Evidence for a neuromuscular circuit involving hypothalamic interleukin-6 in the control of skeletal muscle metabolism

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Hypothalamic interleukin-6 (IL6) exerts a broad metabolic control. Here, we demonstrated that IL6 activates the ERK1/2 pathway in the ventromedial hypothalamus (VMH), stimulating AMPK/ACC signaling and fatty acid oxidation in mouse skeletal muscle. Bioinformatics analysis revealed that the hypothalamic IL6/ERK1/2 axis is closely associated with fatty acid oxidation– and mitochondrial-related genes in the skeletal muscle of isogenic BXD mouse strains and humans. We showed that the hypothalamic IL6/ERK1/2 pathway requires the α2-adrenergic pathway to modify fatty acid skeletal muscle metabolism. To address the physiological relevance of these findings, we demonstrated that this neuromuscular circuit is required to underpin AMPK/ACC signaling activation and fatty acid oxidation after exercise. Last, the selective down-regulation of IL6 receptor in VMH abolished the effects of exercise to sustain AMPK and ACC phosphorylation and fatty acid oxidation in the muscle after exercise. Together, these data demonstrated that the IL6/ERK axis in VMH controls fatty acid metabolism in the skeletal muscle.

INTRODUCTION

The hypothalamus regulates metabolic processes (1, 2) and functions of peripheral organs and glands such as the liver (3), white adipose tissue (4), brown adipose tissue (5), thyroid gland (6), and others. Neuronal activity in the ventromedial hypothalamus (VMH) controls energy expenditure and skeletal muscle metabolism, including fatty acid oxidation and glucose metabolism (7, 8). In this context, hormones and neuropeptides, such as leptin and orexin, can modify muscle metabolism in murine models by stimulating VMH neurons (7, 9).

The central action of interleukin-6 (IL6) exerts distinct functions in the metabolism of mammals (10–12). Hypothalamic IL6 has been implicated in controlling energy expenditure (13) and food consumption (10, 11). The metabolic improvements induced by physical exercise require the hypothalamic IL6 action in obese rodents (14). Here, we hypothesized that hypothalamic IL6 might control skeletal muscle metabolism. Thus, we sought to determine the existence of a neuromuscular circuit involving hypothalamic IL6 action in the VMH nucleus toward skeletal muscle metabolism.

RESULTS

Hypothalamic IL6 stimulates fatty acid metabolism in skeletal muscle

An interaction graph using the Genotype-Tissue Expression (GTEx) human database (15) from 81 individuals revealed a positive association (red lines) between hypothalamic IL6 gene expression and lipid metabolism– and mitochondrial-related genes in the skeletal muscle (Fig. 1A). No correlation was observed between the hypothalamic IL6 gene and glucose metabolism– and protein synthesis– related genes (Fig. 1A). Hypothalamic IL6 gene expression showed a positive correlation (red lines) with lipid metabolism–related
Fig. 1. Hypothalamic IL6 induces fatty acid oxidation in muscle. (A) Interaction and (B) Pearson’s correlation graphs show the correlation between hypothalamic IL6 gene expression and several genes of human skeletal muscle. Gray line represents no correlation. Pearson’s correlation $r \geq 0.5$ and $P < 0.05$ ($n = 81$). n.s., not significant. (C) Factor loading plot: angles more than 90° between gene vectors represent a negative correlation. (D) Interaction and (E) Pearson’s correlation graphs. Pearson’s correlation $r \geq 0.35$ and $P < 0.05$ ($n = 34$). (F) PCA value was obtained using Acadsb, Cpt1c, Mrp41, Pcca, and Uqcr relative expression in the muscle of BXD mice. (G) Heatmap graph using hypothalamic and muscle transcripts from six different BXD mouse strains. (H) Factor loading plot (biplot) by using BXD cohort. (I) Experimental design using male C57BL6/J mice. (J) Oxygen consumption during 3 hours after recombinant IL6 intracerebroventricular (icv) microinjection ($n = 11$ to 14). (K) ACC and AMPK phosphorylation in the soleus muscle mice ($n = 5$). (L) Cpt1β and Ppars mRNA in the soleus muscle 3 hours after recombinant IL6 intracerebroventricular microinjection ($n = 6$ to 10). (M) Palmitic acid oxidation assay in the soleus muscle in mice intracerebroventricularly injected with IL6 ($n = 6$ to 10). (N) Transcription amount (normalized). (O) One-way analysis of variance (ANOVA) was used for statistical analysis in (M). *$P < 0.05$, **$P < 0.01$, ***$P < 0.001$, and ****$P < 0.0001$ versus saline.
genes (Fig. 1B, top panels) and a negative correlation (green lines) with lipid biosynthesis–related genes (Fig. 1B, bottom panels). These data were confirmed by a two-factor analysis (Fig. 1C).

We next used analyses using system data from a mouse genetic reference population derived from crosses of C57BL/6j and DBA/2j inbred strains, named BXD (16), geared toward multiscalar integration of traits. This dataset also showed positive interaction between hypothalamic Il6 gene expression and gene sets involved in fatty acid oxidation in the skeletal muscle (red lines). Conversely, negative interaction (green lines) was observed between hypothalamic Il6 gene expression and expression of some markers of lipid biosynthesis in the skeletal muscle of BXD mice (Fig. 1D, red and E). There was no interaction between hypothalamic Il6 gene expression and glucose metabolism– or protein synthesis–related genes (Fig. 1D).

Principal components analysis (PCA) confirmed a strong positive correlation between hypothalamic Il6 levels and lipid oxidation– and mitochondrial-related genes in the muscle (Fig. 1F). The heat-map graph revealed that specific strains of BXD mice with low levels of hypothalamic Il6 gene expression (BXD63, BXD90, and BXD65a) displayed high levels of transcripts related to skeletal muscle lipid biosynthesis (Fig. 1G). Conversely, BXD families with high levels of hypothalamic Il6 gene expression (BXD43, BXD99, and BXD84) present high lipid oxidation– and mitochondrial-related genes in the skeletal muscle (Fig. 1G). The two-factor loading plots confirmed that the hypothalamic Il6 gene expression is positively correlated with fatty acid oxidation markers (Acadsb, Cpt1c, Mprf41, Pcca, and Uqrc2) and showed a negative correlation with markers of lipid biosynthesis (Sreb1c and Sdci) in the skeletal muscle of strains of genetically diverse BXD mice (Fig. 1H).

Beyond the genetic variation and correlational analyses, we tested our hypothesis experimentally using intracerebroventricular mouse recombinant Il6 microinjection in C57BL6j/mice (Fig. 1I). Recombinant IL6 (200 ng) increased oxygen consumption (Fig. 1L). Recombinant IL6 intracerebroventricular injection increased AMPK phosphorylation in the soleus muscle of both wild-type and TLR4KO mice (Fig. S1E), confirming the specificity of IL6 biological effect in this neuromuscular circuit.

**Central IL6 enhances the muscle lipid metabolism through hypothalamic ERK stimulation**

Previous studies have shown a functional connection between VMH and muscle metabolism (7, 8). In addition, extracellular signal–regulated kinase 1/2 (ERK1/2) in VMH elicits signals from the central nervous system toward the skeletal muscle (20). Curiously, IL6 can induce ERK1/2 phosphorylation in neuronal cells (21, 22). Thus, we monitored IL6/ERK signaling in our model. We found the presence of IL6 receptor (IL6R) in different hypothalamic nuclei, particularly in the arcuate and ventromedial nuclei (Fig. S1, F and G). Next, recombinant IL6 was injected directly into VMH of mice on one side, and a vehicle was injected into the contralateral VMH of the same animal. This experimental approach demonstrated IL6 injection–induced ERK1/2 Thr202/Tyr204 phosphorylation in VMH of mice (Fig. 2A). This finding was confirmed by Western blotting analysis in hypothalamus and isolated VMH samples from IL6 intracerebroventricularly injected mice (Fig. 2, C and D).

Confocal microscopy revealed positive cells for ERK1/2 phosphorylation in IL6Rα–expressing neurons in the VMH of mice (Fig. S2A). We found several phospho-ERK (pERK)–positive stainings in neurons of VMH after recombinant IL6 injection (Fig. 2B). In addition, recombinant IL6 stimulated ERK1/2 phosphorylation in Neuro2a (N2A) cells (Fig. 2E). Together, our data showed the potential of IL6 in stimulating ERK1/2 phosphorylation in neuronal cells.

To test whether the hypothalamic IL6/ERK1/2 axis is involved in skeletal muscle lipid metabolism, PD98059, a mitogen–activated protein kinase (MAPK) kinase (MEK)/ERK signaling pharmacological inhibitor, or saline was intracerebroventricularly administered 30 min before the intracerebroventricular microinjection of recombinant IL6, as detailed in the experimental design (Fig. 2F). Notably, PD98059 abolished hypothalamic IL6-induced ACC and AMPK phosphorylation (Fig. 2G) and fatty acid oxidation (Fig. 2H) in the soleus muscle. We also evaluated the interaction between hypothalamic ERK1/2 signaling and lipid metabolism in the skeletal muscle of humans. The interaction and heatmap graphs confirmed a consistent association between hypothalamic ERK1 and ERK2 gene expression and several genes involved in fatty acid metabolism in human skeletal muscle (Fig. 2, I to K), but not glucose metabolism or protein synthesis (Fig. 2I). Together, bioinformatics and experimental data demonstrated that the hypothalamic IL6/ERK axis controls AMPK/ACC signaling and fatty acid oxidation in the soleus muscle.

**Hypothalamic IL6 stimulates the adrenergic signaling toward the skeletal muscle**

Next, we investigated the involvement of the sympathetic nervous system in this neuromuscular circuit. It has been shown that hypothalamic ERK signaling plays a critical role in controlling sympathetic flow (23, 24). The in silico analysis confirmed that hypothalamic IL6 gene expression is strongly associated with firing rate–related transcripts in mice and humans (Fig. S2, B and C). Next, the denervation experiment was performed. The left hindlimb of mice was denervated, and the contralateral paw was used as control. Two days after surgeries (cannula implant and denervation), the animals received mouse recombinant IL6 (200 ng) intracerebroventricularly
Fig. 2. Evaluation of hypothalamic IL6/ERK axis. (A) Phospho-ERK1/2\(^{Thr202/Tyr204}\) staining in VMH 30 min after vehicle or recombinant IL6 injections. Scale bar, 100 \(\mu\)m. (B) pERK1/2 (red) staining in neurons (green) in VMH (n = 4). Scale bars, 100 \(\mu\)m. Top panels, digital zoom (dZoom). White arrowheads indicate double staining. Scale bars, 10 \(\mu\)m. (C) pERK1/2\(^{Thr202/Tyr204}\) 30 min after recombinant IL6 intracerebroventricular microinjection (200 ng) in the whole hypothalamus (n = 5) and (D) in the ventromedial nucleus (n = 5 to 6). (E) pERK1/2\(^{Thr202/Tyr204}\) in N2A cells after recombinant IL6 incubation (50 ng/ml) for 60 min (n = 5). (F) Schematic view of the experiment. (G) ACC\(^{Ser79}\) and AMPK\(^{Thr172}\) phosphorylation in the soleus muscle (n = 4). (H) Palmitic acid oxidation in the soleus muscle (n = 8 to 15). (I) Interaction and (J) heatmap graphs show the positive correlation between hypothalamic ERK1/2 gene expression and genes related to lipid, glucose, and protein metabolism in the skeletal muscle of humans. Pearson’s \(r \geq 0.35\) and \(P < 0.05\). n = 81. (K) Factor loading plot shows that hypothalamic ERK1/2 gene expressions positively correlate with genes related to fatty acid oxidation in the human muscle as seen on a factor loading plot, where angles less than 90° between gene vectors represent a positive correlation. Unpaired \(t\) test was used in (C) to (E). One-way ANOVA was used for statistical analysis in (G) and (H). *\(P < 0.05\), **\(P < 0.01\), and ***\(P < 0.001\) versus saline and #\(P < 0.05\) and ###\(P < 0.001\) versus IL6.
microinjection, and the soleus muscles were removed, as demonstrated in the experimental design (Fig. 3A). The surgical denervation reduced the ability of the central action of IL6 to phosphorylate ACC and AMPK in the soleus muscle of mice (Fig. 3B), demonstrating that the hypothalamic IL6 requires an intact autonomic nervous system to alter muscle metabolism.

Notably, \( \alpha \)-adrenergic signals mediate the ACC phosphorylation and fatty acid oxidation in the skeletal muscle in response to intrahypothalamic injections of leptin (9) or the synthetic inhibitor of fatty acid synthase C75 (25). On the basis of this information, we sought to determine the involvement of \( \alpha \)-adrenergic signaling in this neuromuscular circuit. The interaction graph revealed that the hypothalamic IL6 gene is positively correlated with the expression of gene set related to \( \alpha \)-adrenergic signaling in the muscle of humans, including \( ADRA2C \), \( CREM \), \( ADCY3 \), \( PDE4B \), \( PDE4D \), and \( AK4 \) genes (Fig. 3C). These data were also confirmed by
Pearson’s correlations (Fig. 3D) and two-component analysis (Fig. 3E). In the BXD cohort, we found that α2 adrenoceptor subunits (Adra2c, Adra2a, and Adra2b), but not α1 subunits, were highly expressed in the skeletal muscle of strains with high expression of hypothalamic Il6 gene expression (fig. S3, A to C).

To investigate whether α2 adrenoceptor subunits are involved in the control of muscle metabolism upon intrahypothalamic IL6 infusion, mice lacking both α2A and α2C adrenoceptors [α2AC knockout (KO)] were generated. Wild-type and α2AC KO mice received an intracerebroventricular microinjection of recombinant IL6, and the soleus muscles were removed 3 hours later for analyses (Fig. 3F). The central microinjection of recombinant IL6 stimulated ACC and AMPK phosphorylation in the muscles of wild-type, but not α2AC double KO, mice (Fig. 3G). Furthermore, recombinant IL6 intracerebroventricular administration failed to stimulate fatty acid oxidation in the skeletal muscle of α2AC KO mice (Fig. 3H). There was no difference in the IL6 serum levels between wild-type and α2AC KO mice or after intracerebroventricular IL6 microinjection (Fig. 3I). Together, these data demonstrate that α2A and α2C receptor subunits are involved in the terminal sympathetic flow in muscle fibers, integrating the peripheral actions of hypothalamic IL6.

**Exercise stimulates central IL6 production and hypothalamic ERK activation and induces prolonged fatty acid oxidation in the skeletal muscle**

After that, we sought to evaluate this neuromuscular circuit under physiological conditions. It has been demonstrated that a single bout of exercise can underpin subsequent resting lipid oxidation for many hours in humans (26–28). Intriguingly, high levels of IL6 were observed in the hypothalamus of exercised fish (29) and mice (14), and IL6 is also produced in the central nervous system of humans in response to exercise (30). Thus, we monitored the contribution of this neuromuscular mechanism in the control of fatty acid oxidation in the skeletal muscle after exercise.

We evaluated the central and peripheral IL6 production and fatty acid metabolism in the skeletal muscle after the acute swimming protocol, as illustrated (Fig. 4A). High levels of Il6 mRNA were found in the hypothalamus immediately (0 hours) after the acute

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**Fig. 4. Exercise stimulates hypothalamic IL6/ERK axis.** (A) Experimental design. (B) Hypothalamic Il6 gene expression immediately after the exercise protocol (n = 5, *P < 0.05 versus Rest). (C) Il6 gene expression in the gastrocnemius muscle after the exercise protocol (n = 5 to 6, *P < 0.05 versus Rest). (D) Serum levels of IL6 (n = 7 to 8, ***P < 0.001 versus Rest). pERK1/2Thr202/Tyr204 in (E) hypothalamus (n = 4) and (F) VMH (n = 5). (G) pERK staining (red) in VMH of mice (n = 4). Scale bars, 100 μm. (H) pERK (red) and NeuN (green) double staining in VMH of exercised mice (n = 5). Scale bars, 100 μm. dZoom from the orange rectangles highlighting double staining in neuronal (white arrows) and nonneuronal cells (yellow arrows) (n = 5). Scale bars, 10 μm. Unpaired t test was used in (B) and (D) to (F). One-way ANOVA was used for statistical analysis in (C). *P < 0.05, ***P < 0.001, and ****P < 0.0001 versus Rest.
after the exercise session, and the muscle samples were analyzed.

This swimming protocol showed that peripheral IL6 was produced later in comparison to hypothalamus, given that the IL6 mRNA in the gastrocnemius muscle was significantly increased 120 min (2 hours) after the acute exercise (Fig. 4C). No difference was found in Il6 mRNA levels in the soleus muscle of exercised mice (Fig. S4D). No changes were found in IL6 serum levels in the first 2 hours after the exercise protocol (Fig. S4E). IL6 serum levels were significantly detected 3 hours after the exercise (Fig. 4D).

Consistent with the hypothalamic IL6 production in response to exercise, ERK1/2Thr202/Tyr204 phosphorylation was also increased immediately after exercise in the hypothalamic tissue (Fig. 4E), including in VMH (Fig. 4F). Immunofluorescence assay confirmed the presence of ERK1/2Thr202/Tyr204 phosphorylation in VMH of exercised, but not control, mice (Fig. 4G). Furthermore, several positive pERK cells were found in neurons of VMH in exercised mice, as indicated by white arrows (Fig. 4H). We also detected the presence of pERK-positive cells in nonneuronal cells of VMH in response to exercise, as indicated by yellow arrows (Fig. 4H).

In terms of fatty acid metabolism in the skeletal muscle in response to exercise, AMPK and ACC phosphorylation were still increased in the soleus muscle 3 hours after exercise (fig. S4F). RNA sequencing analysis was performed to confirm the presence of the oxidative profile in the skeletal muscle 3 hours after the acute exercise. Several genes involved in the mitochondrial function, lipid metabolism, and adrenergic signaling were up-regulated (fig. S4G), whereas, among 99 genes that were up-regulated in response to exercise, almost half of these genes are related to the mitochondrial activity (24 genes), lipid metabolism (15 genes), and adrenergic signaling (7 genes) (fig. S4, G and H).

Swimming exercise protocols are typically used as stressful stimuli for mice (31). IL6-null mice (IL6KO) were submitted to the same swimming exercise protocol to address the interference of swimming-induced stress in this neuromuscular mechanism. In contrast to wild-type mice, the swimming exercise protocol failed to stimulate ERK1/2 phosphorylation in the hypothalamus and flipped to sustain ACC and AMPK phosphorylation 3 hours after exercise in the muscle of IL6KO mice (fig. S5, B and C), confirming that endogenous IL6 levels, but not stress-associated factors, are required to elicit this neuromuscular response. Beyond the swimming exercise, we also monitored the central levels of IL6 and AMPK phosphorylation after a moderate-intensity acute treadmill running protocol (fig. S5D). Like the swimming exercise, treadmill running increased hypothalamic IL6 accumulation (fig. S5E) and induced prolonged AMPK phosphorylation in the soleus muscle (fig. S5F), reinforcing the activation of this neuromuscular system under different exercise models. These data collectively demonstrated that the hypothalamic IL6/ERK axis plays a critical role in controlling muscle metabolism under physiological conditions.

Hypothalamic IL6 controls muscle fatty acid oxidation after exercise

To explore the relevance of IL6 action specifically in the hypothalamus, exercised mice received anti–IL6 antibody (IL6 AB) intracerebroventricular microinjections 30 min before and immediately after the exercise session, and the muscle samples were analyzed 3 hours later, as illustrated in the experimental design (fig. S6A). This experiment revealed that anti–IL6 AB intracerebroventricular microinjections abolished fatty acid oxidation after exercise in the skeletal muscle (fig. S6B).

We confirmed these preliminary findings by using a versatile tool for genome editing, targeting IL6R specifically in VMH of mice using lentiviral vector [short hairpin RNA (shRNA)] microinjection. Five distinct lentivirus clones targeting the hypothalamic IL6R were tested. The TRCN94 lentivirus was the most efficient, reducing hypothalamic IL6R protein content by approximately 40% compared to a nontarget scramble vector (fig. S7A). Next, bilateral microinjections of TRCN94 were then performed to deliver IL6R lentivirus specifically in VMH (IL6R KD VMH). Three days after the lentivirus transduction, food consumption was normalized, and no difference was found in food intake between scramble and IL6R KD VMH mice (fig. S7B). We confirmed that this lentiviral transduction was restricted to VMH, reducing the IL6R content in VMH without affecting the presence of this receptor in adjacent hypothalamic nuclei, including in the arcuate nucleus (Fig. 5, A to D). Seven days after lentivirus microinjection in VMH, the animals were submitted to an acute exercise session. The double-staining assay revealed the abundance of pERK1/2-positive cells in neuronal cells of VMH in the scramble group submitted to acute exercise (Fig. 5E, second panel). However, exercise did not promote the same effect in IL6R KD VMH mice (Fig. 5E, last panel). Quantitative analysis considering pERK1/2-positive cells in neuronal and nonneuronal cells restricted to VMH demonstrated that, while exercise increased pERK1/2-positive cells in hypothalamic cells of scramble group, very few positive cells were found in VMH of IL6R KD VMH exercised mice (Fig. 5F).

High levels of AMPK and ACC phosphorylation were observed 3 hours after exercise in the soleus muscle of control (scramble), but not IL6R KD VMH, mice (Fig. 5G). Last, mass spectrometry analysis was performed 3 hours after the exercise protocol and confirmed that exercise markedly reduced the amount of fraction of saturated, monounsaturated, and polysaturated fatty acid fractions in the skeletal muscle of control (scramble), but not IL6R KD VMH, mice (Fig. 5, H to J, and fig. S7C).

Together, our findings demonstrate that the hypothalamic IL6/ERK axis in VMH drives fatty acid metabolism in the skeletal muscle (fig. S7D), and this neuromuscular circuit seems to be critical to underpin fatty acid oxidation in the skeletal muscle after exercise.

DISCUSSION

We reported that hypothalamic IL6/ERK signaling induces fatty acid oxidation in the skeletal muscle. We demonstrated that ERK signaling in VMH mediates IL6-induced AMPK and ACC phosphorylation and fatty acid oxidation in oxidative muscles. Evidence for the involvement of the α-adrenergic system in the connection of the central action of IL6/ERK signaling to fatty acid metabolism in the skeletal muscle was reported. Notably, we demonstrated that α2A and α2C receptors play a crucial role in this physiological mechanism (fig. S7D). The physiological relevance of these findings was observed in fatty acid metabolism in the skeletal muscle after exercise.

Several studies have demonstrated that the central or peripheral action of IL6 controls energy expenditure and lipid metabolism in animals and humans (13, 32–34). Wallenius and colleagues (13) reported that central, but not peripheral, IL6 injection increased energy expenditure in mice. Moreover, the microinjection of recombinant
Fig. 5. Hypothalamic IL6 controls muscle fatty acid oxidation after exercise in mice. (A) Illustration for VMH microinjections and a representative picture of a coronal view of a mouse brain, demonstrating the blue dye staining confirming the anatomical localization of bilateral microinjections in VMH (bottom). KD, knockdown. (B) IL6R (red) in the arcuate nucleus and VMH 7 days after VMH lentivirus injection (n = 5 to 6). Scale bars, 100 μm. (C) dZoom from the orange rectangles highlighted in (B) shows the presence of ΔIL6R (red). Scale bars, 50 μm. DAPI, 4′,6-diamidino-2-phenylindole. (D) IL6R-positive cells in the arcuate and ventromedial nuclei (n = 5 to 6, **P < 0.01 versus scramble). (E) Top panels: Colocalization of ERK1/2 phosphorylation (red) in neurons (green) of VMH (n = 3 to 5). Scale bars, 50 μm. Bottom panels: dZoom from the orange rectangles highlighted in the top panels shows the presence of pERK1/2 in neuronal cells. Scale bars, 10 μm. (F) Positive cells for pERK1/2 in neuronal and nonneuronal cells in VMH of mice (n = 6, *P < 0.05 versus scramble at rest). (G) ACC and AMPK phosphorylation in the soleus muscle (n = 3, *P < 0.05 versus scramble at rest). (H to J) Determination of fatty acid fractions in the gastrocnemius muscle (n = 4 to 6). Unpaired t test was used in (D) and (F). One-way ANOVA was used for statistical analysis in (G) and (H).
IL6 into the lateral cerebral ventricle increased the body temperature in rats (35). The IL6-deficient mouse displayed lower oxygen consumption during exercise (36). The present study demonstrated that recombinant IL6 intracerebroventricular injection increased oxygen consumption and fatty acid oxidation in the soleus muscle of mice. These findings collectively illustrate the critical role of the central IL6 in the control of energy homeostasis and peripheral metabolism.

VMH neurons have been extensively associated with skeletal muscle metabolism. For instance, the central action of orexin A in VMH induced glucose uptake in the skeletal muscle of mice (7). The stimulation of the hypothalamic ERK1/2 pathway has also been involved in controlling glucose metabolism in the skeletal muscle. ERK and its upstream kinase MEK regulate the leptin-induced increase in glucose uptake and insulin sensitivity in oxidative fibers in the skeletal muscle of mice through activation of the melanocortin system in VMH (20). Our study demonstrated that specific ERK1/2 activation in VMH through the IL6 action induced AMPK and ACC phosphorylation and fatty acid oxidation in the soleus muscle of mice. Beyond VMH, IL6 also plays a critical role in controlling metabolism by acting in other hypothalamic nuclei. For example, in the arcuate nucleus, IL6 action is associated with adiposity control in mice (37). Moreover, IL6 activity in the paraventricular nucleus (PVN) induces several beneficial metabolic effects even under conditions of leptin resistance (10). These studies strongly suggest that hypothalamic IL6 controls various metabolic effects by acting in specific hypothalamic nuclei.

In our study, we found that physical exercise stimulated ERK phosphorylation in neuronal and nonneuronal cells in the hypothalamus of mice. Consistent with these data, positive pERK neurons were detected in different hypothalamic nuclei of wild-type, but not IL6-deficient, mice after a stressful stimulus (38). These data establish that hypothalamic IL6/ERK signaling activation can be observed in different physiological conditions. However, it is essential to mention that IL6 can induce ERK1/2 phosphorylation of neuronal cells through IL6 trans-signaling (21, 22). The IL6 trans-signaling pathway occurs when the IL6 and soluble IL6R (sIL6R) complex (IL6/sIL6R) activates glycoprotein 130 (gp130) (39). IL6 trans-signaling mediates several metabolic effects of IL6 in peripheral tissues and the central nervous system (10, 39). For instance, Timper and colleagues (10) reported that IL6 controls energy and glucose homeostasis through gp130 expression in the PVN of obese animals, independently of IL6R. However, the role of IL6 trans-signaling was not monitored in the present study.

Hormonal and neural factors mediate the interaction between the central nervous system and muscle metabolism and function. In this context, the sympathetic outflow plays an essential role in the control of the activity of peripheral tissues, such as adipose and hepatic tissue and skeletal muscle (7, 40, 41). It has been shown that denervation of the sciatic, femoral, and obturator nerves abolishes the increase in AMPK phosphorylation in the skeletal muscle observed in intracerebroventricular leptin-injected mice (9). Our results demonstrated that the hypothalamic IL6/ERK axis induced fatty acid oxidation in the skeletal muscle through sympathetic outflow and the α2-adrenergic pathway, involving the α2A and α2C receptors. These findings agree with results reported by others since the stimulation of the α-adrenergic pathway, but not the β-adrenergic pathway, is involved in AMPK activation in the skeletal muscle (9). However, the precise mechanism by which α2A and α2C receptors induce AMPK and ACC phosphorylation in the skeletal muscle is unclear.

We and others previously demonstrated that acute exercise promoted a prolonged AMPK and ACC phosphorylation in the skeletal muscle of rodents (42–44). Similar data were found in different studies involving exercised humans (45–47). In addition, a single bout of exercise can underpin subsequent resting lipid oxidation for many hours in humans (26–28). These findings seem to be strongly associated with the physiological phenomenon informally called “afterburn.” Human studies showed that AMPK/ACC signaling remained activated in the skeletal muscle for 60 and 120 min (47, 48). Our results demonstrated that both IL6 intracerebroventricular injection and exercise increase AMPK and ACC phosphorylation and fatty acid oxidation in the skeletal muscle of mice up to 3 hours afterward. Curiously, tocilizumab, an IL6R blocker, induces abnormal lipid metabolism accumulation and disrupts fatty acid mobilization in the adipose tissue of lean and obese exercised subjects (49, 50). Together with our findings, these data demonstrate that targeting IL6 signaling may disturb the lipid metabolism affecting metabolic health.

Previously, we demonstrated that different types of acute exercise, such as swimming and treadmill running, increased the Il6 mRNA levels in the hypothalamus of mice (14) Similarly, stress protocols up-regulated Il6 mRNA expression in rat hypothalamus (38, 51). In addition, a high capacity of hypothalamic microglial cells of mice to produce IL6 under several acute or chronic circumstances has been demonstrated (52). Furthermore, human brain IL6 production was confirmed after acute prolonged exercise (30). Collectively, these studies strongly suggest the ability of hypothalamic cells to produce IL6 under physiological and pathophysiological conditions. Regarding the source of IL6 in exercised mice, a time-course study revealed the rapid accumulation of Il6 mRNA in the hypothalamus, confirming the ability of hypothalamic cells to produce IL6 in response to exercise. Recently, an interesting study using in situ hybridization method reported that IL6 could be produced in neurons, astrocytes, and microglia in the lateral parabrachial nucleus of mice (12). In the present study, we demonstrated that exercise rapidly increased Il6 mRNA levels in the hypothalamus and stimulated ERK phosphorylation in neuronal and nonneuronal cells; however, we did not assess which cell type is responsible for the hypothalamic production of IL6 in response to exercise. In addition, beyond the central production, we may consider that muscle-derived IL6 can also reach the central nervous system.

Last, our study demonstrated that the hypothalamic IL6/ERK axis increases AMPK and ACC phosphorylation and stimulates fatty acid oxidation in the skeletal muscle. Pharmacological, physiological, and genetic approaches revealed that the α-adrenergic system mediates the communication between the hypothalamic IL6/ERK1/2 axis and AMPK/ACC signaling in the muscle. The findings point out that the reported neuromuscular circuit plays an important role in regulating fatty acid metabolism in the skeletal muscle.

MATERIALS AND METHODS

N2A cells

N2A cells, derived from mouse neurons, were cultivated in Dulbecco’s modified Eagle’s medium (DMEM)/HamF12 (1:1) supplemented with 10% (v/v) fetal bovine serum (FBS) and streptomycin/penicillin (100 U/ml). Cells were cultivated at 37°C in a humidified atmosphere containing 5% CO2. N2A cells were subjected to neurite differentiation before treatments by incubation with medium containing...
A plantation was aimed at the third ventricle of mice weighing 25 to 30 g of 14 days of age. After cranial trichotomy and antisepsis, a 1-cm intraparietal incision was performed at the stereotaxic coordinates, and a 1-cm intraparietal incision was confirmed by a positive drinking response after administration of angiotensin II (40 ng); animals that did not drink water after angiotensin injection were not included in the experiment.

**Intracerebroventricular microinjections**

**Recombinant IL6**

Intracerebroventricular IL6 mouse recombinant microinjection (200 ng) from Sino Biological (50136-MNAE) was performed between 5:00 and 6:00 p.m., as previously described (14).

**IL6 neutralizing antibody**

Animals were randomly selected for intracerebroventricular microinjection containing saline or rabbit antiserum against IL6 (IL6 AB). Anti–IL6 AB (100 ng) from Santa Cruz Biotechnology was performed 30 min before and immediately after the exercise protocol, as previously published (14).

**PD98059**

Mice were randomly selected for intracerebroventricular microinjection containing saline or PD98059 (60 μM), a selective, cell-permeable inhibitor of the MEK, as previously described (55). PD98059 was performed 30 min before the intracerebroventricular recombinant IL6 microinjection.

**VMH IL6R lentivirus transfection**

Five different shRNA-based lentiviral clones [The RNAi Consortium (TRC); titer, 10^{6}] targeting IL6R [TRCN 0000057 (LV1), TRCN 0000059 (LV2), TRCN 0000089 (LV3), TRCN 0000093 (LV4), and TRCN 0000094 (LV5)] or scramble (SHC 016 V, pLKO.1-puro nonmammalian shRNA control) (SCR) from Sigma-Aldrich (St. Louis, MO, USA) were tested for IL6R knockdown in preliminary experiments, as previously described (56).

**Reagents and antibodies**

Reagents for SDS–polyacrylamide gel electrophoresis (SDS-PAGE) and immunoblotting were from Bio-Rad Laboratories (Hercules, CA). Tris, aprotinin, adenosine triphosphate (ATP), dithiothreitol, phenylmethylsulfonyl fluoride, Triton X-100, Tween 20, glycerol, and bovine serum albumin (BSA) (fraction V) were from Sigma Chemical Co. (St. Louis, MO). Nitrocellulose paper (BA85, 0.2 mm) was from Schleicher & Schuell (Keene, NH). Ketamine hydrochloride was from Cristália (Itapira SP, Brazil). The chemiluminescent kit was from Bio-Rad Laboratories (Hercules, CA).

**Immunoblotting**

After the respective treatments, animals were anesthetized. Samples from the hypothalamus and soleus muscle were obtained, minced...
coarsely, and homogenized immediately in solubilization buffer containing 100 mM tris (pH 7.6), 1% Triton X-100, 10 mM Na3VO4, 100 mM NaF, 10 mM Na2P2O7, 4 mM EDTA, 150 mM NaCl, aprotinin (0.1 mg/ml), and phenylmethylsulfonyl fluoride (35 mg/ml) using a Polytron PTA 20S generator (model PT 10/35, Brinkmann Instruments, Westbury, NY) operated at maximum speed for 30 s and clarified by centrifugation. All the samples from all groups were subjected to SDS-PAGE and blotted onto a nitrocellulose membrane. The membranes were incubated for 12 hours at 4°C with primary antibody after blockade with 5% nonfat milk in tris-buffered saline with Tween 20 (TBST) (10 mM tris, 150 mM NaCl, and 0.02% Tween 20) for 90 min at room temperature. After the secondary antibody incubation, the signal was detected for 60 min in a 3% nonfat milk–TBST solution, treated with 2 ml of SuperSignal West Pico Chemiluminescent Substrate, exposed to photosensitive RX film from Kodak, or visualized using G:Box from SynGene. Band intensities were quantified by optical densitometry using UN-SCAN-IT gel 7.1 (Silk Scientific Inc.). Ponceau staining from Sigma-Aldrich (St. Louis, MO, USA) was also used to monitor the loading control for each sample.

**mRNA isolation and real-time PCR**

Total RNA was extracted using TRIzol reagent (Life Technologies), according to the manufacturer’s recommendations. Total RNA was rendered genomic DNA free by digestion with ribonuclease-free deoxyribonuclease (RQ1, Promega, Madison, WI, USA). Real-time polymerase chain reaction (PCR) and mRNA isolation were performed using a commercial kit (GAPD, 4352338E) for mouse and TaqMan gene expression assays for mouse IL6 (Mn00446190_m1), and using the following primers: Cpt1β, 5′-TCGAAATCAAGATGCTCATCCT-3′ (forward) and 5′-GAGATGTCCACCTTGCAGTAGTTG-3′ (reverse); Ppara, 5′-TTAGAGGAGAGCCAAGTTGAAGTTC-3′ (forward) and 5′-GAGATGTCCACCTTGCAGTAGTTG-3′ (reverse); Il6ra, 5′-TGCAAGCTCAGTCGATACCG-3′ (forward) and 5′-TGCTTCACTCCTGCAGAGGCAT-3′ (reverse).

**Oxygen consumption determination**

O2 consumption was measured using an indirect open circuit calorimeter (CLAMS, Oxymax Deluxe System, Columbus Instruments, Columbus, OH, USA). Mice were adapted 1 day before the experiment. Measurements were performed for 3 hours after intracerebroventricular IL6 microinjection.

**Palmitate oxidation in soleus muscle**

The animals were euthanized, and their soleus muscles were isolated and incubated as previously described (57). Briefly, mice were sacrificed by cervical dislocation, and their soleus muscles were carefully and quickly isolated, kept under resting tension with stainless steel clips, and preincubated in a buffered Krebs-Ringer bicarbonate (TKBR) containing 5.6 mM glucose (pH 7.4) for 30 min in a water bath to 35°C with 95% O2 and 5% CO2, with continuous shaking (90 oscillations/min). After this period, the muscles were transferred to other vials containing the same buffer but added with [U-14C]palmitate (0.2 μCi/ml) and 100 μM unlabeled palmitate. The latter was previously diluted in ethanol (80 mM solution) before adding the incubation buffer. NaOH solution (2 M) was added to a microtube isolated from the inside of the bottle to adsorb 14CO2. The muscles were incubated for 2 hours under the same conditions. At the end of the incubation, muscles were briefly washed in cold TBKR (4°C), dried on filter paper, frozen in liquid nitrogen, and subsequently weighed. The flask was hermetically sealed to measure the oxidation of [U-14C]palmitate as previously described (58).

**Mass spectrometry analysis for fatty acid determination**

For fatty acid profile determination, 150 μl of tissue homogenates was taken in a screw cap glass tube containing 25 μg of internal standard (tridecanoic acid, C13:0). After that, 1 ml of 0.5 M NaOH-methanol was added, and the sample was boiled at 100°C for 15 min. After cooling the samples, 2 ml of BF3-methanol was added, and the sample was boiled at 100°C for 20 s. The sample was again cooled to room temperature, and 1 ml of isoctane was added. The tubes were shaken, and 5 ml of saturated NaCl solution was added. After phase separation, the supernatant was collected and evaporated with nitrogen gas to concentrate the fatty acid methyl esters (FAMEs). FAME was resuspended in 50 μl of hexane for chromatographic analysis (59). Chromatographic analyses were performed using a gas chromatograph–mass spectrometer (model GCMS-QP2010 Ultra, Shimadzu). A fused silica capillary column (Stabilwax; length, 30 m; internal diameter, 0.25 mm; thickness, 0.25 μm; Restek, USA) was used to inject 1 μl of the sample at 250°C. High-grade pure helium (He) was used as the carrier gas with a constant flow rate of 1.3 ml/min with a split injection of 2:1. The oven temperature was programmed from 80° to 175°C at a rate of 5°C/min, followed by another gradient of 3°C/min to 230°C, which was maintained for 10 min. Mass conditions were as follows: ionization voltage, 70 eV; ion source temperature, 200°C; full scan mode in the mass range of 35 to 500 with a velocity of 0.2 s per scan.

**Immunofluorescence staining**

Ad libitum-fed animals were perfused with 4% paraformaldehyde transcardially. The brains were immersed in 10% (w/v) sucrose and, after 24 hours, transferred to 30% (w/v) sucrose. After that, they were embedded in optimal cutting temperature (OCT) compound (Sakura Finetek, Torrance, CA) and cut into 20-μm coronal sections using a cryostat. The sections were incubated in blocking solution in PBS (0.2% of Triton X-100) and 5% BSA for 2 hours at 21°C and then incubated overnight at 4°C in mouse anti-NeuN (1:300; Millipore, MAB377), anti-IL6Rα (1:200; Santa Cruz Biotechnology, sc660), anti–phospho-p44/42 MAPK (ERK1/2) (Thr 202 /Tyr 204 ) (D13.14.4E) MAB377, anti-IL6R (1:200; Santa Cruz Biotechnology, sc-393592). After washing in PBS, sections were placed in secondary donkey anti-mouse fluorescein isothiocyanate (1:500; Santa Cruz Biotechnology, sc 2010) and goat anti-rabbit Alexa Fluor 546 (1:500; Thermo Fisher Scientific, CA, USA) for 2 hours and mounted with VECTASHIELD Mounting Medium with 4′,6-diamidino-2-phenylindole (DAPI) (H-1200, Vector Laboratories Inc., Burlingame, CA, USA). The images were acquired with a confocal laser microscope (LSM 780, Zeiss, Jena, Germany). Digital/electronic zoom of highlighted areas was performed when necessary.

**Confocal microscopy**

Briefly, after fixation with 4% paraformaldehyde, the slides were washed with PBS, then incubated for 10 min with glycine (0.1 M), and treated for 30 min with 4% BSA solution (Sigma-Aldrich). The slides then were incubated overnight at 4°C with a recombinant mouse immunoglobulin G (IgG) antibody against IL6Rα (D-8, sc-274259, Santa Cruz Biotechnology) and a recombinant rabbit IgG.
antibody against phospho-p44/42 MAPK (ERK1/2, Cell Signaling Technology), both diluted in 1% BSA. The slides were washed and incubated for 2 hours with secondary antibodies (Alexa Fluor 488-conjugated, catalog number A32723, Thermo Fisher Scientific, Waltham, MA, USA; Alexa Fluor 594-conjugated donkey anti-rabbit IgG, catalog number A21207, Thermo Fisher Scientific, Waltham, MA, USA) diluted 1:500 in 1% BSA. Cells were washed and stained with DAPI (catalog number SC3598, Santa Cruz Biotechnology, Dallas, TX, USA) for nuclear labeling. Microscopic images were acquired using Zeiss LSM 880 with Airyscan on an Axio Observer 7 inverted microscope (Carl Zeiss, Jena, Germany) with a C Plan-Apochromat 63×/1.4 oil differential interference contrast objective lens with 4× optical zoom. Before visual inspection of the images, raw .czi files were automatically processed into deconvoluted Airyscan images using Zen Black 2.3 software. DAPI staining was visualized with conventional confocal imaging using a 405-nm laser line for excitation and a pinhole set to 1 Airy unit.

**Arc and VMH cell counting**
Sections from the immunofluorescence assay were used to determine ERK1/2 phosphorylation in VMH of mice transfected with IL6R shRNA (LV5) or scramble. Images were obtained, isolating only Arc or VMH in the images. Five images per group were counted blindly by two independent researchers using ImageJ software 1.48v. The average of each image was used to obtain the group mean.

**Hypothalamic VMH dissection**
The microdissections were performed according to previous studies (9, 60, 61). Fresh whole brains were dissected using a specific steel matrix for mouse brains. Slices (1 mm) were cut following the sagittal orientation. The cuts were made using ultrathin blades (6 × 10⁻³ mm thick), and the first sections from the midline of the brain were used for the microdissection of VMH/DMH (dorsomedial hypothalamus) using a microscope and following the mouse brain atlas coordinates. VMH was quickly frozen in liquid nitrogen for further protein analysis.

**Denervation experiment**
For the denervation experiments, mice were anesthetized with ketamine (100 mg) and diazepam (0.07 mg) (0.2 ml/100 g body weight). The procedure started when corneal and paw pain reflexes were abolished. Denervation and cannula implantation surgeries were performed in the same procedure. The left hindlimb muscles were carefully denervated by excising approximately 4 mm of the sciatic nerve. The animals were used in the experiment 48 hours later, and the contralateral paw was used as control.

**BXD hypothalamus microarray processing**
Correlation analyses were performed using hypothalamic mRNAs of the BXD inbred family [Hypothalamus Affy MoGene 1.0 ST (Nov10) and EPFL/LISP BXD CD Affy Mouse Gene 1.0 ST (Dec11) RMA] (16), except for fig. S2B. These datasets are accessible on GeneNetwork (www.genenetwork.org) (62). The Pearson’s and Spearman’s correlation graphs were built using Prism graph, and the heatmap graph was obtained using the GENE-E software. For fig. S2C, we accessed the normalized, log-transformed (with offset 1) RNA sequencing gene read counts in the hypothalami of 170 deceased human subjects from the GTEx project (15) version 8 through accession number dbGaP phs000424.v8.p2 (see www.gtexportal.org/home/documentationPage for a complete overview of the data and its processing pipeline). We regressed these preprocessed gene expressions in GTEx hypothalami for the human homologs of the firing rate–related genes on the IL6 preprocessed gene expressions through ordinary least squares regression. The P values associated with IL6 expression were corrected with the Benjamini-Hochberg false discovery rate procedure and reported as q values. All individual values used in bioinformatics analysis are described in supplementary tables.

**Statistical analysis**
The results were expressed as means ± SD. Immunoblot results were presented as direct comparisons between the groups. Data were analyzed by Student’s t test or one-way analysis of variance (ANOVA), as appropriate, with post hoc Bonferroni or Tukey tests for multiple unpairwise comparisons of the means. The level of significance adopted was P < 0.05. Statistica 6.0 software was used for the analysis. Pearson’s and Spearman’s correlations were used for the bioinformatics analysis, and r and P values are described in the figures or figure legends. The number of animals/samples used in each experiment is described in the figure legends.

**SUPPLEMENTARY MATERIALS**
Supplementary material for this article is available at https://science.org/doi/10.1126/sciadv.abm7355

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