The brain plays a major role in homeostatic processes ranging from control of body temperature and fat mass to blood pressure and volume. Tight regulation of the circulating glucose level is similarly crucial for survival, and since the brain relies almost exclusively on glucose as a fuel source, it seems counterintuitive to think that the brain does not also play an important role in glucose homeostasis. Based on overwhelming evidence supporting the endocrine pancreas as the primary controller of the blood glucose (BG) level, however, the notion of a key role for the brain was discounted decades ago. Yet recent findings are beginning to chip away at the foundation of the prevailing, islet-centered view of glucose homeostasis. This perceptual shift is being driven not by evidence against a role for the endocrine pancreas in glucose homeostasis, but by evidence that the endocrine pancreas is part of a larger regulatory system, the activity of which is integrated with other critical homeostatic control systems governed by the brain. Integration of glucose homeostasis with energy homeostasis and thermoregulation This countervailing narrative begins with recognition that the amount of insulin secreted in response to a glucose challenge can be dynamically regulated by both humoral and autonomic inputs. Pancreatic islets are richly innervated by both sympathetic and parasympathetic fibers, with the former capable of powerfully […]

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Integration of glucose homeostasis with energy homeostasis and thermoregulation

This countervailing narrative begins with recognition that the amount of insulin secreted in response to a glucose challenge can be dynamically regulated by both humoral and autonomic inputs. Pancreatic islets are richly innervated by both sympathetic and parasympathetic fibers, with the former capable of powerfully inhibiting glucose-stimulated insulin secretion (GSIS) and the latter having the opposite effect (1, 2). More importantly, growing evidence that physiologically important changes in both insulin secretion and tissue glucose utilization can occur in the absence of any change in the BG level suggests that pancreatic β cell function can be regulated as part of a larger system for controlling glucose homeostasis.

For an illustration of this concept, consider that across much of the planet, mammals are confronted with swings in environmental temperature on a daily basis that pose a substantial homeostatic challenge. Meeting this challenge requires activation of diverse metabolic and autonomic responses involving three distinct homeostatic systems — glucose homeostasis, energy homeostasis, and thermoregulation — that must be integrated seamlessly if body temperature, body fat stores, and BG levels are to be maintained within narrow physiological limits.

In response to cold exposure, heat production must increase in a rapid and sustained manner if hypothermia is to be avoided, and the sympathetic nervous system (SNS) plays a key role in driving this process (3, 4). Specifically, activation of SNS outflow to thermogenic tissues (e.g., brown and white adipose tissue, skeletal muscle), driven by thermoregulatory neural circuits situated in the hypothalamic preoptic area, increases heat production via a mechanism that is highly reliant on oxidation of glucose as a substrate (3, 4). Beyond preserving core temperature, two additional challenges thus confront the cold-exposed animal: (a) how to preserve energy balance in the face of markedly increased rates of energy expenditure and (b) how to preserve stable glycemia in the face of markedly increased rates of glucose utilization.

As a result of integration across these three regulatory systems, cold exposure increases food intake in a manner that — somehow — precisely offsets the increase in energy expenditure to preserve energy balance and body fat mass (5). At the same time, insulin secretion is reduced in a manner that precisely offsets the diversion of glucose into thermogenic tissues, thereby averting decreased BG levels (6). Consistent with a role for the brain in this effect, pharmacological blockade of α-adrenergic receptors rapidly reverses the cold-induced inhibition of β cell function (6), implying that reduced insulin secretion, like the thermogenic response to cold, is driven by SNS activation. Thus, the brain orchestrates highly coordinated changes across multiple homeostatic systems that collectively enable heightened thermogenic needs to be met while ensuring that body temperature, body fat mass (5), and BG levels remain virtually unchanged (ref. 6 and Figure 1).

Coordinated regulation of these homeostatic systems via a classical negative feedback loop seems improbable, since this would require temperature, fat mass, and BG level to change before adaptive responses could be mounted, and changes in these variables were not observed (5, 6). An alternative possibility is that these responses are governed by feed-forward control mechanisms that can be engaged rapidly in anticipation of future need and thereby maintain homeostasis (7). To our knowledge, the capacity for this type of regulation is unique to the brain.

Relevance to pathogenesis of diabetes and associated metabolic impairment

At least early in the development of type 2 diabetes (T2D), the BG level appears to be regulated in the usual manner, even as it rises out of the normal range (8). This type of regulatory defect is observed in both essential hypertension and obesity, in the sense that these diseases are characterized by elevated levels of blood pressure and body fat mass, respectively. What is distinctive about each of these disorders
is that although there is an increase in the level of the regulated variable — blood glucose, blood pressure, or body fat mass — the underlying homeostatic control mechanisms appear to function normally. Since these three disorders cluster together as part of the metabolic syndrome, the possibility of a shared regulatory defect can be considered.

What mechanisms drive the defense of hyperglycemia in T2D? While β cell dysfunction clearly plays a role, an important unanswered question is whether this reflects a cell-autonomous defect or is instead imposed upon β cells by the brain (analogous to the reduction in GSIS observed during cold exposure). Consistent with the latter notion is that sympathetic inhibition of insulin secretion is increased in patients with T2D (9).

Studies in mice have identified a distinct subset of neurons in the hypothalamic ventromedial nucleus (VMN) that, when activated, not only induce diabetes-range hyperglycemia, but also completely block GSIS (10). If the brain were to perceive the BG level to be lower than it truly is, it conceivably could mount responses (including GSIS inhibition) that raise the defended BG level, which serves as a readout for whether neuroglucopenia was experimentally induced neuroglucopenia. Indeed, the response to experimentally induced neuroglucopenia establishes this to be the case.

Neuroglucopenia is induced by administration of a nonmetabolizable glucose analog (e.g., 2-deoxy-d-glucose), which is transported into cells but cannot be metabolized further, thereby disrupting cellular glucose metabolism. In response, the brain rapidly raises the BG level, which serves as a readout for whether neuroglucopenia was in fact achieved (11). Moreover, the aforementioned VMN neurons are implicated as drivers of this hyperglycemic response (12). These observations collectively support a model whereby defective brain glucose sensing contributes to the pathogenesis of hyperglycemia in T2D, analogous to the effect of impaired leptin sensing in driving excessive accumulation of body fat.

The progressive nature of β cell dysfunction in T2D, culminating in overt β cell failure, would at first glance seem to challenge this model of disease pathogenesis, since it is not immediately clear how this progression might result from a defect that does not reside within the β cell itself. Despite a decades-long search, however, a cell-autonomous basis for progressive β cell failure remains to be identified. Moreover, most endocrine cell types become severely atrophic and dysfunctional if they are subjected to continuous inhibition over long time intervals. Investigation into the contribution to β cell dysfunction made by tonic inhibition arising from the brain, perhaps aggravated by worsening metabolic status (e.g., hyperglycemia and associated glucose toxicity) and/or genetic susceptibility, is a key priority for future study.

Does T2D pathogenesis involve aberrant activity of hypothalamic glucoregulatory neurocircuits, and is this capable of raising the defended level of glycemia? Although our understanding of glucoregulatory neurocircuits is in its infancy, available evidence indicates that (i) fuel-sensing neurocircuits are concentrated in the mediobasal hypothalamus (MBH) and (ii) some of these circuits are overactive in rodent models of diabetes. Among these are GABAergic neurons situated in the arcuate nucleus that express both agouti-related peptide (Agrp) and neuropeptide Y (NPY) (referred to as Agrp neurons) (7). These neurons are physiologically important regulators of both food intake and glycemia, and they are tonically inhibited by humoral signals that convey information regarding the status of either stored fuel (e.g., leptin) or fuel available for immediate use (glucose) (13, 14). Consequently, these neurons are activated by low plasma levels of either leptin or glucose, and in otherwise normal mice, this activation is sufficient to both stimulate food intake and elevate the BG level into the diabetic range, while
conversely, silencing of these neurons is sufficient to ameliorate hyperglycemia in diabetic db/db mice (13). That these neurons are activated across rodent models of diabetes (15–17) makes them an attractive candidate mediator of the defense of hyperglycemia in T2D. The contribution made by other glucoregulatory neurons (e.g., in the VMN) to diabetic hyperglycemia is under active investigation.

**Therapeutic implications**

Since Agrp neurons are activated by hypoglycemia/neuroglucopenia (18), it seems paradoxical that they should also be activated in diabetic, hyperglycemic animals, and yet this is clearly the case (15–17). To explain this paradox, we hypothesize that brain sensing of glucose and other fuels is impaired in T2D and that hypothalamic glucoregulatory neurocircuits are activated as part of a compensatory response that drives an increase in BG level (in part by inhibiting GSIS). This model of T2D pathogenesis predicts that correcting the underlying defect should normalize glycemia in diabetic animals. Notable in this regard is the sustained antidiabetic action induced by central administration of FGF1 (19–22). In rodent models of T2D, remission of hyperglycemia can be sustained for weeks or months following a single intracerebroventricular injection of FGF1. The underlying mechanism remains under active study, but instead of simply lowering the BG level, FGF1 appears to act on MBH neurocircuits to reset glycemia in the normal range. Such an effect would not seem possible unless (a) the brain plays a key role in establishing the BG level and (b) a defect in this system contributes to the pathogenesis of hyperglycemia in these animal models.

**Conclusion**

The notion that glucose homeostasis is governed primarily by the pancreas, rather than the brain, has come under increasing scrutiny in the wake of findings that simply cannot be explained by this model. Particularly noteworthy is evidence that in rodent models of T2D, BG can be restored to normal for weeks or months by targeting of brain systems controlling glucose homeostasis. Fortunately, recent advances in neuroscience offer an unprecedented ability to map and functionally characterize the relevant neurocircuits in rodent models (7). We should seize upon this opportunity to advance our understanding of how glucose homeostasis is regulated by the brain, identify the contribution made by defects in this regulatory system to the pathogenesis of T2D, determine whether such findings substantially translate to humans, and if so, investigate whether these insights offer novel approaches to more effective disease treatment.

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