Correction

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A hypothalamic circuit that controls body temperature

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The homeostatic control of body temperature is essential for survival in mammals and is known to be regulated in part by temperaturesensitive neurons in the hypothalamus. However, the specific neural pathways and corresponding neural populations have not been fully elucidated. To identify these pathways, we used cFos staining to identify neurons that are activated by a thermal challenge and found induced expression in subsets of neurons within the ventral part of the lateral preoptic nucleus (vLPO) and the dorsal part of the dorsomedial hypothalamus (DMD). Activation of GABAergic neurons in the vLPO using optogenetics reduced body temperature, along with a decrease in physical activity. Optogenetic inhibition of these neurons resulted in fever-level hyperthermia. These GABAergic neurons project from the vLPO to the DMD and optogenetic stimulation of the nerve terminals in the DMD also reduced body temperature and activity. Electrophysiological recording revealed that the vLPO GABAergic neurons suppressed neural activity in DMD neurons, and fiber photometry of calcium transients revealed that DMD neurons were activated by cold. Accordingly, activation of DMD neurons using designer receptors exclusively activated by designer drugs (DREADDs) or optogenetics increased body temperature with a strong increase in energy expenditure and activity. Finally, optogenetic inhibition of DMD neurons triggered hypothermia, similar to stimulation of the GABAergic neurons in the vLPO. Thus, vLPO GABAergic neurons suppressed the thermogenic effect of DMD neurons. In aggregate, our data identify vLPO-DMD neural pathways that reduce core temperature in response to a thermal challenge, and we show that outputs from the DMD can induce activity-induced thermogenesis.

thermoregulation | preoptic area | dorsomedial hypothalamus | fiber photometry | energy expenditure

Adult mammals, including humans, precisely maintain core body temperature ($T_{\rm core}$) within a narrow range. This system is essential for survival because a significant deviation of $T_{\rm core}$ can adversely affect cellular metabolism. Changes in environmental temperature activate homeostatic responses that, in turn, regulate energy expenditure, adaptive thermogenesis, and physical activity in an attempt to maintain near-optimal $T_{\rm core}$. Previous studies have shown that the preoptic area (POA) of the hypothalamus plays an important role in maintaining a stable $T_{\rm core}$ via afferent inputs from skin thermoreceptors. The direct sensing of changes in skin temperature, in turn, activates POA efferent signals that control thermal effector organs (1, 2). For example, in response to a heat stress, warmth receptors within the dorsal root ganglion provide excitatory input to POA neurons via relay neurons in the dorsal lateral parabrachial nucleus (3). In addition, recordings from in vivo models and in slice preparations identify a group of intrinsic warm-sensitive neurons (WSNs) (20-40%) that may enable animals to directly detect brain warmth and promote heat loss (4). Thermosensitive transient receptor potential channel M2 (TRPM2), which is expressed in the POA, may be part of the heat sensor in WSNs that can limit fever and induce hypothermia (5). Furthermore, activation of glutamatergic neurons in several POA subareas results in severe hypothermia, whereas activation of subsets of POA GABAergic neurons

has been found to have only a minimal or no effect on $T_{\rm core}$ (5, 6). However, the precise nature of the POA projections that play a role in suppressing thermogenesis have not been elucidated. Thus, although several relevant brain regions have been suggested to act in concert with the POA to regulate $T_{\rm core}$ (1, 2), including the dorsomedial hypothalamus (DMH), the periventricular nucleus, and the raphe pallidus nucleus, functional connections between the POA and other thermoregulatory regions are largely unknown.

In the present study, we began by defining sites in which cFos staining was induced by changes in environmental temperature. These studies identified temperature-activated neurons in the ventral part of the lateral preoptic nucleus (vLPO) and the dorsal part of the DMH (DMD). We confirmed that DMD neurons responded to cold by using fiber photometry, and tested the ability of these and POA neurons to affect $T_{\rm core}$, energy expenditure (EE), and physical activity by using optogenetics (7) or designer receptors exclusively activated by designer drugs (DREADDs) (8). These functional studies identified neural pathways in which heat-activated GABAergic neurons in the vLPO inhibit cold-activated neurons in the DMD to suppress thermogenesis and lower $T_{\rm core}$.

Results

Activation of GABAergic POA Neurons Is Sufficient To Drive Hypothermia.

The POA is known to receive input from cold- and heat-sensitive neurons (1). However, although recent data have identified some of the neural substrates for temperature sensing (5, 6), components of the neural circuit(s) that respond to thermal challenges have not been

Significance

Thermal homeostasis is essential for survival in mammals. Although it is known that temperature-sensitive neurons in the hypothalamus can control body temperature, the precise neural types and dynamics of neurons responding to changes in environmental temperature are not well defined. In this study, we identified subsets of temperature-activated neurons in two hypothalamic nuclei, the preoptic area (POA) and the dorsomedial hypothalamus (DMH), and showed that modulating their activity can lead to alterations in core temperature. The data further suggest that heat-activated GABAergic neurons in the POA reduce the activity of cold-activated neurons in the DMH, which function to increase thermogenesis and physical activity. These data identify a neural circuit that controls core temperature and thermogenesis.

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fully elucidated. For example, there are conflicting data concerning the potential role of GABAergic neurons in the POA (2, 5, 6). To identify these key neural populations, we characterized the pattern of cFos activation following a thermal challenge (SI Appendix, Fig. S1). We found heat-induced cFos expression in the medial preoptic area (MPO) and the vLPO, and it colocalized with the GABAergic marker GAD67 (Fig. 1A and SI Appendix, Fig. S5A). Both the MPO and the vLPO have been suggested to play important roles in thermoregulation (1, 2, 9-11). To test the function of the vLPO neuronal population, we targeted expression of channelrhodopsin-2 (ChR2) fused with eYFP to GABAergic neurons within the vLPO by injecting Credependent adeno-associated virus (AAV) 5 viruses into the vLPO of Vgat-IRES-Cre driver mice (Vgat stands for vesicular GABA transporter) (Fig. 1B). In slice recordings, we confirmed that the delivery of blue light to ChR2-expressing vLPO^{Vgat} neurons resulted in neural excitation (Fig. 1C). In vivo, we found that light-induced activation of vLPO^{Vgat} neurons triggered a rapid reduction in T_{core} with a decrease in physical activity ($\Delta T = -1.8 \pm 0.09$ °C at t = 30 min, mean \pm SEM, Fig. 1 D and E) in freely behaving mice. This effect was specific to ChR2, because injection of control AAV5 viruses (expressing eYFP) did not significantly affect T_{core} or activity.

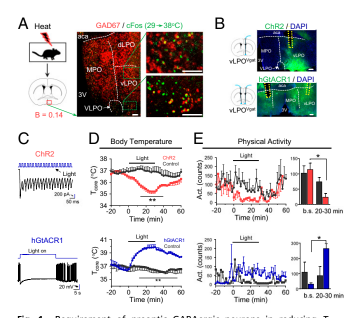


Fig. 1. Requirement of preoptic GABAergic neurons in reducing T_{core}. (A) Heat-induced (38 °C, 2 h) cFos colocalized with the GABAergic marker GAD67 in the vLPO (no. of cFos⁺ and GAD67⁺/no. of cFos⁺ = 36.3 \pm 2.4%, n = 3). The dashed white lines indicate boundaries between subregions. (B) Scheme of optogenetic modulation and viral expression of ChR2 (excitatory) or hGtACR1 (inhibitory) in vLPOVgat neurons. The dashed yellow lines indicate the positions of optical inserts. (C) Slice recordings of neurons expressing ChR2 (Upper) or hGtACR1 (Lower). Blue light (blue, 6 mW, 40 Hz) faithfully elicited photocurrents in ChR2-expressing neurons in the vLPO. A blue light pulse (6 mW) completely silenced hGtACR1-expressing neurons in the vLPO. (D and E) T_{core} (D) and activity (E) changes after optogenetic stimulation in mice expressing ChR2 (Upper, n = 4) or hGtACR1 (Lower, n = 3) in vLPO^{Vgat} neurons. Stimulation protocol for ChR2: unilateral light pulses for 2 s (473 nm, 10 mW, 20 Hz, 40% on) followed by a 2-s break, with the sequence repeating for 30 min. For hGtACR1: bilateral light on for 30 s (473 nm, 6 mW) followed by a 90-s break, with the sequence repeating for 30 min. Bar graph of activity changes (average of 10-min interval) are shown in the right. Baselines (b.s.) represents the average counts between t = -30 and -20 min. (20–30 min) represents the average of counts between t=20 and 30 min. (Scale bars: A, 100 μ m; B; 200 μ m.) All data are plotted as mean \pm SEM. The P values compared with control group (eYFP) are calculated based on statistical tests listed in SI Appendix, Table S1. * $P \le 0.05$; ** $P \le 0.01$. aca, anterior commissure, anterior part; B, bregma; dLPO and vLPO, dorsal and ventral part of lateral preoptic nucleus respectively; MPO, medial preoptic nucleus; 3V, third ventricle; VLPO, ventrolateral preoptic nucleus.

Next, we asked whether inhibiting these neurons was sufficient to drive hyperthermia. Using the *Guillardia theta* anion channelrhodopsin 1 (hGtACR1), which robustly silences neural activity in response to blue-yellow light (12), we found that blue light delivery was sufficient to silence neurons in slice recordings (Fig. 1C). Remarkably, light stimulation of mice expressing hGtACR1 in vLPO $^{\text{Vgat}}$ neurons caused severe hyperthermia with elevated activity levels (maximal $T_{\text{core}} = 40.6\,^{\circ}\text{C}$, Fig. 1D and E). Indeed, we needed to minimize the time in which animals received light illumination to prevent hyperthermia-induced death.

We also tested the function of MPO GABAergic neurons by targeted expressing of ChR2 to GABAergic neurons within the MPO of Vgat-IRES-Cre driver mice (*SI Appendix*, Fig. S5B). The GABAergic neurons in the MPO have been suggested to play important roles in thermoregulation (1, 2). Surprisingly, we found that optogenetic activation of these MPO $^{\text{Vgat}}$ neurons did not significantly affect T_{core} or activity (*SI Appendix*, Fig. S5 C and D). Similar to our results, DREADD activation of these neurons has a minimal effect on T_{core} (5).

Thus, our results establish that activation of GABAergic neurons within a preoptic subregion (vLPO) can inhibit thermogenesis, whereas inhibition of these neurons dramatically raises core temperature. We next explored the functional targets of these neurons.

Critical Role of the POA→DMD Connection in Reducing T_{core}. We thought that vLPO^{Vgat} neurons might project to the DMH because the DMH is known to participate in thermoregulation, and because we observed that thermal stimuli induced strong cFos staining within the DMD (SI Appendix, Fig. S1). To test whether vLPO^{Vgat} neurons directly innervate DMD neurons, we first performed anterograde tracing from these neurons by injecting ChR2 into vLPO^{Vgat} neurons (Fig. 24) and found staining of nerve terminals in the DMD (Fig. 2B). We then performed retrograde labeling by injecting the retrograde protein, cholera toxin B subunit (CTb; ref. 13) into the DMD and found it labeled many neurons in the POA, including heat-activated neurons (as indicated by induction of cFos) in the vLPO (Fig. 2C).

We tested the function of this vLPO \rightarrow DMD projection by stimulating vLPO $^{\text{Vgat}}$ terminals in the DMD after viral injection of ChR2 into the vLPO (Fig. 2D). We found that stimulation with blue light triggered a significant reduction of T_{core} ($\Delta T = -2.3 \pm 0.6$ °C at t = 60 min, mean \pm SEM, Fig. 2E) along with a decrease in activity (Fig. 2F). The magnitude of the effect was similar to that observed after direct stimulation of Vgat neurons in the vLPO. Thus, stimulation of vLPO $^{\text{Vgat}}$ nerve terminals in the DMD recapitulated the phenotype observed when vLPO $^{\text{Vgat}}$ cell bodies were stimulated (Fig. 1 D and E, Upper).

These functional data suggest that vLPOVgat neurons inhibit DMD neurons and reduce core temperature and thermogenesis. We next set out to confirm this inhibition directly by recording inhibitory postsynaptic currents (IPSCs) in DMD neurons after vLPO vgat terminals were stimulated. This recording is important because our data could also be explained by inhibition of axon fibers that pass through the DMD without directly innervating DMD neurons. In slice preparations, we confirmed that stimulation of vLPOVgat terminals expressing ChR2 by blue lightinduced IPSCs in the DMD neurons and that these currents were blocked by a GABA_A receptor antagonist, bicuculline (Fig. 2G). The onset of these currents was <3 ms after light delivery, which is within the time range of monosynaptic transmission (14). Furthermore, to identify which types of DMD neurons are synaptically connected to the vLPO^{Vgat} neurons, we used the recording pipette to isolate mRNA from individual, light-responsive DMD neurons immediately after recordings and analyzed gene expression by single-cell reverse transcription PCR (RT-PCR) (15). These singlecell RT-PCR data showed that there were both Vgat⁺ and Vglut2⁺ postsynaptic neurons in the DMD, with the majority being Vgat+ (Fig. 2H and SI Appendix, Fig. S6).

Several reports (16–21) have suggested that DMD neurons can regulate thermogenesis, but the inputs to these neurons have not been elucidated. Our studies in anterograde and retrograde

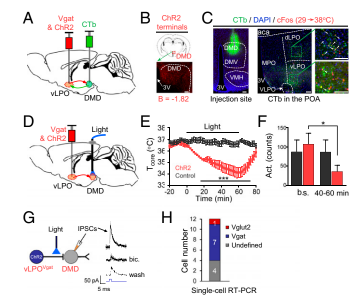


Fig. 2. Critical role of the vLPO^{Vgat} \rightarrow DMD connection in reducing T_{core} . (A) Scheme for anterograde tracing using ChR2 and retrograde tracing using CTb. (B) vLPO^{Vgat & ChR2} terminals in the DMD. (C) vLPO neurons were retrogradely labeled by CTb injected in the DMD, which colocalized with heatinduced cFos. White arrows indicate the colocalization of cFos and CTb. (Scale bars: 100 µm.) (D) Scheme for terminal optogenetic stimulation in the DMD after ChR2 injection into vLPO^{Vgat} neurons. Both injection and stimulation were bilateral. (E and F) Bilateral terminal optogenetic stimulation in the DMD-reduced T_{core} (E) and activity (F) (n = 5). Illumination protocol: light pulses for 2 s (473 nm, 10 mW, 20 Hz, 40%) followed by a 2-s break, with the sequence repeating for 1 h. Bar graph of activity changes (average of 20-min interval) are shown in F. Baselines (b.s.) represents the average of counts in between t = -30 and -10 min. (40–60 min) represents the average of counts between t = 40 and 60 min. (G) Induction of inhibitory postsynaptic currents (IPSCs) in DMD neurons by light stimulation (blue, 6 mW, 2 ms) of ChR2expressing terminals projected from vLPO^{Vgat} neurons. IPSCs were blocked by bicuculline (bic.), and partially recovered after wash. Thick lines indicate the mean, whereas the shaded areas indicate SD. (H) Single-cell RT-PCR analysis of recorded cells. All data are plotted as mean \pm SEM (except in G); P values compared with control group (532 nm) are calculated based on statistical tests listed in *SI Appendix*, Table S1. * $P \le 0.05$; *** $P \le 0.001$. DMD and DMV, dorsal and ventral part of dorsomedial hypothalamic nucleus respectively; VMH, ventromedial hypothalamic nucleus.

tracing, optogenetic activation of terminals from vLPOVgat neurons in the DMD, slice recordings, and single-cell RT-PCR suggest the possibility that there are functional connections between the vLPO^{Vgat} neurons and DMD^{Vgat} or DMD^{Vglut2} neurons to regulate core temperature. We next tested these connections directly by monitoring and modulating the activity of neurons in the DMD.

Fiber Photometry Reveals That DMD Neurons Are Cold Activated. The observation that inhibitory neurons that project to the DMD cause hypothermia suggests that neurons within the DMD may be cold sensitive (via either direct or indirect inputs) and act to raise core temperature (which would explain why inhibiting their inhibitors would raise T_{core} ; Fig. 1D). We first tested this hypothesis by using the calcium reporter GCaMP6f (22) to directly measure neural Ca²⁺ signals by fiber photometry after changes in environmental temperature. Similar experimental setups have yielded stable recordings from the dorsal raphe (23) (Fig. 3A). We recorded Ca²⁺ signals from both glutamatergic (DMD^{Vglut2}) and GABAergic (DMD^{Vgat}) neurons separately by injecting a Cre-dependent version of GCaMP6f into Vglut2-IRES-Cre and Vgat-IRES-Cre mice, respectively (Fig. 3B). We controlled floor temperature (T_{floor}) of mice with a Peltier controller (Fig. 3C). The DMD^{Vglut2} neurons showed significant Ca²⁺ signals

in response to cooling (25–13 °C, maximal response, $\Delta F/F_{0 \text{ max}}$ = $16.5 \pm 2.6\%$, mean \pm SEM; Fig. 3D, Left), but not to warming (25– 38 °C, Fig. 3D, Center). The responses occurred within seconds of the temperature shift and diminished quickly in cold. Approximately 2 min following the temperature shift (t = 110-120 s), mean neuronal activity levels $(\Delta F/F_{0 (110-120 \text{ s})})$ were not higher than baseline (Fig. 3D, Right).

We also observed an even larger Ca²⁺ increases in response to cooling in DMD^{Vgat} neurons (25–13 °C) ($\Delta F/F_{0 \text{ max}} = 22.7 \pm 2.8\%$, mean \pm SEM; Fig. 3*E*, Left). This Ca²⁺ signal increased rapidly (within seconds) after the temperature shift and diminished more slowly than in DMD vglut2 neurons. $\Delta F/F_{0(110-120s)}$ was significantly larger than baselines (Fig. 3E, Right). We fitted the response curve by using a sigmoidal function and found the full width at half maximum (FWFM or $T_{1/2 \text{ max}}$) of the DMD $^{\text{Vgat}}$ neurons was longer than that of DMD $^{\text{Vglut2}}$ neurons ($T_{1/2 \text{ max}} = 22.6 \text{ s}$ and 16.5 s, reserving the SV $^{\text{Vglut2}}$ neurons ($T_{1/2 \text{ max}} = 22.6 \text{ s}$ and 16.5 s, reserving the SV $^{\text{Vglut2}}$ spectively; SI Appendix, Fig. S8 A and B).

Taken together, these results indicate that both GABAergic and glutamatergic neurons in the DMD are activated by cooling. The response of DMD^{Vgat} neurons lasted longer than that of DMD vglut2 neurons. All of our temperature stimuli-evoked calcium responses appeared to result from periphery sensory input, rather than from a slow change in body temperature, based on

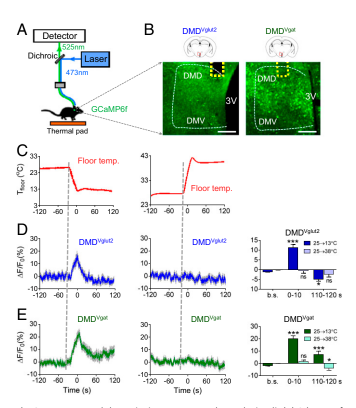


Fig. 3. DMD neural dynamics in response to thermal stimuli. (A) Scheme of fiber photometry setup. (B) The expression of GCaMP6f in the DMD driven by Vglut2-IRES-Cre or Vgat-IRES-Cre. The dashed yellow lines indicate the positions of optical inserts. The dashed white lines indicate the boundary of the DMH. (Scale bars: 200 μ m.) (C) Controlled floor temperature (T_{floor}) with a Peltier controller. (Left) Cooling traces (25-13 °C). (Right) Heating traces (25–38 °C). (D) Cooling, but not warming-activated DMD^{Vglut2} neurons. Thick lines indicate the mean, whereas the shaded areas indicate SEM. $\Delta F/F_o$ represents change in GCaMP6f fluorescence from the mean level before the floor temperature change, b.s. represents baseline, which is the averaged $\Delta F/F_0$ between t=-20 and -120 s. (0-10) and (110-120 s) represent the averaged $\Delta F/F_0$ between t = (0-10 s) and t = (110-120 s), respectively (n = 3). (E) Cold, but not warmth, activated DMD Vgat neurons (n = 4). All data are plotted as mean \pm SEM. The P values, compared with baselines, are calculated based on statistical tests listed in SI Appendix, Table S1. * $P \leq 0.05$; ***P ≤ 0.001; ns, not significant. DMD and DMV, dorsal and ventral part of dorsomedial hypothalamic nucleus, respectively; 3V, third ventricle.

the fast-rising kinetics. The calcium responses occurred within seconds after the temperature shift (Fig. 3 D and E), whereas the changes in $T_{\rm core}$ happened minutes after the temperature shift (SI Appendix, Fig. S2).

DMD Neurons Promote Thermogenesis. The finding that cooling leads to abrupt Ca²⁺ transients in DMD Vglut2 neurons suggests that these neurons might play a functional role in cold-induced thermogenesis. We next tested this by using DREADDs to activate DMD Vglut2 neurons remotely, allowing us to measure energy expenditure (EE) without being impeded by optical fibers required for optogenetics. As predicted, we found that injection of the ligand clozapine N-oxide (CNO) into mice expressing designed human M3 muscarinic receptor coupled to Gq (hM3D) (8) in DMD Vglut2 neurons strongly increased $T_{\rm core}$, along with an increase in EE and physical activity ($\Delta T = 1.4 \pm 0.3$ °C, $\Delta EE = 47.3 \pm 5.1\%$ at t = 100 min, mean \pm SEM; Fig. 4.4–D). These increases are direct evidence that activation of DMD Vglut2 neurons promotes thermogenesis, which is consistent with previous studies showing that the DMD may send excitatory (or glutamatergic) projections to promote cold- or febrile-induced thermogenesis (16, 17, 24, 25).

The findings that cooling induced calcium transients in DMD^{Vgat} neurons (Fig. 3E) and that cold-induced cFos colocalized with GAD67 in the DMD (SI Appendix, Fig. S7A) suggest that these neurons could also be important for regulating coldinduced thermogenesis. We tested this by using both DREADDs and optogenetics to activate DMD^{Vgat} neurons. We found that injection of CNO, but not saline, into mice expressing h3MD in DMD^{Vgat} neurons resulted in a slow, yet long-lasting increase in T_{core} , with an increase in EE and activity ($\Delta T = 1.3 \pm 0.2 \,^{\circ}\text{C}$, $\Delta EE = 47.4 \pm 7.5\%$ at t = 100 min, mean \pm SEM; Fig. 4 E–H). We fitted the response curve with a sigmoidal function and calculated maximum change rate (k) and full width at half maximum (FWFM) (SI Appendix, Fig. S8 C and D). For the same dose of CNO, the k for the T_{core} , EE, and activity response curves were smaller for DMD^{Vgat} neural activation than was observed after DMD^{Vglut2} neural activation. However, the FWFWs were longer after DMD^{Vgat} neural activation versus activation of DMD^{Vglut2} (SI Appendix, Fig. S8 C and D). The kinetics of these biological responses matched that seen by recording from these neurons, which indicates that cold-induced Ca²⁺ transients of DMD^{Vgat} neurons diminished more slowly than that of DMD^{Vglut2} neurons (Fig. 3 D and E and SI Appendix, Fig. S8 A and B). Also, we observed an increase in T_{core} and activity after optogenetic stimulation of mice expressing ChR2 in DMD^{Vgat} neurons (SI

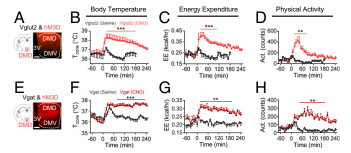


Fig. 4. Activation of DMD neurons increases body temperature. (A) Schematic and representative images of viral expression of hM3D in DMD Vglut2-IRE5-Cre+ neurons. (B-D) Activation of DMD^{Vgut2} neurons (n = 5) by CNO injection (i.p. at t = 0, 1.5 mg/kg body weight) resulted in significant increases in $T_{\rm core}$ (B), EE (C), and activity (D). (E) Schematic and representative images of viral expression of hM3D in DMD Vgat-IRE5-Cre+ neurons. (F-H) Activation of DMD^{Vgat} neurons (n = 8) by CNO injection (i.p. at t = 0, 1.5 mg/kg body weight) resulted in significant increases in $T_{\rm core}$ (F), EE (G), and activity (H). (Scale bars: 100 μ m.) All data are plotted as mean \pm SEM. The P values, compared with the saline group, are calculated based on statistical tests listed in SI Appendix, Table S1. **P \leq 0.01; ***P \leq 0.001.

Appendix, Fig. S8 B–D). In aggregate, these data show that activation of both glutamatergic and GABAergic neurons in the DMD can increase $T_{\rm core}$, EE, and activity, although the kinetics of the biological responses to activation of these neurons were different.

Inhibition of DMD Neurons Is Sufficient To Drive Hypothermia. The finding that vLPO Vgat inputs to the DMD lower core temperature led us to predict that optogenetic inhibition of DMD neurons would have a similar effect. This prediction was tested by injecting AAV9 viruses expressing hGtACR1 into the DMD of Vglut2-IRES-Cre or Vgat-IRES-Cre driver mice (thereby driving hGtACR1 expression in DMD Vglut2 or DMD Vgat neurons, respectively). We found that blue light stimulation of mice expressing hGtACR1 in DMD Vglut2 neurons resulted in a significant reduction in T_{core} , along with a decrease in activity ($\Delta T = -1.3 \pm 0.2$ °C at t = 60 min, mean \pm SEM; Fig. 5 A and B), similar to optogenetic activation of the vLPO Vgat neurons (Fig. 1 D and E). Similarly, blue light stimulation of mice expressing hGtACR1 in DMD Vgat neurons resulted in significant reductions in T_{core} , along with a decrease in activity ($\Delta T = -2.5 \pm 0.5$ °C at t = 60 min, mean \pm SEM; Fig. 5 C and D). Taken together, these results define elements of a POA→DMH neural circuit in the hypothalamus (including vLPO Vgat →DMD Vglut2 and vLPO Vgat →DMD Vgat connections) that regulate thermogenesis.

Discussion

The maintenance of a stable core temperature is essential for survival. Our study elucidates a central mechanism through which changes in core temperature (in response to alternations in ambient temperature or other stimuli) elicit a set of adaptive thermogenic responses that defend $T_{\rm core}$. In aggregate, we find that in response to a heat challenge, heat-activated GABAergic neurons in the vLPO directly inhibit the activity of cold-activated glutamatergic and GABAergic neurons in the DMD to lower $T_{\rm core}$, and do so (in part) by suppressing EE and activity. Thus, we have elucidated pathways for controlling body temperature and activity-induced thermogenesis.

In connection with our study, previous studies have suggested that the vLPO is an important site for thermoregulation. Local warming of the vLPO causes paw vasodilation in rats (9), suggesting the existence of WSNs in this area that can drive hypothermia. Also, the vLPO is labeled when a retrograde tracer (pseudo rabies virus) is injected into the interscapular brown adipose tissue (BAT) of rats (26), suggesting its involvement in controlling BAT thermogenesis. Interestingly, a recent study discovered that a heat-sensitive channel, TRPM2, in the POA (including the vLPO) may be part of the WSN heat sensor to limit fever (5). Thus, it will be interesting to see whether TRPM2 is important for vLPO neurons to detect brain warmth and lower $T_{\rm core}$.

We found that heat activated a subset of GABAergic neurons in the vLPO, which then lowered T_{core} . Although a similar (or stronger) effect was observed after activation of glutamatergic neurons in several preoptic subregions [MPA (6), and MPO (5)], activation of GABAergic neurons in these areas has a small effect on T_{core} , and the role of GABAergic neurons in the vLPO was not studied previously. We found that activation of GABAergic neurons in the vLPO is sufficient to induce hypothermia (Fig. 1 B and D). However, this activation does not include the ventrolateral preoptic nucleus (VLPO), which is important for sleep regulation (27, 28) but may be dispensable for thermoregulation, because its lesion has a minimal effect on T_{core} (27). The vLPO GABAergic neurons comprise a functionally confirmed GABAergic population in the POA. In addition, we found that inhibiting vLPO GABAergic neurons induced fever-level hyperthermia, suggesting that these neurons can control T_{core} in either direction (Fig. 1D), and that these neurons provide a key entry point for studying the neural mechanisms controlling thermogenesis. Furthermore, we found heat can also activate a subset of glutamatergic neurons in the vLPO, which can drive severe hypothermia and hypoactivity (SI

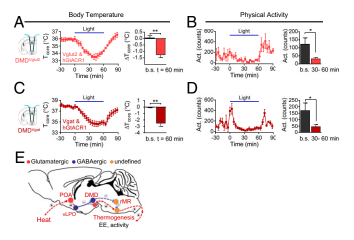


Fig. 5. Optogenetic inhibition of DMD neurons induces hypothermia. (A and B) Bilateral inhibition of DMD^{Vgut2} neurons via hGtACR1 resulted in significant decreases in $T_{\rm core}$ (A) and activity (B) (n=4). $\Delta T_{\rm core}$ represents the $T_{\rm core}$ changes from the mean level before light delivery (t=-30 to -10 min). The baseline (b.s.) (average of t = -30 to -10 min) and t = 60 min are in the bar graph. The average of activity in 30-min intervals between t = -40 and -10 min (baseline, b.s.) and between t=30 and 60 min are shown in the bar graph. Stimulation protocol: light on for 30 s (473 nm, 10 mW) followed by a 90-s break, with the sequence repeating for 1 h. (C and D) Bilateral inhibition of DMD^{Vgat} neurons (n = 4) via hGtACR1 resulted in significant decreases in T_{core} (C) and activity (D). Stimulation protocol is the same as in A. (E) Model for heat-induced suppression of thermogenesis. Sold line represents the connection verified in the current study. Dash lines represents proposed connections based our data and other reports. (+), activation. (-), inhibition. All data are plotted as mean \pm SEM. The P values, compared with baseline (b.s.), are calculated based on statistical tests listed in SI Appendix, Table S1. * $P \le 0.05$; ** $P \le 0.01$. DMD, dorsal part of dorsomedial hypothalamic nucleus; POA, preoptic area; rMR, rostral medullary region; vLPO, ventral part of lateral preoptic nucleus.

Appendix, Fig. S3). Thus, vLPO GABAergic neurons might receive local inputs from vLPO glutamatergic neurons or other preoptic glutamatergic neurons as proposed (1, 2) (such as the MnPO, because the MnPO receives inputs from the periphery (3) and sends glutamatergic outputs specifically to the vLPO; SI Appendix, Fig. S4). These inputs must be functionally confirmed.

In addition to the POA, the DMH is also an important site for thermoregulation. Although a POA

DMH connection has been proposed to play a role in thermoregulation (1, 2), direct functional confirmation of this connection is lacking. For example, pioneering studies suggest that there are anatomical connections between DMH neurons and MnPO or MPO neurons, and DMH thermogenic neurons are differentially regulated by GABAergic and glutamatergic inputs (18, 24, 25). However, it is equally likely for these inputs to arise from preoptic subregions (such as the MnPO or MPO) vs. other brain regions. In addition, activation of GABAergic neurons in subregions including the MnPO and MPO have a small effect on T_{core} (refs. 5 and 6 and SI Appendix, Fig. S5). We thus wondered whether vLPO^{Vgat} neurons innervate DMD neurons and reduced $T_{\rm core}$ in response to heat by inhibiting DMD neurons. To test this hypothesis, we combined anterograde and retrograde tracing, terminal optogenetic stimulation, IPSC recording, and single-cell RT-PCR (Fig. 2) to show that vLPO^{Vgat} DMD connections are critical for bidirectionally regulating $T_{\rm core}$ and activity. Interestingly, we found that the vLPO^{Vgat} neurons preferentially innervated DMD^{Vgat} neurons, compared with DMD^{Vglut2} neurons (Fig. 2*H*).

Previous studies have shown that DMD neurons, especially glutamatergic neurons, play an important role in promoting thermogenesis (16–18, 29, 30). However, the response of these neurons to thermal stimuli was unknown. Here, we recorded calcium dynamics in the DMD by using fiber photometry and found, surprisingly, that both glutamatergic and GABAergic neurons were activated by cooling (25–13 °C; Fig. 3 D and E), but that these neurons responded with different dynamics. The glutamatergic neurons responded more abruptly and their response decayed more quickly compared with the GABAergic neurons (SI Appendix, Fig. S5). Consistent with these kinetics, we found that increases in T_{core} , EE, and activity levels resulting from CNO-mediated activation of DMD glutamatergic neurons arose more rapidly and decayed more quickly than those elicited by CNO-mediated activation of DMD GABAergic neurons (Fig. 4 and SI Appendix, Fig. S8).

The role played by GABAergic neurons in the DMD in thermoregulation has not been studied previously, although the DMD contains more GABAergic than glutamatergic neurons (31). Here, we have shown unequivocally that these GABAergic neurons are essential for controlling thermogenesis. Their activation strongly promoted increases in $T_{\rm core}$, EE, and physical activity (Fig. 4 $E\!-\!H$ and SI Appendix, Fig. S5), whereas their suppression reduced T_{core} and activity (Fig. 5 C and D). We therefore have uncovered a type of thermogenic neuron.

Several reports have suggested that neurons expressing the leptin receptor (LepR) in the DMD are important for thermogenesis (19–21). Cold-induced cFos expression colocalizes with LepR (21), and activation of these LepR neurons increases T_{core} and EE (19). Our immunostaining results suggest that DMD^{LepR} neurons contain both glutamatergic and GABAergic types, with the majority being glutamatergic (SI Appendix, Fig. S9). Thus, DMD^{LepR} neurons might be a downstream target of the vLPO^{Vgat} neurons.

It remains unclear how DMD neurons are connected with premotor neurons to direct thermogenesis. Early observations suggested that glutamatergic neurons may project directly to premotor neurons within the rostral medullary region (rMR) to promote thermogenesis (16-18). It has also been reported that rMR premotor neurons receive inhibitory inputs from other medullary regions (32, 33). Thus, DMD Vglut2 neurons may directly innervate rMR premotor neurons to promote thermogenesis, and DMD^{Vgat} neurons might disinhibit rMR neurons by suppressing their inhibitory inputs.

Delineating the specific neural cell types involved in thermoregulation is a key step toward understanding these critical neural circuits. Using the PhosphoTRAP approach (34), which enables the immunoprecipitation of translational ribosomes via phosphorylated ribosomal protein S6 (a marker of neural activity), we (SI Appendix, Fig. S10) and Tan et al. (35) independently discovered that brain-derived neurotrophic factor (BDNF), a classic neurotrophic factor (36), is transcriptionally activated following heat challenge and is a novel marker for heat-activated neurons. It is interesting to see that heat affects the expression of a neurotrophic factor, suggesting that long-term heat challenge may affect nerve growth and cause remodeling of thermoregulatory networks. The exact role played by BDNF in this context must be further tested. BDNF-expressing neurons in the ventromedial preoptic nucleus (VMPO) can drive hypothermia without affecting physical activity (35). Thus, the circuits involving VMPO neurons and vLPO vgat neurons may act coordinately to regulate EE, BAT thermogenesis, activity, and vasodilation to lower T_{core} .

Our results establish a neural circuit (Fig. 5E) for regulating heat loss behaviors, in which environmental heat indirectly activates POA glutamatergic neurons (refs. 1–3, 5, and 6 and SI Appendix, Fig. S3), which may, in turn, activate a population of GABAergic neurons in the vLPO (Fig. 1 and SI Appendix, Fig. S4). The vLPO GABAergic neurons inhibit thermogenic neurons in the DMD to suppress EE and activity, thereby lowering $T_{\rm core}$ (Figs. 2, 4, and 5). The DMD neurons include both glutamatergic and GABAergic subtypes and are cold activated (Fig. 2 and 3). DMD glutamatergic neurons may send excitatory input to premotor neurons in the rMR to stimulate thermogenesis (16– 18). DMD GABAergic neurons might disinhibit rMR premotor neurons by suppressing their inhibitory inputs (32, 33).

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Materials and Methods

Mice. Animal care and use conformed to institutional guidelines of ShanghaiTech University, Shanghai Biomodel Organism Co., and governmental regulations. All experiments were performed on adult mice (8–16 wk old). Mice were housed under controlled temperature (22–25 °C) in a 12-h reverse light/dark cycle (light time, 8 PM to 8 AM) with a standard chow diet [4% (wt/wt) fat SPF Rodent Feed] and ad libitum drinking water. The following mice strains were from Jackson Laboratory (USA): C57BL/6J (000664); Vglut2-IRES-Cre (016963); Vgat-IRES-Cre (028862); Ai14 (007914); LepR-Cre (008320); Gad67-GFP (006340).

AAV Vectors. We used Cre-inducible AAV vectors (titer >10¹²) from the following sources: the Vector Core at the University of North Carolina at Chapel Hill (AAV5-EF1a-DIO-hChR2(H134R)-eYFP, AAV5-EF1a-DIO-eYFP, AAV5-hSyn-DIO-hM3D-mCherry, and AAV5-hSyn-DIO-hM4D-mCherry), the Vector Core at the University of Pennsylvania (AAV5-Syn-Flex-GCaMP6f), and Shanghai Taitool Bioscience Co. (AAV9-hSyn-GtACR1-P2A-EGFP). The latter construct was a gift by Minmin Luo, National Institute of Biological Sciences, Beijing, originally described in ref. 12.

Surgeries. A detailed description is provided in *SI Appendix, SI Materials and Methods*. We delivered 0.2 μ L (unless specified) of AAV virus through a pulled-glass pipette and a pressure microinjector (Nanoject II, 3-000-205A, Drummond). The fiber-optic inserts (200 μ m i.d., AniLab Co.) were chronically implanted (200 μ m above viral injection sites) and secured with dental cement.

Behavioral Assays. A detailed description is provided in *SI Appendix, SI Materials and Methods.* T_{core} , EE, and activity were monitored by animal monitoring system with temperature telemetry (Columbus, with G2 E-Mitter transponders).

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Immunohistochemistry. A detailed description is provided in SI Appendix, SI Materials and Methods.

Fiber Photometry. A detailed description is provided in *SI Appendix, SI Materials and Methods*. The GCaMP6f fluorescence signals were acquired with a fiber photometry system (Fscope, Biolinkoptics, China). The floor temperature was controlled by a Peltier controller by customized Labview code (National Instrument) described in ref. 37.

Electrophysiological Recordings and Single-Cell RT-PCR. A detailed description is provided in *SI Appendix, SI Materials and Methods*.

PhosphoTRAP and mRNA Sequencing. A detailed description is provided in *SI Appendix, SI Materials and Methods*. The procedure of PhosphoTRAP was similar as described in ref. 34 with some modification.

RNA Fluorescent In Situ Hybridization. A detailed description is provided in *SI Appendix, SI Materials and Methods.* RNA probes against Vgat, GAD67, Vglut2, and BDNF were constructed according to the description on Allen Brain Atlas (www.brain-map.org).

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