The function of the thyroid gland is to produce the thyroid hormones T3 and T4, which regulate gene transcription throughout the body (1). In medical practice, the thyroid becomes an issue when its size or shape becomes abnormal or when it produces too much or too little hormone. Thus, we typically think of the thyroid with reference to the clinical states of goiter, or hyper- or hypothyroidism. But what is the physiology of the thyroid when the gland and the entire hypothalamic-pituitary-thyroid axis are intact? As first year medical students ask each year, Why exactly do we have a thyroid, at all?

The usual presentation of thyroid physiology does not stress dynamic changes in hormone levels. Unlike insulin or cortisol levels, which are widely understood to fluctuate in response to food ingestion and stress, respectively, thyroid hormones are typically thought to be maintained at a basal level of hormone that keeps the metabolic machinery humming at the proper rate. In our opinion, this static view of thyroid physiology is mistaken.

**Nutrition and thyroid hormones.** In addition to changes in thyroid hormone that occur in development, from tadpoles to mammals, thyroid hormone levels are subject to major physiologic regulation during the transition from the fed to the starved state. In the well-studied rodent model, starvation rapidly suppresses T4 and T3 levels (2, 3). The benefit of this suppression is clear: Starvation represents a severe threat to survival, and, in rodents, the capacity to survive without nutrition is measured in days. Because thyroid hormones set the basal metabolic rate, a drop in thyroid hormone levels should reduce the obligatory use of energy stores. As long as hypothyroidism does not impair the ability to obtain food, this adaptation would be expected to enhance survival. Because animals in the wild are thought to commonly experience periods of starvation, the thyroid response to starvation should be viewed as a major aspect of the regulatory biology of the thyroid gland.

The thyroid system is regulated at multiple levels, one or more of which might account for nutritional adaptation. First, thyrotropin-releasing hormone, a neuropeptide produced in the paraventricular nucleus (PVN) of the hypothalamus, controls the release of thyroid-stimulating hormone (TSH) from the pituitary. TSH acts on receptors on the thyroid to promote synthesis and release of the thyroid hormones T4 and T3. In addition, a family of deiodinases metabolize the less-active T4 to the more-active T3 or to the inactive reverse T3. In primary hypothyroidism, when T3 and T4 levels fall because of a defect within the thyroid, a 2-part compensatory system kicks in. In the PVN of the hypothalamus, TRH expression increases because of the lack of negative feedback by thyroid hormones (4). In the pituitary thyrotroph, TSH production increases due to both increased TRH production and decreased negative feedback by thyroid hormones on the genes encoding TSH subunits (5). The increased TSH serves to drive the failing thyroid and is the most sensitive test for the diagnosis of thyroid failure.

**Figure 1**

Maintenance of the thyroid axis by leptin through actions on the TRH neuron in the PVN of the hypothalamus. A sufficient level of leptin signaling is needed to maintain TRH expression in the hypothalamic PVN, which is necessary for normal production of TSH and production of thyroid hormones by the thyroid. Two mechanisms may be involved. In one, leptin regulates arcuate neurons expressing proopiomelanocortin (POMC) (induced by leptin) and AgRP (suppressed by leptin). These arcuate neurons project to TRH neurons, where they influence TRH expression by antagonistic actions of α-MSH (stimulatory) and AgRP (inhibitory) on MC4Rs. Leptin may also act directly on TRH neurons through leptin receptors on these cells. In the absence of leptin signaling, the feedback loop between T4/T3 and the hypothalamus-pituitary-thyroid system is lost. Hence, although levels of T4/T3 may be low, TRH and TSH levels remain suppressed.
Starvation appears to act, at least in part, by suppressing TRH expression in the PVN—although, interestingly, TRH continues to be expressed in the remainder of the central nervous system (CNS), which plays no role in regulating pituitary TSH production (2). TSH production falls, and simultaneously the pattern of glycosylation on newly synthesized TSH is altered so that the TSH that is produced is of reduced bioactivity (6). Thus, as a consequence of starvation, T4 and T3 levels fall, leading to central hypothyroidism.

Leptin and the regulation of thyroid function by the brain. The mechanism by which the brain orchestrates this adaptation is now becoming increasingly clear. The dominant, and perhaps sufficient, signal to the brain that suppresses TRH expression in the PVN is a drop in the level of the hormone leptin. This 16-kD hormone is expressed predominantly in adipose cells, and its absence, as in the ob/ob mouse, produces severe obesity (7). Initially viewed primarily as a hormone designed to prevent obesity, a substantial body of work now suggests that leptin also signals the switch from the fed to the starved state (3, 8). A fall in leptin acts through the hypothalamic TRH neuron to increase appetite, decrease energy expenditure, and modify neuroendocrine function in a direction that favors survival. The consequences of falling leptin include suppression of reproduction, linear growth, and the thyroid axis, as well as activation of the stress axis (3). As with the mouse gene, mutation of the human leptin receptor gene can also cause obesity with central hypogonadism and hypothyroidism (9).

As reviewed in ref. 10, much effort is now directed to understanding the precise neural circuits through which leptin brings about these effects on appetite and neuroendocrine function. With regard to thyroid activity, a crucial question is whether falling leptin levels are sensed directly by the leptin receptor (the ObRb isoform; ref. 11) found in TRH neurons, or indirectly, through one or more distinct leptin-responsive neurons that communicate with the TRH neuron. A recent study suggested that an indirect pathway might exist. Legradi et al. (12) chemically ablated the arcuate nucleus in rats and observed that starvation failed to suppress thyroid levels. Because the treatment they used, neonatal administration of monosodium glutamate, leaves the TRH neurons of the PVN intact, this finding suggests that leptin regulates input from the arcuate to the TRH neurons in the PVN (12).

The paper by Kim et al. in this issue of the JCI adds chemical specificity to this model (13). These authors have used in vivo and in vitro approaches to reveal a role for the melanocortin pathway in mediating the nutritional response of the TRH neuron to leptin. The melanocortin 4 receptor (MC4R) is stimulated by the latter but inhibited by the former (14, 15), so the pathway can be antagonized by decreasing the agonist α-MSH, by increasing the agonist AgRP, or by a loss in receptor function. Indeed, each of these events causes obesity (16, 17). The data reported here strongly suggest that the melanocortin pathway plays an important role in the regulation of the thyroid axis by leptin as well, perhaps by promoting contacts between functionally antagonistic leptin-regulated neurons in the arcuate nucleus and TRH neurons in the PVN. Kim et al. (13), demonstrate that α-MSH increases TSH levels when it is administered centrally to living rats and that it stimulates TRH release when added to hypothalamic slices. Furthermore, AgRP blocks release of TRH by antagonizing α-MSH and thereby opposing the action of leptin. A recent report identified α-MSH in nerve terminals innervating TRH neurons in the PVN and demonstrated that this hormone prevents the fasting-induced suppression of Pro-TRH gene expression (18). Thus, the central melanocortin system can regulate the thyroid axis and is well positioned to mediate the actions of leptin on the thyroid axis.

Many endocrine pathways are subject to regulation at several levels, and the suppression of thyroid function in starvation appears to be no exception. Preliminary data suggest that TRH neurons are direct targets of leptin as well, because leptin activates TRH promoter constructs in transfected cells (19). Further studies using different approaches will be needed to examine this question. However, it should be noted that AgRP, which ectopically overexpress the agouti protein, a homologue of AgRP that also antagonizes MC4R signaling, do not have hypothyroidism. This might suggest that the acute and chronic melanocortin blockade affect thyroid function differently. In any event, the precise role of the melanocortin pathway in control of the thyroid axis under a variety of physiologic and pathophysiologic circumstances remains to be determined.

Whatever the precise wiring mechanism by which it regulates the TRH neuron, leptin’s ability to orchestrate changes in the thyroid axis in rodents during the transition from the fed to the starved states is established, and this nutritional response should probably be viewed as a key but underappreciated evolutionary function of the thyroid hormone system. Starvation is less immediately threatening for humans than for mice, because humans have greater energy stores compared with their metabolic rate. Accordingly, suppression of the thyroid axis during starvation in humans occurs more slowly and is of smaller magnitude. Studies of the possible role of leptin and the central melanocortin system in regulation of the hypothalamic-pituitary-thyroid axis in humans are in their infancy. It is unknown, for example, whether the leptin and melanocortin pathways regulate the thyroid axis under physiologic states other than from starvation or in response to severe illness, another state in which the thyroid axis may be severely suppressed (20). As leptin and reagents that modify the melanocortin system become available for clinical research studies, many new insights are sure to emerge.


