Central control of brown adipose tissue thermogenesis

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Thermogenesis, the production of heat energy, is an essential component of the homeostatic repertoire to maintain body temperature during the challenge of low environmental temperature and plays a key role in elevating body temperature during the febrile response to infection. Mitochondrial oxidation in brown adipose tissue (BAT) is a significant source of neurally regulated metabolic heat production in many species from mouse to man. BAT thermogenesis is regulated by neural networks in the central nervous system which responds to feedforward afferent signals from cutaneous and core body thermoreceptors and to feedback signals from brain thermosensitive neurons to activate BAT sympathetic nerve activity. This review summarizes the research leading to a model of the feedforward reflex pathway through which environmental cold stimulates BAT thermogenesis and includes the influence on this thermoregulatory network of the pyrogenic mediator, prostaglandin E2, to increase body temperature during fever. The cold thermal afferent circuit from cutaneous thermal receptors, through second-order thermosensory neurons in the dorsal horn of the spinal cord ascends to activate neurons in the lateral parabrachial nucleus which drive GABAergic interneurons in the preoptic area (POA) to inhibit warm-sensitive, inhibitory output neurons of the POA. The resulting disinhibition of BAT thermogenesis-promoting neurons in the dorsomedial hypothalamus activates BAT sympathetic premotor neurons in the rostral ventromedial medulla, including the rostral raphe pallidus, which provide excitatory, and possibly disinhibitory, inputs to spinal sympathetic circuits to drive BAT thermogenesis. Other recently recognized central sites influencing BAT thermogenesis and energy expenditure are also described.

Keywords: brown adipose tissue, thermogenesis, thermoregulation, sympathetic nerve activity, preoptic hypothalamus, fever, rostral raphe pallidus, rostral ventromedial medulla

INTRODUCTION
Thermogenesis, the production of heat energy, is an essential component of the homeostatic repertoire to maintain body temperature during the challenge of low environmental temperature. The heat generated during pyrogen-stimulated thermogenesis in brown adipose tissue (BAT) also contributes to fever, a controlled elevation in body temperature that reduces pathogen viability and stimulates immune cell responses. Since energy consumption during thermogenesis can involve oxidation of lipid fuel molecules, regulation of thermogenesis in response to metabolic signals can also contribute to energy balance and regulation of body adipose stores. Indeed, there is increasing interest in devising pharmacological approaches to maintaining the activation of BAT energy consumption as a metabolic furnace to burn the excess calories stored in the white adipose tissue of the obese (Clapham, 2011).

Thermogenesis can occur to a greater or lesser extent in most tissues, since heat generation is a by-product of the inefficiency of mitochondrial ATP production and of ATP utilization. However, CNS thermoregulatory networks can stimulate thermogenesis in response to a cold environment, to a fall in core body temperature or to the presence of pyrogenic cytokines primarily in three tissues: BAT, skeletal muscle, and the heart. In contrast to the indirect nature of shivering thermogenesis in skeletal muscles that are normally used to produce movement and posture, “non-shivering” thermogenesis in BAT is the specific metabolic function of this tissue and is accomplished by the heat generating capacity of a significant facilitated proton leak across the extensive mitochondrial membranes of the brown adipocytes which occurs because of the high expression of uncoupling protein-1 (UCP1) in BAT mitochondria (Cannon and Nedergaard, 2004). The levels of BAT sympathetic nerve activity (SNA), of ganglion cell norepinephrine release and of β3-adrenergic receptor binding to brown adipocytes determine the level of thermogenesis in BAT by regulating both the activity of lipases, such as hormone-sensitive lipase and adipose tissue triacylglycerol lipase, providing the immediate fuel molecules for BAT mitochondria and the level of expression of BAT mitochondrial UCP1 (Cannon and Nedergaard, 2004). Recently, alternatively activated, adipose tissue macrophages have also emerged as a significant source of norepinephrine for the cold-evoked stimulation of BAT thermogenesis (Nguyen et al., 2011).

Brown adipose tissue has developed as an essential thermoregulatory effector in cold defense in rodents and other small mammals (Golozubova et al., 2006), including infant humans, whose large surface area to body mass ratio suggests that basal metabolism alone would yield insufficient heat to maintain body temperature in cold environments. Recent observations using PET to assess tissue glucose uptake (reviewed in Nedergaard et al., 2007) and...
UCP1 analysis of biopsied tissue have confirmed a remarkable amount of metabolically active, BAT in adult humans (Hany et al., 2002; Cypress et al., 2009; van Marken Lichtenbelt et al., 2009; Virtanen et al., 2009). The functional locations of BAT depots in adult humans bear an interesting similarity to those in rodents: a large BAT pad in the vicinity of the upper thorax, individual pads atop each of the paravertebral sympathetic ganglia, and in the vicinity of the adrenal gland and kidney. The stereotyped, but curious localization of BAT pads, exemplified, for instance, by those over each sympathetic ganglion, suggests that, in addition to the overall defense of core temperature in the cold, BAT may also serve as an undescended function to maintain optimal neuronal and synaptic function in specific locations during situations in which the maintenance of core body temperature is challenged. Similar to its function in smaller mammals, adult human BAT activity (assessed by glucose uptake) is highly responsive to beta-adrenergic agonists (Soderlund et al., 2007) and to environmental temperature (Christensen et al., 2006). In addition, the finding of significantly less metabolically active BAT in obese adult human (van Marken Lichtenbelt et al., 2009) has suggested a heretofore unrecognized interaction among BAT energy expenditure, overall energy homeostasis, and body weight.

This review describes our current understanding of the functional organization of the central neural pathways (Figure 1), including the core thermoregulatory network, regulating the sympathetic outflow to BAT and, in turn, BAT thermogenesis and energy expenditure.

**THERMOREGULATORY CONTROL OF BAT THERMOGENESIS**

The core thermoregulatory network in the central nervous system (Figure 1) comprises the fundamental pathways through which cutaneous and visceral cold and warm sensation and/or reductions or elevations in brain temperature elicit changes in thermoregulatory effector tissues to counter or protect against changes in the temperature of the brain and other critical organ tissues. As with other thermal effectors (reviewed in Morrison, 2011; Morrison and Blessing, 2011), the thermoregulatory control of BAT thermogenesis occurs through a circuit that includes the pathways transmitting ambient temperature signals from thermal receptors in the skin to the hypothalamic networks that receive and integrate them with brain temperature information to activate efferent pathways to the thermal effectors. Each of the synaptic integration sites in the core thermoregulatory pathway constitutes a potential site for the modulation of BAT thermogenesis by non-thermal signals.

The thermoregulatory control of BAT thermogenesis in response to environmental temperature constitutes a feedforward reflex or “central command” drive, in that the stimulus of a cold environment in contact with skin thermal receptors is not, itself, affected by the evoked thermogenic response, but rather, BAT thermogenesis is initiated to counter a “predicted” fall in body core temperature that would result from the exposure to the cold environment. In contrast, the control of BAT thermogenesis by temperature-sensitive neurons in the brain constitutes a negative feedback reflex in that the warm-sensitive, preoptic area (POA) neurons activated by the increases in brain temperature resulting from stimulated BAT thermogenesis act, in turn, to inhibit sympathetic outflow to BAT.

**CUTANEOUS THERMAL RECEPTOR AFFERENT PATHWAY**

**Cutaneous thermoreception**

The thermoregulatory system initiates defensive thermoregulatory responses in response to changes in skin temperature before they affect body core temperature. In this way, exposure to a cold environment can leave core and brain temperatures unaffected or slightly increased (Lomax et al., 1964; Bratinscsak and Palkovits, 2005). Such environmental temperature sensation is mediated through cutaneous thermoreceptors which are located in primary sensory nerve endings distributed in the skin. The molecular mechanisms of cutaneous thermoreception appear to reside in the transient receptor potential (TRP) family of cation channels. The strongest data are in support of the TRPM8 as the cutaneous cold receptor TRP channel: TRPM8 is activated by modest cooling (<27°C; McKemy et al., 2002; Peier et al., 2002), TRPM8-deficient nerve fibers show profound loss of cold sensitivity and TRPM8-deficient mice exhibit a reduced ability to avoid innocuous cold temperatures (Bautista et al., 2007; Colburn et al., 2007; Dhaka et al., 2007). In addition, TRPM8 is activated by menthol (McKemy et al., 2002; Peier et al., 2002) which evokes warm-seeking behavior as well as cold-defensive, physiological responses. TRPV3 and TRPV4 are warm-sensitive TRP channels that are activated by innocuous warm temperatures and are expressed in keratinocytes in skin epidermis. Compared to wild-type mice, mice lacking either TRPV3 or TRPV4 show altered behaviors in discriminating innocuous warm temperatures (Lee et al., 2005; Moqrich et al., 2005).

Intriguing effects on body temperature control and BAT thermogenesis have been reported for agonists and antagonists of TRPV1, a TRP channel that can be activated, in vitro, by a noxious range of heat (>43°C), by protons (pH ≤ 5.9) or by capsaicin (Tominaga et al., 1998). Peripheral or central administration of capsaicin induces hyperthermia (Jancso-Gabor et al., 1970; Hori, 1984), an effect beneficial to people living in hot climates, but is ineffective in TRPV1-deficient mice (Caterina et al., 2000). Furthermore, administration of potent TRPV1 antagonists induces hyperthermia (Gavva et al., 2007) by both increasing metabolism and reducing heat loss from the body surface, but not by evoking warm-seeking behavior (Steiner et al., 2007) and this hyperthermic effect is likely exerted by antagonizing TRPV1 located within abdominal viscera (Steiner et al., 2007; Romanovsky et al., 2009). Therefore, tonic activation of peripheral TRPV1, effected by non-thermal stimuli at body temperatures below the threshold for TRPV1 activation, could provide afferent signals to lower body temperature (Romanovsky et al., 2009), however, TRPV1-deficient mice exhibit no obvious deficit in body temperature control (Szelenyi et al., 2004).

In addition to cutaneous thermoreception, thermoreceptive mechanisms exist in body core structures including the brain (e.g., neurons in the POA), spinal cord, and abdomen. The afferent fibers from cold and warm receptors in the abdominal viscera are included among the splanchnic and vagus nerve afferent fibers and their responses to temperature changes are similar to those of cutaneous thermoreceptors (Riedel, 1976; Gupta et al., 1979). Temperature changes in the spinal cord can affect the activity of thermoregulatory neurons in more rostral areas of the brain.
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(Guieu and Hardy, 1970). TRP channels that are located in the central endings of primary somatosensory fibers in the spinal dorsal horn (Tominaga et al., 1998; Bautista et al., 2007) may sense spinal temperature and could underlie an integration of spinal thermal signals with cutaneous thermal signals at the spinal cord level. Thus, rather than responding directly to changes in environmental temperature, core body thermosensation could play a role (a) in setting the basal tone of thermoregulatory effector efferents including BAT thermogenesis, (b) in enhancing thermoregulatory responses in situations of extreme thermal environments when the feedforward thermoregulatory responses driven by changes in skin temperature have proven inadequate to prevent changes in brain or body core temperature, and (c) in responding to challenges to thermal homeostasis involving shifts in internal body temperature brought about by changes in metabolism (e.g., exercise, hypoglycemia, hypoxia, etc.) or by changes in internal temperature (e.g., intake of cold fluids, hemorrhage, etc.).

**Dorsal horn**

Primary thermal somatosensory fibers deliver thermal information to lamina I neurons in the spinal (or trigeminal) dorsal horn (Craig, 2002; Figure 1). Craig and colleagues have described thermoreceptor-specific cells responding linearly to graded, innocuous cooling or warming stimuli, and not being activated further in the noxious temperature range (Andrew and Craig, 2001; Craig et al., 2001). The spinothalamocortical pathway, in which second-order thermosensory neurons in lamina I ascend to synapse on thalamic neurons that, in turn, project to the primary somatosensory cortex, is responsible for conscious...
Within the neural circuits regulating body temperature, the hypothalamus, including the POA, occupies a pivotal position between the cutaneous sensation of ambient temperature and the motor pathways controlling the engagement of thermal effectors (Figure 1). Befitting its function as a central integrator of the many dimensions of homeostatic space, the hypothalamus is composed of several interconnected populations of neurons, receives a variety of signals relating to behavioral and emotional state, as well as the condition of the body and the interstitial fluid, and has outputs influencing emotional, behavioral, somatic, and autonomic responses. Control of body temperature is but one of a myriad of interrelated homeostatic functions embedded in the hypothalamic matrix. Despite the anatomical and neurochemical complexity of this brain region, and the many factors that can influence body temperature regulation, considerable progress has been made in understanding the functional organization of the hypothalamic network that controls BAT thermogenesis.

**Lateral parabrachial nucleus**

Neurons in the external lateral subnucleus (LPBel) of the LPB and projecting to the median subnucleus (MnPO) of the POA are activated following cold exposure (Bratinscsak and Palkovits, 2004; Nakamura and Morrison, 2008b), while those in the dorsal subnucleus (LPBd) are activated in response to skin warming (Bratinscsak and Palkovits, 2004; Nakamura and Morrison, 2010). The discharge rate of single, MnPO-projecting LPBel neurons recorded in vivo increased markedly in response to skin cooling in a manner paralleling the skin cooling-evoked increases in BAT SNA (Nakamura and Morrison, 2008b). In contrast, single, MnPO-projecting LPBd neurons were excited by skin warming in parallel with the simultaneous inhibition of BAT SNA (Nakamura and Morrison, 2010). The critical role of LPB neurons in transmitting cutaneous, and possibly visceral, thermal sensory information to the hypothalamus to drive BAT thermogenic responses is demonstrated by the elimination of BAT responses to alterations in skin temperature following experimental inactivation of local neurons or blockade of local glutamate receptors in the LPB (Kobayashi and Osaka, 2003; Nakamura and Morrison, 2008b). Similarly, glutamate or other stimulations of LPBel or LPBd neurons can evoke BAT sympathetic and thermogenic responses that parallel those evoked during decreases or increases, respectively, in skin temperature (Nakamura and Morrison, 2008b, 2010). Thus, both cold and warm cutaneous thermostensive signals that are transmitted from spinal dorsal horn or trigeminal neurons to the POA by separate populations of LPB neurons (Figure 1) are essential for eliciting rapid responses in BAT thermogenesis to defend body temperature from a variety of thermal challenges. Although nociceptive inputs play only a minor role (Nakamura and Morrison, 2008b), we do not know what other signals are integrated with cutaneous cold affenter inputs to LPBel neurons in the feedforward pathway contributing to drive BAT thermogenesis during environmental cold challenges.

**Hypothalamic mechanisms in the thermoregulatory control of BAT thermogenesis**

Within the neural circuits regulating body temperature, the hypothalamus, including the POA, occupies a pivotal position
FIGURE 2 | Inhibition of neurons in the median preoptic nucleus (MnPO) or blockade of GABA\textsubscript{A} receptors in the medial preoptic nucleus (MPO) prevents skin cooling-evoked BAT thermogenesis. (A) Before and after injection of saline (SAL) vehicle into the MnPO [inset: typical injection site (arrow) in the MnPO; 3v, third ventricle; ox, optic chiasm; ac, anterior commissure], episodes of skin cooling evoke increases in BAT sympathetic nerve activity (SNA), BAT temperature (TBAT), expired CO\textsubscript{2} (Exp CO\textsubscript{2}), and heart rate (HR), with no change in arterial pressure (AP). Following nanoinjection of the inhibitory transmitter, glycine (GLY), into the MnPO, skin cooling no longer increases these thermoregulatory parameters. Modified with permission from Nakamura and Morrison (2008a). (B) The skin cooling-evoked increases in thermoregulatory parameters, including BAT SNA and TBAT, are unaffected by nanoinjection of saline vehicle into the MPO, but these increases are reversed by blockade of GABA\textsubscript{A} receptors in MPO with nanoinjection of bicuculline (BIC). Modified with permission from Nakamura and Morrison (2007).

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The MPO or LPO, are activated (express Fos protein) in response to putative warm-sensitive neurons; Nakayama et al., 1961, 1963). Increased their discharge during local hypothalamic warming (i.e., spontaneous discharge at thermoneutral temperatures that are consistent with a model (Figure 1) in which warm-sensitive POA neurons that are tonically active at thermoneutral temperatures, integrate cutaneous and local thermal information, and send inhibitory projections from the MPO to suppress BAT thermogenesis. As ambient temperature is reduced below the thermoneutral zone, such as to a normal “room” temperature of ~23°C, the discharge rate of warm-sensitive POA neurons is reduced, thereby disinhibiting BAT sympathoexcitatory neurons in more caudal brain sites such as the dorsomedial hypothalamic (DMH) or the rostral raphe pallidus (rRPa; see below) whose increased activity then drives BAT thermogenesis (Figure 1). Thus, the firing rates of warm-sensitive projection neurons in the MPO, potentially the principal neurophysiological substrate underlying the thermoregulatory “balance point” (Romanovsky, 2004), are determined by both thermosensory afferent signals from the skin and by the effect of local brain temperature. Other inputs to POA warm-sensitive neurons, such as those postulated from “temperature-insensitive” neurons (Boulant, 2006b), are expected to influence the level of BAT thermogenesis by affecting the basal discharge level of warm-sensitive projection neurons in the MPO or their sensitivity to local and cutaneous thermal signals.

**Warm-sensitive neurons in POA integrate cutaneous and core thermal signals to provide inhibitory regulation of BAT thermogenesis**

The conceptual foundation of our current understanding of the role of the hypothalamus in normal body temperature regulation and in the elevated body temperature during fever is the existence of a class of hypothalamic neurons which have intrinsic temperature sensitivity: in the absence of synaptic inputs, their discharge frequency increases as the temperature of their local environment increases. The neurophysiological mechanism underlying the thermosensitivity of warm-sensitive neurons in the POA is thought to reside in a warming-dependent facilitation of the rate of rise of a depolarizing prepotential, due to an heat-induced increase in the inactivation rate of an A-type potassium current, which shortens the intervals between action potentials and thereby increases their firing rates (Boulant, 2006a). Warm-sensitivity could also arise from a heat-induced membrane depolarization that allows warm-sensitive neurons to reach their discharge threshold potential and then determines their discharge frequency (Kobayashi et al., 2006), although the channel mediating such a depolarization has not been identified. Although neurons whose spontaneous discharge frequency is altered by changing the temperature of their local environment exist throughout the CNS, those in the POA and anterior hypothalamus have been most intensely studied because thermoregulatory responses, perhaps with the exception of certain thermoregulatory behaviors (Almeida et al., 2006), are dependent on the integrity of POA neurons. The preeminent importance of central warm-sensitive neurons for the maintenance of normal body temperature can also be appreciated from the relative position of mammalian resting body temperatures well above the freezing point of water, but only a few degrees below the temperature at which proteins begin irreversible denaturation (Romanovsky, 2007).

Initial, in vivo recordings in the POA identified neurons with spontaneous discharge at thermoneutral temperatures that increased their discharge during local hypothalamic warming (i.e., putative warm-sensitive neurons; Nakayama et al., 1961, 1963). The POA contains warm-sensitive neurons whose tonic discharge is also reduced by skin cooling and whose thermosensitivity to preoptic temperature is increased when the skin is cooled (Boulant and Hardy, 1974). In subsequent recordings in the POA in hypothalamic slices, the majority of thermosensitive neurons were warm-sensitive (Boulant and Dean, 1986) and the majority of these were GABAergic (Lundius et al., 2010). Further, either skin cooling or direct cooling of the local environment of POA neurons evokes sympathetic thermogenesis in BAT (Imai-Matsumura et al., 1984) and transections immediately caudal to the POA elicit large increases in BAT temperature (Chen et al., 1998). These findings are consistent with a model (Figure 1) in which warm-sensitive POA neurons that are tonically active at thermoneutral temperatures, integrate cutaneous and local thermal information, and send inhibitory projections from the MPO to suppress BAT thermogenesis. As ambient temperature is reduced below the thermoneutral zone, such as to a normal “room” temperature of ~23°C, the discharge rate of warm-sensitive POA neurons is reduced, thereby disinhibiting BAT sympathoexcitatory neurons in more caudal brain sites such as the dorsomedial hypothalamic (DMH) or the rostral raphe pallidus (rRPa; see below) whose increased activity then drives BAT thermogenesis (Figure 1). Thus, the firing rates of warm-sensitive projection neurons in the MPO, potentially the principal neurophysiological substrate underlying the thermoregulatory “balance point” (Romanovsky, 2004), are determined by both thermosensory afferent signals from the skin and by the effect of local brain temperature. Other inputs to POA warm-sensitive neurons, such as those postulated from “temperature-insensitive” neurons (Boulant, 2006b), are expected to influence the level of BAT thermogenesis by affecting the basal discharge level of warm-sensitive projection neurons in the MPO or their sensitivity to local and cutaneous thermal signals.

**The dorsomedial hypothalamus contains BAT sympathoexcitatory neurons**

The observation that transection of the neuraxis immediately caudal to the POA increases BAT SNA and BAT thermogenesis (Chen et al., 1998) suggests that the efferent output of the POA is inhibitory to BAT thermogenesis. However, transections made just caudal to the hypothalamus do not increase basal levels of BAT thermogenesis in normothermic animals (Rothwell et al., 1983) and, in fact, reverse PGE2-evoked increases in BAT SNA and BAT thermogenesis (Morrison et al., 2004; Rathner and Morrison, 2006). Thus, although a long inhibitory pathway from POA warm-sensitive neurons to medullary BAT sympathetic premotor neurons may contribute to the regulation of BAT thermogenesis, a source of excitatory drive to BAT thermogenesis must exist between the POA and the rostral midbrain.

Convincing data support a role for neurons in the DMH/dorsal hypothalamic area (DMH/DA) in the control of BAT thermogenesis (Dimicco and Zaretsky, 2007), potentially as BAT sympathoexcitatory neurons antecedent to medullary BAT sympathetic premotor neurons in rRPa (Figure 1). Administration of endotoxin or cold exposure increases the expression of Fos in neurons of the DMH/DA (Elmqquist et al., 1996; Cano et al., 2003; Sarkar et al., 2007). Disinhibition of DMH/DA neurons increases BAT SNA (Cao et al., 2004; Figure 3A) and thermogenesis (Zaretskaia et al., 2002), suggesting a tonic GABAergic inhibitory input to BAT thermogenesis-promoting neurons in the DMH/DA (Figure 1). This tonic GABAergic input to neurons in the DMH/DA may originate in the POA since POA-derived GABAergic axon swellings make close appositions with DMH neurons, including those that project to the rRPa (Nakamura et al., 2005). In addition, inhibition of neurons in the DMH/DA or blockade of local glutamate receptors in the DMH/DA reverses febrile and cold-evoked excitations
of BAT SNA (Figure 3B) and stimulations of BAT thermogenesis (Zaretskaia et al., 2003; Madden and Morrison, 2004; Morrison et al., 2004; Nakamura et al., 2005; Nakamura and Morrison, 2007). Thus, the glutamate-driven activity of neurons in the DMH/DA is required for cold-evoked and febrile activations of BAT thermogenesis, following their disinhibition by a reduction in their GABAergic inhibitory inputs from POA (Figure 1). Although the source of the required excitatory input to DMH/DA neurons accounting for their activity following disinhibitory stimuli has yet to be determined, it could arise from cold-sensitive or thermally insensitive neurons in the POA (Boulant, 2006b) or perhaps from an input from the periaqueductal gray (PAG; de Menezes et al., 2009).

Neurons in the DMH/DA do not project directly to BAT sympathetic preganglionic neurons (SPNs), but their monosynaptic projection to the rostral ventromedial medulla (Hermann et al., 1997; Samuels et al., 2002; Nakamura et al., 2005; Yoshida et al., 2009), including the principal site of BAT sympathetic premotor neurons in the rRPa (see below), has been implicated in mediating the effects of DMH/DA neurons on BAT thermogenesis. Glutamate receptor activation in the rRPa is necessary for the increase in BAT SNA and BAT thermogenesis evoked by disinhibition of neurons in the DMH/DA (Cao and Morrison, 2006). Neurons in the DMH/DA that are retrogradely labeled from tracer injections into the rRPa express Fos in response to BAT thermogenic stimuli such as endotoxin, cold exposure or stress (Sarkar et al., 2007; Yoshida et al., 2009), and some DMH/DA neurons that project to the rRPa receive close GABAergic appositions from neurons in the MPO (Nakamura et al., 2005).

In addition to the evidence for a direct monosynaptic pathway from the DMH/DA to the rRPa, both anatomical and physiological evidence suggest a role for neurons in the PAG in determining the level of BAT thermogenesis, potentially by influencing the output from the DMH. Some DMH/DA neurons projecting to the caudal PAG (cPAG) express Fos in response to cold exposure (Yoshida et al., 2005) and some neurons in the cPAG are multisynaptically connected to BAT (Cano et al., 2003), presumably including those that project directly to the raphe (Hermann et al., 1997). Neurons in the cPAG express Fos in response to cold (Cano et al., 2003), although these may not project to the rRPa (Yoshida et al., 2009). Excitation of neurons in cPAG increases BAT temperature, without a concomitant increase in core temperature (Chen et al., 2002), while similar excitation of neurons in the lateral and dorsolateral PAG (dl/lPAG) of conscious rats increases core temperature, the latter being dependent on activity within the DMH (de Menezes et al., 2009). In contrast, in anesthetized and paralyzed rats, skin cooling-evoked stimulation of BAT thermogenesis was unaffected by muscimol injections into the cPAG (Nakamura and Morrison, 2007). The area of the rostral ventromedial PAG (rvmPAG) contains neurons with an inhibitory effect on BAT thermogenesis that are capable of reversing the BAT thermogenesis evoked by PGE2 injections in to POA or by disinhibition of neurons in DMH/DA (Rathner and Morrison, 2006). Clearly, there is a need for further investigation of the pathways transmitting the sympathetic drive for BAT thermogenesis from the hypothalamus to medullary BAT premotor neurons, and of the roles of various regions of the PAG in regulating BAT thermogenesis.

THE ROSTRAL RAPHE PALLIDUS CONTAINS BAT SYMPATHETIC PREMOTOR NEURONS

Within the hierarchical organization of the central thermoregulatory network, neurons in the rostral ventromedial medulla, centered in the rRPa and extending into nearby raphe magnus nucleus and over the pyramids to the parapyramidal area (PaPy; Bamshad et al., 1999; Oldfield et al., 2002; Cano et al., 2003; Yoshida et al., 2003), play key roles as BAT sympathetic premotor neurons – providing an essential excitatory drive to BAT SPNs in the thoracolumbar spinal cord, which, in turn, excite sympathetic...
ganglion cells innervating the BAT pads (Figure 1). A comparison of the localization of Fos induced by cold exposure which activates BAT thermogenesis with the locations of retrogradely labeled neurons following virus inoculations of BAT provided function-based evidence that the rRPa (Figure 4A) and the ventromedial parvicellular subdivision of the paraventricular hypothalamic (PVH) nucleus are the two potential premotor populations having principal roles in mediating the descending regulation of the spinal sympathetic circuit controlling BAT thermogenesis during cold defense (Cano et al., 2003). Further functional studies have clearly identified the preeminent role of BAT sympathetic premotor neurons in rRPa in the cold-defense activation of BAT thermogenesis; however, the role of the Fos-expressing neurons in PVH remains unknown.

Brown adipose tissue sympathetic premotor neurons in the rRPa receive a potent glutamatergic excitation, as well as GABAergic inhibitory inputs, with the latter predominating under warm conditions to reduce BAT thermogenesis. Relief of this tonically active, GABAergic inhibition as well as an increase in glutamate-mediated excitation, including that from the DMH (Cao and Morrison, 2006), contributes to the cold-evoked and febrile increases in BAT premotor neuronal discharge that drives BAT SNA and BAT heat production. NANOinjections into rRPa of agonists for either NMDA or non-NMDA glutamate receptors evoke brief, but intense activations of BAT SNA (Madden and Morrison, 2003), indicating that neurons in rRPa capable of increasing the sympathetic drive to BAT express NMDA and non-NMDA subtypes of glutamate receptors. That nanoinjections of bicuculline into the rRPa evoke intense activations of BAT SNA (Figure 4B) and BAT energy expenditure (Morrison et al., 1999) that are reduced by glutamate receptor antagonists suggests the existence either of an ongoing or bicuculline-activated excitatory input to rRPa BAT sympathetic premotor neurons.

Conversely, inhibition of neuronal activity or blockade of glutamate receptors in the rRPa reverses the increases in BAT SNA and BAT thermogenesis elicited by a variety of thermogenic stimuli, including skin cooling (Figure 4C) and fever (Nakamura et al., 2002; Madden and Morrison, 2003; Morrison, 2003; Nakamura and Morrison, 2007; Ootsuka et al., 2008). Inhibition of rostral ventromedial medullary neurons produces dramatic falls in body temperature in conscious rats (Zaretsky et al., 2003), consistent with an active contribution of BAT sympathetic premotor neurons in the rRPa and BAT thermogenesis to the maintenance of core temperature in a room temperature environment. Other thermogenic stimuli whose activation of BAT thermogenesis is reversed or prevented by inhibition of neural activity in the rRPa include disinhibition of neurons in the DMH (Cao et al., 2004) or in the lateral hypothalamus (Cerri and Morrison, 2005); activation of central mu-opioid receptors (Cao and Morrison, 2005), central melanocortin receptors (Fan et al., 2007) or preoptic CRF receptors (Cerri and Morrison, 2006), and systemic administration of the adipose tissue hormone, leptin (Morrison, 2004). Thus, the rRPa and PaPy regions of the ventromedial medulla contain the principal populations of BAT sympathetic premotor neurons that provide the final common medullospinal pathway (Figure 1) for the BAT sympathoexcitatory drive to the spinal network controlling BAT SNA and that are both necessary and sufficient for the
BAT thermogenic responses to thermoregulatory (Figure 1) and febrile stimuli and to a variety of neurochemical mediators that influence body temperature.

That vesicular glutamate transporter 3 (VGLUT3)-expressing and serotonin-containing neurons in the rostral ventromedial medulla are functionally related to the activation of cold-defensive, BAT thermogenesis is indicated by the findings that a significant percentage of VGLUT3-containing neurons in the rRPa express Fos in response to cold exposure or intracerebroventricular (ICV) PGE$_2$ (Nakamura et al., 2004a) and that physiologically identified serotonergic neurons in the rRPa increase their firing rate in response to PGE$_2$ administration or cold exposure (Martin-Cora et al., 2000).

Rats maintained with a brainstem transection just rostral to the superior colliculus (Nautiyal et al., 2008) or with bilateral cuts just caudal to the POA (Blatteis and Banet, 1986) can mount relatively normal cold-defense responses, leading to the conclusion that the POA is not essential for the integration of autonomic thermoregulatory responses in the rat and that circuits caudal to the transection are sufficient to convey cutaneous thermal information to the premotor neurons in the rRPa necessary to evoke cold-defense responses. Why the pathways proposed by these investigators to explain these effects in transected rats are ineffective in intact rats following neuronal inhibition of hypothalamic sites in intact animals (Zaretskaia et al., 2003; Madden and Morrison, 2004; Nakamura and Morrison, 2007, 2008a) remains unexplained. These results may point to an effect of anesthesia or to a response to the transection injury, since they could not be repeated acutely following nearly identical transections rostral to the colliculi in anesthetized rats (Osaka, 2004). Thermally sensitive neurons have been recognized in several sites caudal to the POA and these neurons, rather than cutaneous thermal receptors, may be engaged in eliciting cold-defense responses in rats with transections caudal to the POA. Overall, while these occasional data derived from transected preparations are curious, their relevance to normal thermoregulatory mechanisms in intact animals remains unknown.

**SPINAL SYMPATHETIC MECHANISMS INFLUENCING BAT THERMOGENESIS**

The discharge of BAT SPNs that determines the level of BAT SNA and BAT thermogenesis, as well as the rhythmic bursting characteristic of BAT SNA, is governed by their supraspinal and segmental inputs as well as those to the network of spinal interneurons that influence BAT SPN excitability. A significant fraction of the BAT sympathetic premotor neurons in rRPa, identified following viral retrograde tracing, are glutamatergic and/or serotonergic neurons. Spinally projecting neurons in the rRPa region can contain phenotypic markers for (a) the VGLUT3, potentially indicative of glutamatergic neurons (Nakamura et al., 2004a; Stornetta et al., 2005); (b) serotonin (5-HT) or tryptophan hydroxylase, a synthetic enzyme for 5-HT (Cano et al., 2003; Nakamura et al., 2004a; Stornetta et al., 2005), and (c) glutamic acid decarboxylase-67 (GAD-67), a marker for GABAergic neurons (Stornetta et al., 2005). Consistent with these findings, 5-HT-containing (Bacon and Smith, 1988; Vera et al., 1990) and VGLUT3-containing terminals synapse on SPNs (Stornetta et al., 2005) or make close appositions with SPN dendrites (Nakamura et al., 2004a,b). In addition, IML-projecting dendrites (Nakamura et al., 2004a; Madden and Morrison, 2006) and blockade of glutamate receptors in the upper thoracic IML suppresses the increase in BAT thermogenesis evoked by bicuculline injection into rRPa (Nakamura et al., 2004a). Putative serotonergic neurons in the rRPa increase their firing rate in response to cold (Martin-Cora et al., 2000; Nason and Mason, 2006) or PGE$_2$ administration (Nason and Mason, 2006) and blockade of spinal serotonin receptors reverses the cold-evoked activation of BAT SNA (Madden and Morrison, 2010). Serotonin in the IML can activate BAT SNA and BAT thermogenesis and potentiates the BAT SNA response to NMDA injections into the IML (Madden and Morrison, 2006), such that prior application of serotonin into the IML allows a subsequent subthreshold dose of NMDA to evoke a marked increase in BAT SNA (Madden and Morrison, 2006). The significant role of serotonin-containing neurons in normal cold-defense responses is also supported by the finding that mice that lack almost all central serotonergic neurons show blunted BAT thermogenesis during cold exposure (Hodges et al., 2008). The mechanisms of the interaction between glutamatergic and serotonergic neurotransmission in the IML remain to be elucidated.

Viral inoculations of interscapular BAT (Can et al., 2003) consistently label a population of spinal interneurons in the vicinity of the IML. Spinal GABAergic interneurons would appear to be among this population since they influence the discharge of SPNs (Deuchars et al., 2005). That such interneurons could receive inputs from the BAT premotor area in the rostral ventromedial medulla is suggested by the demonstration that VGLUT3- and GAD-67-containing terminals synapse on GABAergic neurons in the IML (Stornetta et al., 2005), providing a potential anatomical substrate for a pathway that increases the activity of SPNs through disinhibition. 5-HT-containing terminals form close appositions with GABAergic interneurons in the central autonomic area (Conte et al., 2007) and serotonergic inputs activating 5-HT$_{1A}$ receptors on GABAergic neurons in the IML have been postulated to explain the potentiation of excitatory inputs to BAT SPNs by exogenously applied 5-HT (Madden and Morrison, 2006, 2008a).

Spinal catecholamine release may also modulate the activity of BAT SPNs, considering the excitatory and inhibitory effects of catecholamines on functionally unidentified SPNs (Coote et al., 1981; Miyazaki et al., 1989) and the observation of a dense dopamine beta hydroxylase innervation of SPNs that are synaptically connected to BAT (Can et al., 2003). Substance P terminals also innervate SPN's (Vera et al., 1990), substance P excites the majority of SPNs (Cammack and Logan, 1996), and intrathecal substance P affects thermoregulation (Dib, 1987), although the latter may be due to an effect on thermal afferent processing in the dorsal horn.
Fever is a defended elevation in body temperature that plays a significant role in the acute phase reaction stimulated by endogenous pyrogens released during infection or inflammation. PGE2, which is synthesized in the brain vasculature and in peripheral tissues in response to immune signals (Elmquist et al., 1997; Matsumura et al., 1998; Yamagata et al., 2001), is a powerful endogenous pyrogenic mediator that binds to EP3 receptors in the POA, particularly the MPO and MnPO (Scammell et al., 1996; Nakamura et al., 2000, 2002; Lazarus et al., 2007), to activate BAT thermogenesis in concert with other thermoregulatory effectors to produce a sustained increase in core body temperature. Intravenous PGE2 is also effective in eliciting the BAT thermogenic component of the febrile response (Ootsuka et al., 2008).

Although we have only a rudimentary understanding of the microcircuity of the POA thermoregulatory network, we have proposed (Nakamura et al., 2004a, 2005) a model, incorporated in Figure 1, for the essential POA mechanisms through which application of PGE2 into the POA elicits the BAT thermogenic component of the febrile response. As described above for cold-defense responses, under normal conditions, EP3 receptor-expressing POA neurons, potentially including the population of warm-sensitive POA neurons that control BAT thermogenesis, maintain a tonic GABAAergic inhibition of sympathoexcitatory, BAT thermogenesis-promoting neurons in the DMH, and potentially in the rRPa. During infection, PGE2, produced locally and/or systemically, binds to these EP3 receptors, reducing the activity of BAT-controlling, warm-sensitive neurons in POA, which, in turn, leads to disinhibition of BAT thermogenesis-promoting neurons in DMD/DA and subsequently in rRPa, to drive BAT sympathetic outflow and BAT heat production contributing to an elevated core body temperature. Although the resulting increase in the local temperature of the POA would normally elicit more rapidly rising prepotentials in POA warm-sensitive neurons to inhibit BAT activation, this is offset by the membrane hyperpolarization elicited by EP3 receptor occupancy, thereby allowing a sustained activation of BAT thermogenesis and a maintained fever.

Several experimental findings are consistent with such a model. Binding of PGE2 to EP3 receptors can inhibit neuronal activity by coupling to inhibitory GTP-binding proteins (Narumiya et al., 1999) and the tonic activity of most warm-sensitive neurons in the POA is inhibited by the E-series of PGs (Schoener and Wang, 1976; Ranels and Griffin, 2003). A population of EP3 receptor-expressing POA neurons multisynaptically innervates BAT (Yoshida et al., 2003) and a subpopulation of EP3 receptor-expressing neurons in the POA projects directly to the DMD (Nakamura et al., 2002, 2005). Interestingly, this subpopulation is separate from the subpopulation of EP3 receptor-expressing POA neurons projecting to the rRPa, although a population of POA neurons that do not express EP3 receptors does send bifurcating axonal projections to both DMD and the rRPa (Nakamura et al., 2009). The majority of EP3 receptor-expressing POA neurons are GABAergic (Nakamura et al., 2002), as are warm-sensitive neurons in POA (Lundius et al., 2010). Antagonizing GABA_A receptors in the DMH evokes a fever-like stimulation of BAT thermogenesis (Morrison, 1999; Morrison et al., 1999; Zaretskai et al., 2002; Gao et al., 2004) that is also reversed by blockade of glutamate receptors in the rRPa (Cao and Morrison, 2006). Inhibition of POA neurons with a muscimol nanoinjection elicits hyperthermic, cardiovascular, and neuroendocrine responses similar to those evoked by a PGE2 nanoinjection into the same site (Zaretsky et al., 2006). In addition, ICV PGE2 application reduces cAMP level in the POA and ICV administration of an inhibitor of phosphodiesterase, a degradation enzyme for cAMP, blunts fever evoked by intra-POA PGE2 application (Steiner et al., 2002). A marked increase in core temperature and a tachycardia can be elicited by injection of PGE2 into the paraventricular nucleus of the hypothalamus (PVH) or into the parotine parabrachial nucleus (PBN; Skibicka et al., 2011), although whether neurons in these two sites play a role in fever generation remains to be determined. In this regard, the demonstration that elimination of EP3-R selectively in the POA is sufficient to prevent LPS fever (Lazarus et al., 2007) indicates that EP3-R in PVH or in PBN is not sufficient to support a febrile response or possibly that PGE2 is not increased in these regions during a febrile stimulus such as LPS.

**OREXIN IN rRPa INCREASES BAT THERMOGENESIS**

Orexins (hypocretins; de Lecea et al., 1998; Sakurai et al., 1998) are synthesized exclusively in widely projecting (Peyron et al., 1998) neurons in the perifornical area of the lateral hypothalamus (PeF-LH). Orexin neurons influence a variety of functions, including the regulation of food intake (Sakurai et al., 1998). ICV administration of orexin also increases activity and body temperature (Monda et al., 2001). Loss of orexin neurons leads to the disordered sleep patterns of narcolepsy, which is often accompanied by defective energy and metabolic homeostasis, including a high risk for obesity (Kok et al., 2003; Hara et al., 2005) and the potential for altered thermoregulation (Plazzi et al., 2011). Interestingly, systemic orexin, perhaps secreted from the placenta, is required for BAT differentiation and its absence during early development leads to obesity arising from inadequate BAT energy expenditure (Selayah et al., 2011).

A role for orexin neurons in the PeF-LH in the regulation of BAT thermogenesis has recently been described (Tupone et al., 2011). An anatomical substrate for the influence of orexinergic neurons on BAT thermogenesis is provided by the demonstration that the PeF-LH contains orexinergic neurons that are synaptically coupled to BAT, as shown with viral, trans-synaptic retrograde tracing (Oldfield et al., 2002; Berthoud et al., 2005; Tupone et al., 2011), and that project to the rRPa, as shown with retrograde transport of CTb following injections into the rRPa region containing BAT sympathetic premotor neurons (Tupone et al., 2011). In anesthetized rats, both injections of orexin into rRPa (Figure 5C) and glutamatergic activation of orexinergic neurons by nanoinjections of NMDA into the PeF-LH (Figure 5C) produced long lasting activations of BAT SNA and BAT thermogenesis, accompanied by marked increases in energy expenditure, indicated by sustained increases in expired CO2 that paralleled the increases in BAT heat production (Figure 5A). Curiously, the strong stimulation of BAT SNA and BAT thermogenesis elicited by injection of orexin into the rRPa or by activation of neurons in the PeF-LH required an ongoing, basal level of BAT SNA, generated in this case by maintaining the rats at a slightly cooled core body temperature. When the rats were warmed to eliminate any basal discharge on the sympathetic...
FIGURE 5 | Orexin in the rostral raphe pallidus (rRPa) or activation of neurons in the perifornical lateral hypothalamus (PeF-LH) produces a prolonged increase in BAT sympathetic nerve activity (SNA) and BAT thermogenesis in cool, but not warm rats. (A) Under cool conditions (core temperature <37°C) in which a low level of basal BAT SNA is present, nanoinjections of orexin-A (Orx-A, left panel, dashed line) in the rRPa or of N-methyl-D-aspartate (NMDA, right panel, dashed line) into the PeF-LH elicited prolonged increases in BAT SNA, BAT temperature (TBAT), and expired CO2 (Exp CO2). (B) Under warm conditions (core temperature >37°C) in which no basal BAT SNA was present, nanoinjections of Orx-A (left panel, dashed line) in rRPa or of NMDA (right panel, dashed line) into the PeF-LH failed to evoke significant changes in BAT SNA, TBAT or Exp CO2. (C) Representative histological sections illustrating nanoinjection sites in the rRPa (left panel, white arrowhead) and in the PeF-LH (right panel, white arrowhead). Note that the NMDA injection sites in the PeF-LH were located in the midst of many neurons immunohistochemically labeled for Orexin-A (red). Modified with permission from Tupone et al. (2011).
nerves to BAT, neither injection of orexin into rRPa nor activation of PeF-LH neurons elicited increases in BAT SNA (Tupone et al., 2011; Figure 5B).

These anatomical data demonstrate not only an orexenergic projection from neurons in the PeF-LH to the site of BAT sympathetic premotor neurons in the rRPa, but also a synaptic connection between orexin-containing neurons in PeF-LH and BAT sympathetic premotor neurons. Physiologically, the requirement for an ongoing level of BAT SNA, and thus an activation of BAT sympathetic premotor neurons in rRPa, in order for orexin in rRPa to evoke large and sustained increases in BAT SNA and BAT thermogenesis is consistent with a role for orexin in rRPa to change the gain of the response of BAT sympathetic premotor neurons to their activating, presumably glutamatergic excitatory inputs.

The physiological conditions which activate the orexin neurons in PeF-LH that project to rRPa to modulate the gain of the synaptic drive to BAT sympathetic premotor neurons and facilitate BAT thermogenic responses remains to be determined. Also unknown are the other potential projection targets of the orexin neurons that influence BAT thermogenesis. However, a potential role of orexin neurons in feeding could suggest a simultaneous increase in energy consumption in BAT under conditions of a high level of stored calories. Alternatively, the activation of orexin neurons in awake or aroused states could suggest an accompanying facilitation of BAT thermogenesis to increase brain and core temperatures to optimize performance. In this regard, it would be of interest to determine if the level of activity in orexin neurons parallels the ultradian oscillations in the BAT thermogenesis (Ootsuka et al., 2009). Conversely, a reduction in the activity of the orexenergic input to rRPa may contribute to the reductions in BAT thermogenesis, energy consumption, and body temperature under conditions of sleep, hibernation or starvation.

A ROLE FOR THE VENTROMEDIAL HYPOTHALAMUS IN BAT THERMOGENESIS REMAINS UNCLEAR

The seminal study of Hetherington and Ranson (1940) initiated the interest in the ventromedial hypothalamus (VMH) in energy homeostasis and several succeeding studies have suggested that the VMH affects energy expenditure via sympathetic activation of BAT thermogenesis. Electrical stimulation of the VMH increases BAT thermogenesis (Perkins et al., 1981; Minokoshi et al., 1986) and microinjection of glutamate (Yoshimatsu et al., 1993) or triiodothyronine (Lopez et al., 2010) into VMH increases BAT SNA. Conversely, lesions of the VMH attenuate BAT SNA (Nijjima et al., 1984; Sakaguchi et al., 1988). However, the methodologies used in these studies, including electrical stimulation, electrolytic lesions, and large injection volumes (i.e., 100–200 nl for microinjection studies without appropriate anatomical control experiments) significantly limit the conclusions that can be drawn from the resulting data. Furthermore, the close proximity of the VMH to other regions involved in the sympathetic regulation of BAT thermogenesis, including the DMH, the arcuate nucleus and the lateral hypothalamus, and the frequent absence of histological data in these previous studies precludes any conclusions on the role of neurons in the VMH in the control of BAT thermogenesis and BAT energy expenditure. An additional confound is that trans-synaptic retrograde tracing studies have consistently failed to identify any significant population of neurons in the VMH following injection of pseudorabies virus in BAT, even at long post-inoculation times (Bamshad et al., 1999; Oldfield et al., 2002; Cano et al., 2003; Yoshida et al., 2003). Nonetheless, a recent study found impaired diet-induced thermogenesis and lower levels of UCP1 in BAT in mice lacking PI3K specifically in the SF-1 containing neurons of the VMH (Klockener et al., 2011), suggesting a role for VMH neurons in the regulation of BAT. Whether this alteration in BAT UCP1 expression is mediated by a direct effect of altered VMH neuronal discharge on the sympathetic input to BAT, and if so, how this observation might be reconciled with the lack of viral tracer labeling of VMH neurons following inoculation of BAT, remain interesting areas for future study.

INHIBITORY REGULATION OF BAT THERMOGENESIS

HYPOXIA INHIBITS BAT THERMOGENESIS

The brainstem contains the pathways mediating the inhibition of BAT thermogenesis in response to arterial hypoxia, a reflex to restrict oxygen consumption in the face of reduced oxygen availability or compromised oxygen diffusion and transport in the blood. Systemic hypoxia or bolus systemic injections of sodium cyanide produce a prompt and complete reversal of the BAT SNA activations evoked by hypothermia and by PGE2 in the POA and this response to hypoxia is eliminated by section of the carotid sinus nerves or by inhibition of second-order arterial chemoreceptor sensory neurons in the commissural region of the nucleus of the tractus solitarius (NTS; Madden and Morrison, 2005). Interestingly, hypoxia also eliminates the BAT SNA activation resulting from bicuculline nanoinjection into the rRPa, suggesting that activation of a GABAergic input to BAT sympathetic premotor neurons in rRPa is unlikely to mediate the hypoxic inhibition of BAT thermogenesis. Indirect evidence points to a possible role for a spinal inhibitory mechanism. Similar to arterial hypoxia, disinhibition of neurons in the rostral ventrolateral medulla (RVLM) reduces the BAT SNA activation following bicuculline into the rRPa (Morrison et al., 1999) and both anatomical (Stornetta et al., 2004) and electrophysiological (Deuchars et al., 1997) studies support the existence of a bulbohypothalamic inhibitory pathway to SPNs from the RVLM. The pathway for the hypoxic inhibition of BAT metabolism between the NTS and the BAT SPNs remains to be investigated.

HYPOGLYCEMIA AND GLUCOPRIVATION INHIBIT BAT THERMOGENESIS

Similar to the hypoxia-evoked inhibition of thermogenesis, hypoglycemia and glucoprivation cause hypothermia (Freinkel et al., 1972; Mager et al., 1976), at least in part by inhibiting sympathetically mediated metabolism in BAT (Egawa et al., 1989a; Madden, 2012). This neurally regulated decrease in metabolism reduces cellular oxidative demands during conditions of reduced availability of metabolic fuel. The importance of this adaptive response, which spares scarce glucose resources for use by critical tissues such as the brain, at the expense of thermoregulation, is demonstrated by the observation that prevention of hypothermia during severe hypoglycemia results in increased mortality rates (Buchanan et al., 1991).

Glucoprivation selectively within the lateral hypothalamus reduces BAT SNA by ∼25% (Egawa et al., 1989b), whereas systemic...
glucoprivation can completely inhibit BAT SNA (Egawa et al., 1989a; Madden, 2012). More recently we have demonstrated not only that the glucoprivic agent, 2-deoxy-D-glucose (2-DG), reversed the cooling-evoked activation of BAT SNA, but that selective glucoprivation with local nanoinjection of 5-thio-D-glucose (5-TG) within the ventrolateral medulla (VLM) also completely reversed the cooling-evoked activation of BAT SNA (Madden, 2012). Interestingly, intravenous administration of 2-DG did not attenuate the activation of BAT SNA following pontomedullary transection, suggesting that the hypoglycemia-evoked inhibition of BAT SNA is not mediated solely by circuits within the medulla (Madden, 2012). The NPY/catecholaminergic projection from the VLM to the PVH may play an important role in the glucoprivic inhibition of BAT SNA, consistent with the role of VLM neurons in the glucoprivic inhibition of BAT SNA, the ability of neuronal activation in the PVH to inhibit BAT SNA (see below) and the established role of the NPY/catecholaminergic input from the VLM to the PVH in other counterregulatory responses to hypoglycemia (Ritter et al., 2011). Identifying the respective contributions of hypothalamic and medullary regions to the reduced BAT thermogenesis during conditions of reduced availability of metabolic fuel, as well as the characterization of the specific neural pathways by which hypoglycemia influences thermoregulatory neural circuits await further research.

**NEURONS IN THE PARAVENTRICULAR HYPOTHALAMIC NUCLEUS INHIBIT BAT THERMOGENESIS**

The PVH plays a major role in energy homeostasis, influencing both food intake and energy expenditure. Although the parvicellular connection of neurons in the PVH to BAT (Bamshad et al., 1999; Oldfield et al., 2002; Cano et al., 2003; Yosida et al., 2003) strongly supports a role for these neurons in the sympathetic regulation of BAT thermogenesis, their influence on the regulation of BAT thermogenesis has been controversial. Initially, neurons in the PVH were thought to play a role in the excitation of BAT SNA, since neurons in the dorsal PVH with direct projections to the spinal sympathetic preganglionic cell column are activated during fever (Zhang et al., 2000) and lesions of PVH attenuate fever (Horn et al., 1994; Caldeira et al., 1998; Lu et al., 2001). Curiously, cold-evoked BAT thermogenesis was unaffected by lesions of the PVH (Lu et al., 2001). In contrast, activation of neurons in the PVH has recently been demonstrated to inhibit BAT SNA and BAT thermogenesis (Madden and Morrison, 2009). Unilateral disinhibition of neurons in PVH with nanoinjections of bicuculline or their glutamatergic activation with NMDA injections completely inhibits increases in BAT SNA and BAT thermogenesis elicited by skin and core cooling (Figure 6A), by injections of PGE2 into the MPO that mimic fever, or by disinhibition of neurons in the DMH/DA (Madden and Morrison, 2009). Although activation of PVH neurons could attenuate the increases in BAT SNA and BAT thermogenesis evoked by injections of NMDA into the rRPa, those resulting from bicuculline injections into rRPa were unaffected by disinhibition of PVH neurons (Figure 6B), consistent with the PVH-evoked inhibition of BAT SNA being mediated by GABA_A receptors in the rRPa. That neurons in the PVH provide an inhibitory influence on BAT SNA is also supported by the observations that NPY presynaptically inhibits GABA release onto PVH neurons (Cowley et al., 1999) and microinjection of NPY into the PVH decreases BAT SNA (Egawa et al., 1991). These apparent controversies in the relation of PVH neurons to BAT thermogenesis, particularly during fever, might be explained by the presence of subpopulations of PVH neurons mediating contrasting effects or by a role of PVH neurons during fever that involves the stimulation of other fever-supporting effector systems such as the cutaneous vasculature or hormone release, whereas the inhibition of BAT thermogenesis by PVH neurons contributes to a non-febrile homeostatic function.

Controversy also exists concerning the role of melanocortin receptor activation in the PVH in energy expenditure and activation of BAT thermogenesis. Selective rescue of melanocortin-4 receptor (MC4R) expression in neurons of the PVH (and the medial amygdala) in mice lacking expression of MC4R failed to normalize (elevate) their oxygen consumption to wild-type levels (Balthasar et al., 2005). Based on these data it was suggested that PVH MC4Rs do not mediate the energy expenditure effects of melanocortins. In contrast, other groups have demonstrated that microinjection of melanocortin receptor agonists into the PVH increases core and BAT temperatures (Song et al., 2008; Skibicka and Grill, 2009). These effects of melanocortin receptor activation could be mediated by activation of presynaptic MC4Rs, which have been shown to potentiate GABAergic inputs to PVH neurons (Cowley et al., 1999). Indeed, this explanation would reconcile this controversy, since the rescue of MC4R in the study of Balthasar et al. would only rescue the postsynaptic MC4R in PVH neurons and not those located presynaptically and potentially responsible for the effects of exogenously administered melanocortin receptor agonists. This explanation is also consistent with our data indicating that the activity of neurons in the PVH is inhibitory to BAT SNA. The physiological conditions which would stimulate the BAT sympathoinhibitory output from the PVH are unknown, but may include hypoglycemia and chronic intermittent hypoxia. Another interesting possibility is that neurons in the PVH provide a tonic inhibition of BAT thermogenesis and release from this inhibition under specific conditions, such as changes in dietary composition, may activate BAT SNA and BAT energy expenditure.

**PONTINE RETROBULLAR FIELD NEURONS MAINTAIN A TONIC INHIBITION OF BAT THERMOGENESIS**

The existence of a tonically active inhibition of BAT thermogenesis from mid-pontine neurons (Figure 1) was indicated by the large increases in BAT and/or core temperatures that followed transections of the neuraxis in the vicinity of the pontomedullary junction, but which were absent if transections were made rostral to the pons, but caudal to the DMH (Rothwell et al., 1983; Amini-Sereshki and Zarrindast, 1984). Inactivation of neurons in the vicinity of the pontine reticular field produced a similar stimulation of BAT thermogenesis (Shibata et al., 1999). Neither the exact location of the neurons mediating this inhibition nor the physiological basis for its control has been determined.

**INHIBITION OF BAT THERMOGENESIS FROM NEURONS IN THE VENTROLATERAL MEDULLA AND THE NUCLEUS OF THE SOLITARY TRACT**

The VLM and the NTS contain neurons that comprise fundamental cardiovascular and respiratory regulatory circuits, including
the baroreceptor and arterial chemoreceptor reflex pathways regulating local vasoconstrictor and cardiac sympathetic premotor neurons, as well as the respiratory generating network and its regulation by central chemoreceptors. A role for neurons in these same regions in regulating BAT thermogenesis has been recently described (Cao et al., 2010). Disinhibition of neurons in the VLM with a unilateral nanoinjection of bicuculline elicits a prompt and complete inhibition of the increased BAT SNA and BAT thermogenesis due to skin and core cooling (Figure 7A), to injections of PGE2 into the MPO, to disinhibition of neurons in DMH/DA or

FIGURE 6 | Disinhibition of neurons in the paraventricular hypothalamic (PVH) nucleus inhibits the increases in BAT sympathetic nerve activity (SNA) evoked by cooling, but not those evoked by disinhibition of neurons in the rostral raphe pallidus (rRPAs). (A) Nanoinjection of bicuculline (BIC) into the PVH completely reversed the increases in BAT SNA, BAT temperature (TBAT), expired CO2 (Exp CO2) produced by whole body cooling. Core body temperature (TCORE) was also reduced; heart rate (HR) and arterial pressure (AP) were increased by BIC in PVH. (B) The increases in BAT SNA, TBAT, expired CO2, TCORE, HR, and AP evoked by nanoinjection of BIC into the rRPAs are not affected by nanoinjection of BIC into the PVH. Modified with permission from Madden and Morrison (2009).

FIGURE 7 | Activation of neurons in the ventrolateral medulla (VLM) or in the nucleus of the solitary tract (NTS) inhibits BAT sympathetic nerve activity (SNA) and BAT thermogenesis. (A) Cooling-evoked increases in BAT SNA, BAT temperature (TBAT), and expired CO2 (Exp CO2) were reversed following unilateral nanoinjection of either the glutamate receptor agonist, NMDA, or the GABAA receptor antagonist, bicuculline (BIC), into the rostral VLM (rVLM). Core temperature (TCORE) was also reduced by activation of rVLM neurons. (B) Increases in BAT SNA, TBAT, and Exp CO2 evoked by BIC nanoinjection into the rostral raphe pallidus (rRPa) were reversed by bilateral nanoinjections of BIC into the medial NTS. Modified with permission from Cao et al. (2010).
the rRPAs, or to pontomedullary transection (Cao et al., 2010). The VLM region from which inhibitions of BAT SNA could be evoked corresponds to the locations of the A1 and C1 cell groups in the region ventral to the nucleus ambiguous. Similar inhibitions of BAT SNA were also produced by bilateral injections of bicuculline into the medial NTS (Figure 7B). However, application of leptin and TRH into the NTS elicits increases in BAT temperature (Hermann et al., 2006). Whether the inhibitions of BAT thermogenesis evoked throughout this rostro-caudal extent of the VLM represent activation of a homogeneous inhibitory mechanism or stimulations of different mechanisms at different rostro-caudal levels remains to be determined, as do the mechanisms through which the VLM and the NTS inhibitions of BAT SNA is effected and the physiological circumstances under which such inhibitions of BAT thermogenesis would normally be elicited. In this regard, both hypoxia and hypoglycemia are strong inhibitory signals for BAT thermogenesis (see above) and the commissural NTS contains second-order chemoreceptor sensory neurons mediating an inhibition of BAT SNA (Madden and Morrison, 2005), and the VLM contains neurons sensitive to glucopenia (Ritter et al., 2001), which also inhibits BAT SNA (Madden and Morrison, 2008b).

SUMMARY

Brown adipose tissue thermogenesis is regulated primarily by a core thermoregulatory neural network which responds to the feedforward afferent signals from cutaneous and core body thermoreceptors and to feedback signals from brain thermosensitive neurons to alter the level of activation of the sympathetic outflow to BAT. We have summarized the research leading to a model (Figure 1) of the thermoregulatory reflex pathway through which environmental cold stimulates thermogenesis and includes the influence on this thermoregulatory network of the pyrogenic mediator, PGE₂, to increase body temperature during fever. The cold thermal afferent circuit from cutaneous thermal receptors, through second-order thermosensory neurons in the dorsal horn of the spinal cord ascends to activate neurons in the LPBel which drive GABAergic interneurons in the MnPO to inhibit warm-sensitive, inhibitory output neurons of the MPO. Through the resulting disinhibition of thermogenesis-promoting neurons in the DMH, BAT sympathetic premotor neurons in the rostral ventromedial medulla, including the rRPAs, are stimulated to increase the excitation to and responsiveness of the spinal circuits controlling BAT SPN discharge to drive BAT thermogenesis.

Hypoxia and hypoglycemia are strong metabolic regulators of BAT thermogenesis. The activity of neurons in several brain regions can influence the level of BAT thermogenesis and thus could modulate the thermoregulatory control of BAT, although their discharge does not appear to be required to mediate the skin cooling-evoked stimulation of BAT thermogenesis. These include an orexigenic excitation of BAT thermogenesis from neurons in the PeF-LH and inhibitory regulation of BAT thermogenesis by neurons in the PVH, the pontine reticular field, the VLM and the NTS.

ACKNOWLEDGMENTS

Support of the research that contributed to this review came from National Institutes of Health grants R01NS40987 (Shaun F. Morrison), R01DK57838 (Shaun F. Morrison), F32DK065401 (Christopher J. Madden), R56DK082558 (Christopher J. Madden), and an American Heart Association Scientist Development Grant (Christopher J. Madden).

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