TH modulated thermogenic activity of BAT was thought to resemble an inverse U-shaped curve with its maximum roughly within the euthyroid range (50). This model was then refined by studies in hyperthyroid animals with moderate T4 doses (<200 µg/kg per day), which lead to an elevated body temperature after 2 weeks of T4 treatment, but no changes in interscapular BAT (iBAT) thermogenesis, revealed by IR thermography. Furthermore, UCP1 protein levels were not altered, but Ucp1 mRNA expression was decreased (21), suggesting that iBAT was on the verge of being turned down. This trend was continued in mice treated with T3, which reached a higher level of hyperthyroidism, where BAT was in fact less active in IR thermography, PET-CT scans, and lipid uptake assays (22). However, when the treatment regimen reaches high pharmacological doses of T4 (>600–1000 µg/kg per day) over a period of 4–5 weeks, this physiological downregulation of iBAT thermogenesis is overridden. Here, an increased NE response, both at thermoneutrality as well as during cold exposure, was observed (51). Likewise, increased Ucp1 mRNA content in BAT was also observed in mice with orally administered T3 (~600µg/kg per day) at room temperature (52).

Together, this can be concluded that the extent of hypo- or hyperthyroidism has a severe effect on BAT function. Mild hypothyroidism will cause an increased sympathetic tone, which can activate BAT for compensation of reduced obligatory thermogenesis and elevated heat loss; however, when hypothyroidism becomes more severe, BAT function collapses. Likewise, mild to moderate hyperthyroidism will cause a reduction in BAT activity, presumably because the overall increase in metabolism elevates obligatory thermogenesis and thereby reduces the need for facultative thermogenesis in BAT. However, when hyperthyroidism reaches excessive levels, BAT becomes activated again (Fig. 1C) – a puzzling conundrum that can probably be explained by the central actions of TH in the brain and on the body temperature set point.

Central effects of TH

Besides the direct actions of TH on peripheral organs such as liver, heart, and BAT, TH may also provoke central actions indirectly mediating peripheral alterations. A first hypothesis, the altered set point hypothesis by Nedergaard et al. in 1997, proposed that the body temperature set point is centrally regulated and influenced by TH (51). More recently, López et al. presented findings on the interplay of the SNS and BAT thermogenesis upon central intracerebroventricular administration of T3 in rats. T3 decreases levels of the AMP-activated protein kinase in the ventromedial hypothalamus (VMH), which in turn increases SNS activity stimulating BAT thermogenesis. Additionally, thermogenic markers, such as Ucp1, hormone sensitive lipase, and Dio2 are upregulated in BAT, leading to an increased thermogenic capacity of BAT as a consequence of central T3 administration (53, 54). More importantly, body temperature was also raised, an effect that was not observed in mice lacking UCP1, demonstrating that BAT is the primary driver of body temperature elevation induced by central T3 (55, 56). While this was found to be dependent on TRα1, other studies observed an important role of TRβ in the VMH in this context, although BAT function specifically was not studied in great detail (57). However, given that TRα1 is the predominant isoform in neurons (58) and the levels of hypothalamic TRβ protein are low (59), the precise contribution of TRβ to the central control of BAT thermogenesis remains yet to be established in this context.

That central T3 action is crucial for body temperature regulation was recently confirmed in a study employing MCT8 and OATP1C1 double knockout mice. These animals lack two transporters required for the uptake of TH into the brain, with the consequence that their brains are severely hypothyroid (60). Despite their peripherally elevated T3 levels, these animals did not show an elevated body temperature (61), demonstrating that this classic feature of hyperthyroidism indeed requires the central actions of TH.

However, the connection between centrally acting T3 and its effects on peripheral organs might still be more complex. Ucp1 knockout mice treated systemically with T3 or T4 did in fact show an elevated body temperature (22, 62), suggesting that the body temperature elevation in hyperthyroidism does not depend on active BAT thermogenesis. More importantly, the authors showed that T3 treated mice had elevated heat loss over the tail, suggesting that the animals actually needed to dissipate heat to maintain the body temperature, indicative of
excess heat production in the periphery, presumably the muscle (22). Interestingly, however, it was also shown that blocking TRα1 signaling exclusively in the muscle also does not interfere with the elevation of body temperature upon thyroid hormone treatment (63).

Collectively, these data suggest that T3 acts in the brain to elevate the body temperature set point. Under normal conditions, the simultaneous action of T3 in the muscle suffices to meet this new set point – and under certain conditions even beyond, as the T3 treated mice dissipated excess heat (22). However, if a thermogenic impairment occurs such as lack of UCP1 in BAT or defective TRα1 in muscle, other mechanisms seem to be activated to achieve the body temperature elevation.

Open questions

After decades of research involving TH and its regulatory role in BAT and subsequently on BAT thermogenesis, a lot has been learnt on the underlying mechanisms, ranging from actions in muscle, white adipose tissue and BAT for thermogenesis over tail heat loss to central adjustments of thermoregulation and behavior (Fig. 2). However, the complex interplay of TH with all these systems and their interactions among each other is far from being understood. Many open questions remain to be addressed in this area, especially with the connections to other systemic effects induced by TH.

What is the reason for different body weight phenotypes during hyperthyroidism in mice, rats, and humans?

A current paradox in the field is the different body weight phenotypes in rats, mice, and humans in hyperthyroidism. Hyperthyroid rats and humans are metabolically more active leading to a reduction of body weight, whereas hyperthyroid mice gain weight over time despite an elevated body temperature and higher metabolic demands (36, 62, 64, 65). Similarly, hypothyroid mice develop a leaner phenotype than euthyroid controls, although energy expenditure and overall daily activity are significantly reduced. A recent study suggested that mouse hypothyroidism triggers an upregulation of different markers involved in skeletal muscle-mediated adaptive thermogenesis, as well as the upregulation of fatty acid oxidation to compensate for the well-established cold intolerance of hypothyroid mice (64). However, it remains to be seen whether this thermogenic effect in muscle is maintained at more severe degrees of hypothyroidism, as BAT thermogenesis is also stimulated in mild hypothyroidism, or fails at more severe conditions (see earlier). To solve this conundrum more studies comparing rats and mice at different TH levels are urgently needed, especially when considering that primary brown adipocytes of both species also respond differently with regard to Ucp1 mRNA expression (66).

What are the precise contributions of the central and peripheral TH effects for BAT thermogenesis and body temperature regulation?

However, to date the most striking enigma evolves around the contribution of TH on central and peripheral actions, and in particular, the dose, treatment duration, and temperature dependency of both effects. Severe hyperthyroidism induced by high doses of T3 recruits BAT via increased UCP1 protein and enhances the thermogenic capacity of BAT (54, 55); however, low to moderate doses of T3 or T4 do not seem to have this effect on BAT thermogenesis (21, 22). Nevertheless, central TH actions are absolutely required to increase body temperature (61). Consequently, further experiments aiming at dissecting central and peripheral (direct BAT) effects in a dose- and time-dependent manner are urgently required to address this conundrum.
What do Ucp1 expression levels and UCP1 protein tell us about the thermogenic capacity of BAT?

With the discovery of UCP1 in the 1980s and the subsequent connection to BAT thermogenesis, Ucp1 expression and UCP1 protein are normally used to describe the thermogenic capacity of BAT while the responsiveness of BAT to NE is less often discussed. However, Ucp1 mRNA and sometimes even UCP1 protein abundance can be insufficient to describe the real thermogenesis (67). Particularly, in the response to TH, these parameters can often be discordant (22, 61). It is therefore highly recommended to provide physiological data, for example, IR thermography or energy expenditure also in response to NE, in addition to the molecular data. If needed, the latter could in addition to Ucp1/UCP1 include intracellular cAMP levels, histology, proteins of the oxidative phosphorylation chain, or even NE-turnover measurements (22) to provide a more complete picture on the state of BAT thermogenesis. As these data are largely missing for many of the older studies (for obvious technical reasons), reevaluation of seemingly established concepts might be necessary.

When does ‘browning’ of WAT contribute to thermogenesis or energy expenditure?

Since BAT and its thermogenic capacity is often discussed in the context of obesity and as a possible target to assist with weight loss or to lower the risk for cardiovascular diseases and diabetes type II, it is of high interest if white adipose tissue (WAT) can be triggered to gain brown fat features – a process called ‘browning’. Browning of WAT is usually associated with greater amounts of mitochondria and increased levels of Ucp1 expression and UCP1 protein, which theoretically implies an enhanced thermogenic capacity. In the context of TH research, browning was observed in several studies (34, 56, 68). However, recent data suggested that, at least with T3, browned WAT is metabolically not active (22). Nevertheless, given that a more specific activation, by only targeting TRα1, evoked metabolic benefits (68), presumably due to the lack of parallel TRα1 activation in brain and muscle, there may still be some value in targeting TH signaling in WAT.

Which peripheral tissue mediates the increase in body temperature in response to hyperthyroidism?

In the light of the recent studies, it seems plausible that TH primarily elevates body temperature by increasing the central set point as in pyrexia (22, 62), which is at least partially fueled by peripheral TH action in muscle (22, 39). However, the higher body temperature is observed in conditions with inactive BAT or browned WAT including Ucp1 knockout mice (22, 62), as well as in mice with blocked skeletal muscle TRα1 signaling (63), which suggests that neither BAT nor muscle are absolutely required for the effect but likely can compensate for each other. Interestingly, alternative and UCP1-independent mechanisms of BAT thermogenesis could play a role in this context (69, 70); however, these pathways were not upregulated in hyperthyroid mice (22). Alternatively, one could speculate that under these rather pathological conditions, other mechanisms are recruited to defend the elevated body temperature, including reduced conductance or altered behavior (39). Additional carefully designed studies will be required to dissect the individual contributions.

What are the contributions of heat dissipation via the tail for adjusting facultative thermogenesis mediated by BAT?

The vast majority of studies on TH and body temperature regulation have exclusively been looking at thermogenesis. However, a recent study has shown that TH is also important for tail artery function, which regulates heat dissipation over the tail surface (71). If tail vasoconstriction is impaired, as seen in defective TRα1 signaling, heat is lost and additional thermogenesis is initiated to defend body temperature. This clearly demonstrates that BAT activation can occur as a secondary consequence of heat conductance defect, which is not only restricted to defective vasoconstriction but could also include fur properties, breathing rate, or behavioral changes (72). Again, these contributing factors have not been taken into consideration in previous studies, which may therefore require reevaluation of established concepts.

Take home message

Despite decades of TH research that unveiled the highly regulated interactive triangle of TH action, NE sensitivity and BAT thermogenesis, several aspects in this context are still incompletely understood. Although we know that TH and NE signaling alone are insufficient to induce UCP1-mediated thermogenesis and only the synergy of both stimuli provides an adequate BAT effect, the dose- and time-dependent contributions of both factors remain to be investigated. More recent research additionally complicated the regulatory mechanisms by revealing that central actions of TH also influence BAT thermogenesis.
and moreover the body temperature set point, which in turn leads to secondary thermoregulatory adaptations in several systems. This includes, for instance, the adjustment of heat loss, for example, via vasoconstriction over the tail surface in mice, which has rarely been documented in previous studies. Consequently, further studies aiming to investigate the effects of TH on body temperature regulation are urgently required and should consider the most recent findings on centrally acting THs and their influence on tail vasoconstriction to provide a complete picture of the thermoregulatory pathways affected by TH.

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

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