Acute effects of leptin require PI3K signaling in hypothalamic proopiomelanocortin neurons in mice

Jennifer W. Hill, Kevin W. Williams, Chianping Ye, Ji Luo, Nina Balthasar, Roberto Coppari, Michael A. Cowley, Lewis C. Cantley, Bradford B. Lowell, and Joel K. Elmquist

1Division of Hypothalamic Research, Department of Internal Medicine and Department of Pharmacology, University of Texas Southwestern Medical Center, Dallas, Texas, USA. 2Division of Endocrinology, Beth Israel Deaconess Medical Center, Boston, Massachusetts, USA. 3Department of Systems Biology, Harvard Medical School, and Division of Signal Transduction, Beth Israel Deaconess Medical Center, Boston, Massachusetts, USA. 4Division of Physiology, University of Bristol, Bristol, United Kingdom. 5Division of Neuroscience, Oregon National Primate Research Center, Oregon Health and Science University, Beaverton, Oregon, USA.

Normal food intake and body weight homeostasis require the direct action of leptin on hypothalamic proopiomelanocortin (POMC) neurons. It has been proposed that leptin action requires PI3K activity. We therefore assessed the contribution of PI3K signaling to leptin’s effects on POMC neurons and organismal energy balance. Leptin caused a rapid depolarization of POMC neurons and an increase in action potential frequency in patch-clamp recordings of hypothalamic slices. Pharmacologic inhibition of PI3K prevented this depolarization and increased POMC firing rate, indicating a PI3K-dependent mechanism of leptin action. Mice with genetically disrupted PI3K signaling in POMC cells failed to undergo leptin depolarization or increased firing frequency in response to leptin. Insulin’s ability to hyperpolarize POMC neurons was also abolished in these mice. Moreover, targeted disruption of PI3K blunted the suppression of feeding elicited by central leptin administration. Despite these differences, mice with impaired PI3K signaling in POMC neurons exhibited normal long-term body weight regulation. Collectively, these results suggest that PI3K signaling in POMC neurons is essential for leptin-induced activation and insulin-induced inhibition of POMC cells and for the acute suppression of food intake elicited by leptin, but is not a major contributor to the regulation of long-term organismal energy homeostasis.

Introduction

The rising rate of obesity in Western countries has led to intensified efforts to understand the molecular and neuroanatomic mechanisms underlying coordinated control of energy balance in the face of abundant food and high-energy-density diets. One key metabolic signal is the adipocyte-derived hormone leptin. Leptin circulates in proportion to stored body fat, and lack of leptin or long-form leptin receptors in the brain produces severe obesity (1–3). One identified leptin-responsive cell is the proopiomelanocortin (POMC) neuron. POMC cells are located in the arcuate nucleus (ARC) of the hypothalamus and are directly activated by circulating leptin (4, 5). Indeed, direct leptin action on POMC neurons is required for normal body weight homeostasis, because deletion of the leptin receptor in POMC neurons alone causes increased adiposity (6).

Leptin activates a cytokine receptor, inducing JAK/STAT pathway signaling and leading to alterations in transcription of several target genes, including the gene encoding POMC (7, 8). It has been established that STAT3 signaling plays a major role in leptin’s control of energy homeostasis (9). However, many acute responses to leptin do not necessitate altered gene expression. For example, leptin has been shown to depolarize and increase POMC neuronal activity via activation of a nonselective cation current within minutes of leptin application (5). The acute modulation by leptin of cellular activity is conserved in several brain areas and suggests that neurons are sensitive to rapid changes in leptin levels (5, 10–13). Moreover, this change in cellular activity might correlate to a leptin-induced change in physiology or behavior.

Recent attention has focused on the potential role of the PI3K pathway in inducing rapid effects of leptin in hypothalamic neurons (14, 15). In numerous cell types, PI3K acts in a heterodimeric form consisting of 1 85-kDa regulatory (p85) and 1 110-kDa catalytic (p110) subunit. Under normal conditions, signaling by p85-p110 dimers is partially inhibited by an excess of p85 subunits (16). Thus, reduction of p85 levels through deletion of a single isoform can paradoxically upregulate PI3K signaling, while deletion of multiple isoforms prevents catalytic activity and eliminates signaling (17, 18). PI3K is activated by phosphorylated IRS proteins, allowing the enzyme to catalyze the phosphorylation of PtdIns(4,5)P2 (PIP2) to PtdIns(3,4,5)P3 (PIP3). PIP3, in turn, may recruit downstream molecules carrying PIP3-binding domains or modify channel activation (19, 20).

While recent work suggests that leptin activates PI3K signaling in the hypothalamus (1, 2, 14, 21–27), the role of a central leptin-PI3K pathway in short- and long-term energy regulation remains to be established. We therefore tested the hypothesis that PI3K signaling in POMC neurons is necessary for normal leptin responsiveness. We first examined the activity of POMC neurons in the presence of leptin, insulin, and PI3K antagonists. Second, using the cre/loxP system, we eliminated all of the major p85 isoforms expressed in POMC cells of the ARC (28) and investigated the resulting impact on leptin signaling and energy homeostasis.

Nonstandard abbreviations used: ACSF, artificial cerebrospinal fluid; AgRP, agouti-related protein; ARC, arcuate nucleus; NTS, nucleus of the solitary tract; PIP3, PtdIns(3,4,5)P3; POMC, proopiomelanocortin.

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Results
PI3K blockade inhibits leptin-induced activation of POMC neurons. Loose-patch recordings of POMC neurons were performed in intact mice that express GFP under the control of the POMC promoter (referred to herein as WT) (5, 29, 30). Leptin (100 nM) increased the firing rate in 12 of 13 POMC neurons (artificial cerebrospinal fluid [ACSF], 1.2 ± 0.1 Hz; leptin, 3.4 ± 0.4 Hz; n = 12, P = 0.00003; Figure 1A). Similar to previous reports, the leptin-induced increase in firing rate was reversible within approximately 15 minutes (12, 13, 31, 32). The selective PI3K antagonist wortmannin (100 nM) was used to delineate a PI3K contribution to the leptin-induced excitation of POMC neurons. Preapplication of wortmannin blocked the leptin-induced activation of all POMC cells examined (control, 0.9 ± 0.1 Hz; wortmannin and leptin, 1.1 ± 0.1 Hz; n = 6, P = 0.22; Figure 1B). This suggests that leptin directly excites POMC neurons via a PI3K-dependent mechanism.

Whole-cell patch-clamp recordings were made to assess the effects of leptin on membrane potential. In current clamp mode, leptin caused rapid depolarization from rest in 19 of 23 POMC neurons (6.2 ± 0.5 mV; resting membrane potential, −44.8 ± 0.8 mV; n = 19; Figure 2). This depolarization was accompanied by a 26% decrease in the whole-cell input resistance (control, 1,052.3 ± 202.0 MΩ; leptin, 780.2 ± 115.8 MΩ; n = 9, P = 0.008; Figure 3). Extrapolation of the linear slope conductance in control and leptin-containing ACSF revealed a reversal potential of −28.1 ± 2.3 mV (n = 9; Figure 3), which was similar to previous reports that suggested a non-selective cation conductance may be involved in the leptin-induced depolarization of hypothalamic neurons (5, 33).

Selective PI3K inhibitors were then used to determine the role of PI3K in the leptin-induced depolarization of POMC neurons. Wortmannin (100 nM) prevented the leptin-induced depolarization in 4 of 5 POMC neurons (0.7 ± 0.7 mV; n = 5; Figure 2). Similarly, the PI3K inhibitor LY294002 (10 μM) blocked the leptin-induced depolarization in 8 of 10 POMC neurons (0.5 ± 0.8 mV). LY294002 also prevented the leptin-induced decrease in whole-cell input resistance (control, 1,094.1 ± 182.0 MΩ; LY294002 and leptin, 1,015.5 ± 158.3 MΩ; n = 10, P = 0.76). In contrast, a MAPK inhibitor, PD98059 (50 μM), failed to prevent leptin-induced depolarization in 8 of 10 POMC neurons (4.8 ± 0.7 mV; n = 8), which suggests that acute activation of POMC neurons is independent of MAPK signaling. Some neurons were voltage-clamped at membrane potentials between −50 and −70 mV to monitor changes in whole-cell current during bath application of leptin. Leptin application resulted in an inward current in all WT neurons examined (−15.5 ± 3.2 pA, n = 5). In agreement with the leptin effect on membrane potential, the leptin-induced inward current was prevented by LY294002 (−3.4 ± 4.0 pA, n = 5). Together, these pharmacological data indicate that a PI3K-dependent mechanism is required for the leptin-induced activation of POMC neurons.

Development of Pik3r1 POMC KO Pik3r2−/− mice. To generate mice with impaired PI3K signaling in POMC neurons, we crossed mice in which loxP sites flank exon 7 of the Pik3r1 gene (Figure 4A). These mice were crossed with transgenic mice expressing Cre driven by POMc regulatory elements (6). By interbreeding the resulting mice with Pik3r2−/− (35) and B6:129-Gt(Rosa)26Sor (36) mice (The Jackson Laboratory), we were able to produce mice globally deficient in p85α but lacking p85α only in POMC neurons with concomitant GFP reporter expression (Pik3r1 POMC KO Pik3r2−/−). For all experiments, these mice were compared with littermates not expressing POMC-cre (Pik3r1 POMCKO Pik3r2−/−). All mice carried the ROSA26 allele.

In the rodent brain, POMC expression is limited to the ARC of the hypothalamus and the nucleus of the solitary tract (NTS) (36, 37). Cre recombine expression in this POMC-cre model has been previously shown to recapitulate this expression pattern, with greater than 90% of α-melanocyte-stimulating hormone–positive neurons able to drive reporter expression (6). We performed PCR on genomic DNA from the ARC, dorsomedial hypothalamic...
nucleus, paraventricular nucleus, NTS, and control areas of the brain using primers surrounding the targeted exon 7 region. In these animals, deletion of the 103-bp loxP-flanked region of exon 7 of Pik3r1 was limited to the ARC and the NTS (Figure 4B).

We examined whether POMC neurons developed normally in these mice. No change was seen in their number or size (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI32964DS1). Furthermore, resting membrane potential, basal firing rate, and whole-cell input resistance were not altered in POMC neurons of Pik3r1 POMC KO Pik3r2−/− mice (Supplemental Table 1). The whole-cell capacitance of normal POMC and Pik3r1 POMC KO Pik3r2−/− POMC neurons were also statistically similar. These data suggest that POMC neuronal development is not altered in Pik3r1 POMC KO Pik3r2−/− neurons.

Normal corticotroph function in Pik3r1 POMC KO Pik3r2−/− mice. Because hypercortisolism increases food intake, decreases energy expenditure, and can lead to obesity (38–40), we considered the possibility that Pik3r1 gene deletion in POMC-expressing corticotrophs might alter corticosterone release. We therefore examined the corticosterone levels in males seen under basal conditions and psychosocial-induced stress (41) in these mice. Basal levels of corticosterone did not differ between experimental mice and littermate controls (Supplemental Figure 2). We found that 30 min of social confrontation resulted in a similar increase in plasma corticosterone in Pik3r1 POMC KO Pik3r2−/− mice and littermate controls, suggesting normal corticotroph function in the experimental mice.

Targeted POMC Pik3 deletion disrupts insulin and leptin effects on neuronal activity. Insulin activates PI3K in the hypothalamus, and insulin-induced modulation of cellular activity requires PI3K in POMC cells of the arcuate (42). Therefore, we used insulin to further test whether PI3K signaling was disrupted in our mouse model. We used whole-cell patch-clamp recordings in WT mice that express GFP under the control of the POMC promoter. Similar to previous reports (43, 44), 50 nM insulin caused hyperpolarization from rest in 7 of 11 POMC neurons (−7.7 ± 1.2 mV; n = 7; Figure 5). The hyperpolarization was accompanied by a 28% decrease in whole-cell input resistance, from 1.1637 ± 77.7 M in control ACSF to 842.0 ± 45.7 M with insulin treatment (n = 7 from unidentified neurons; P = 0.004, paired t test). Extrapolation of the slope conductance revealed an estimated reversal potential of −95 ± 5 mV (n = 7) for the hyperpolarization, which is near the calculated K+ equilibrium. Similar to previous reports, application of 200 μM tolbutamide reversed the insulin-induced hyperpolarization (n = 3; Figure 5), suggesting that the insulin-induced hyperpolarization of hypothalamic neurons depends on activation of ATP-dependent K+ channel conductance (11).

PI3K signaling was then assessed in POMC neurons from Pik3r1 POMC KO Pik3r2−/− mice by examining the insulin-induced suppression of cellular activity. Insulin (50 nM) failed to influence the membrane potential in 10 of 12 POMC neurons from Pik3r1 POMC KO Pik3r2−/− mice (0.3 ± 0.4 mV; n = 10; Figure 5). Of the 2 remaining cells, insulin application resulted in a small depolarization (2 mV) in one cell and a small hyperpolarization (−2 mV) in the other. These data suggest that PI3K signaling, as measured by acute insulin-induced inhibition of cellular activity, is severely disrupted in ARC POMC neurons from Pik3r1 POMC KO Pik3r2−/− mice. Finally, we tested whether the absence of PI3K signaling in POMC neurons could prevent the observed leptin-induced excitation of cellular activity. In loose-patch recordings, leptin failed to increase unit activity in 4 of 5 POMC neurons from Pik3r1 POMC KO Pik3r2−/− mice (control, 1.3 ± 0.1 Hz; leptin, 1.6 ± 0.2 Hz; n = 5, P = 0.39; Figure 2). Leptin application also failed to influence the membrane potential in 10 of 11 POMC neurons from these mice (0.3 ± 0.2 mV; n = 11; Figure 3). Leptin application in the remaining cell resulted in a small depolarization from rest (3 mV). Moreover, whole-cell input resistance in all POMC cells examined from Pik3r1 POMC KO Pik3r2−/− mice was also unaffected by leptin (control, 1,148.1 ± 111.1 MΩ;
leptin, 1,077.6 ± 98.4 MΩ; n = 9, P = 0.18; Figure 3). Similarly, the leptin-induced inward current was not observed in POMC neurons from Pik3r1 POMCKO Pik3r2–/– mice (–3.0 ± 1.0 pA; n = 6). These data suggest that PI3K signaling is required for the leptin-induced excitation of ARC POMC neurons.

As previously described, in other cell types, deletion of a single PI3K regulatory isoform fails to suppress PI3K signaling (17, 18). Therefore, we also examined the effect of deleting p85α alone from POMC neurons (Pik3r1 POMCKO mice). The basal activity before adding leptin was similar to that seen in p85α-intact POMC neurons (Supplemental Table 1). Importantly, 100 nM leptin acutely activated POMC neurons from both WT and Pik3r1 POMCKO mice (Figure 2). There was also no significant difference between the leptin-induced depolarization and increase in firing rate of neurons from WT and Pik3r1 POMCKO mice. The contrast between the leptin responsiveness of Pik3r1 POMC neurons and Pik3r1 POMC Pik3r2–/– POMC neurons suggests that deletion of both p85β and p85α is required to disrupt PI3K activity in POMC neurons.

Targeted POMC PI3K disruption blunts responses to acute leptin administration. To directly assess leptin sensitivity in vivo, we compared the response of littermate controls and Pik3r1 POMC Pik3r2–/– mice to exogenous leptin administration. Because body fat levels influence leptin sensitivity (45), mice were matched for body weight. Following surgical implantation of guide cannulae aimed at the lateral ventricle, male mice (n = 6–7 per group) from each genotype were fasted overnight and injected i.c.v. with PBS vehicle or 6 μg leptin. As shown in Figure 6, in control mice i.c.v. injection of leptin caused a significant reduction in food intake at 3 h after injection (P = 0.04) and body weight at 24 h after injection (P = 0.002) compared with saline-treated controls. However, in Pik3r1 POMCKO Pik3r2–/– mice, i.c.v. leptin did not significantly affect food intake or body weight compared with vehicle. These results demonstrate that leptin-mediated inhibition of feeding is blunted in the absence of p85α and p85β in POMC.

Energy homeostasis, fertility, and linear growth. Because leptin levels are generally increased in models of impaired leptin signaling, we measured leptin levels in Pik3r1 POMC Pik3r2–/– mice and littermate controls. Impairment of PI3K signaling in POMC cells did not affect circulating leptin levels. To further examine the effect of the targeted p85 deletion on long-term energy homeostasis, we examined body weight and food intake in Pik3r1 POMC Pik3r2–/– mice. Deletion of both α and β subunits in POMC cells failed to influence long-term body weight regulation in males or females fed high-fat chow (Figure 7A). Food intake also did not differ between the groups (Figure 7B). To rule out the possibility that an increased adiposity phenotype was masked by a decline in lean mass, body composition was assessed in diet-induced obese mice using dual-energy X-ray absorptiometry. As shown in Figure 7C, fat percentage and lean mass were unaffected. Metabolic rate and respiratory quotient were similar between control and Pik3r1 POMC Pik3r2–/– mice (Figure 8, A and C). Finally, hypothalamic POMC, NPY, and agouti-related protein (AgRP) mRNA levels were measured in these animals; no significant difference in expression levels was seen compared with controls (Figure 8B). Comparable expression levels were seen in mice 6–8 weeks of age (Supplemental Figure 3). These data demonstrate that PI3K signaling in POMC neurons is not required for long-term energy homeostasis.

In addition to regulating energy balance, leptin is also known to affect hypothalamic control of ovulation and body length (46–49). However, Pik3r1 POMC Pik3r2–/– mice were similar to littermate controls in terms of pups per litter (Pik3r1 POMC Pik3r2–/–, 6.1 ± 0.8, n = 13; control, 6.7 ± 0.5, n = 14), average time to conception (Pik3r1 POMC Pik3r2–/–, 5.6 ± 2.6 d, n = 14; control, 7.2 ± 2.4 d, n = 15), and body length (Pik3r1 POMC Pik3r2–/– males, 9.5 ± 0.1 cm,
These data suggest that, at least in POMC neurons, the PI3K pathway is not required for fertility or linear growth.

**Discussion**

Selective deletion of leptin receptors in POMC neurons results in hyperleptinemia and causes moderate long-term weight gain (6). Whether this obese phenotype is dependent upon altered neuronal excitability or altered gene transcription was not clearly established previously. Moreover, while leptin directly depolarizes POMC neurons in the ARC (5), the cellular mechanism responsible has been a subject of debate. Previous studies showed that leptin activation of protein kinases (e.g., PI3K and MAPK) and subsequent phosphorylation of target proteins occur rapidly and therefore might regulate membrane potential (14, 50). Our present results demonstrated that leptin acutely activates POMC neurons via a PI3K mechanism. In addition, we demonstrated that the POMC PI3K pathway contributes to the acute effects of exogenous leptin on feeding. These results are in accord with previous studies that have demonstrated abolition of leptin’s suppression of feeding following PI3K inhibitor administration (25, 27).

While PI3K impairment in POMC neurons had striking, short-term physiological consequences, the absence of POMC PI3K signaling had no detectable impact on long-term body weight homeostasis. These results are consistent with evidence that other intracellular signaling pathways, such as those targeting the transcription factor

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**Figure 4**

Deletion of exon 7 of Pik3r1 in genomic DNA from hypothalamic nuclei of Pik3r1 POMCKO Pik3r2–/– mice. (A) The 1,275-bp region between primer binding sites becomes 298 bp in length when exon 7 is excised by cre recombinase. (B) Amplified genomic DNA from a Pik3r1 POMCKO Pik3r2–/– and littermate control (Con) mice. Tail DNA from a Pik3r1–/– mouse is shown as a positive control (+). Off, olfactory bulb DNA; HYP, DNA from whole hypothalamus; DMH, dorsomedial hypothalamic nucleus; PVN, paraventricular nucleus.

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**Figure 5**

Insulin hyperpolarizes POMC neurons via a PI3K-dependent mechanism. (A) Current-clamp record depicting the hyperpolarization of a POMC neuron from rest by 50 nM insulin. The hyperpolarization was reversible within 15 min of wash to control ACSF. Downward deflections are responses to rectangular current steps. (B) Current-clamp recording at resting membrane potential showing insulin-induced hyperpolarization (50 nM) followed by tolbutamide (200 μM) blockade of the insulin effect. (C) Sample trace illustrating the absence of insulin-induced hyperpolarization in POMC neurons from Pik3r1 POMCKO Pik3r2–/– mice. (D) Insulin-induced responses of identified POMC neurons from WT and Pik3r1 POMCKO Pik3r2–/– mice. P = 0.02, Pik3r1 POMCKO Pik3r2–/– versus WT.

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$n = 6$; control males, $9.7 \pm 0.2$ cm, $n = 8$). These data suggest that, at least in POMC neurons, the PI3K pathway is not required for fertility or linear growth.
STAT3, mediate many of leptin’s actions on energy homeostasis. For example, Bates and colleagues demonstrated that disruption of leptin activation of STAT3 by a targeted mutation of the leptin receptor causes dramatic hyperphagia and decreased energy expenditure, resulting in profound obesity similar to \( \text{db/db} \) mice (9). In addition, Xu et al. recently demonstrated that inactivating STAT3 in POMC neurons of female mice causes increased body weight and fat mass and reduces hypothalamic POMC expression (51). Interestingly, the authors also found that the responsiveness of these mice to acute leptin administration remained intact. While the precise role of IRS isoforms in leptin signaling via PI3K in POMC neurons has not been established, mice lacking \( \text{Irs2} \) in POMC cells do not display increased body weight, hyperleptinemia, or hyperphagia (43). Thus, our data are consistent with several findings demonstrating a role for both PI3K-dependent and -independent pathways in the actions of leptin in POMC neurons.

Several recent papers have demonstrated that proximity of the FoxO1 and STAT3 binding sites on the POMC promoter permits competition between the 2 factors, with FoxO1 inhibiting STAT3 stimulation of POMC gene transcription (42, 52). In addition, PI3K signaling would be expected to decrease nuclear localization of FoxO1 (52). Because our mice showed no increase in Pomc gene expression, our results suggest that alternate mechanisms maintain normal STAT3 transcriptional activity in mice with disrupted PI3K signaling.

When exposed to chronic hyperleptinemia via chronic central infusion (53), recombinant adeno-associated virus–mediated gene delivery (54), or as a result of diet-induced obesity (55), rodents gradually become unresponsive to the anorexic effects of leptin. A number of causes of this leptin resistance have been proposed, including impaired leptin-induced PI3K signaling (56, 57). However, \( \text{Pik3r1} \) \( \text{POMCKO Pik3r2}^{−/−} \) mice, like WT mice, had both normal body weights and normal leptin levels, which suggests that \( \text{Pik3r1} \) \( \text{POMCKO Pik3r2}^{−/−} \) mice remain leptin sensitive. These findings support the idea that other mechanisms, such as the competition between FoxO1 and STAT3, play a role in maintaining normal leptin responsiveness.

### Figure 6

**Leptin fails to suppress feeding in Pik3r1 POMCKO Pik3r2\(^{−/−}\) mice.**

(A) Food consumption was measured 3 and 24 hours after leptin administration and compared with food consumption in each animal following saline administration. Values were normalized to kg body weight (\( n = 11–15 \)). (B) Body weight 24 hours after injection in Pik3r1 POMCKO Pik3r2\(^{−/−}\) mice and littermate controls lacking POMC-cre expression. Data are mean ± SEM. *\( P = 0.0487 \), **\( P = 0.0022 \) versus saline-injected controls; Student’s \( t \) test.

### Figure 7

**Energy homeostasis, fertility, and linear growth.**

(A) Body weight curves and (B) cumulative food intake of Pik3r1 POMCKO Pik3r2\(^{−/−}\) mice and littermate controls lacking POMC-cre expression on high-fat chow. (C) Body fat percentage and lean mass in 8-month-old male and female diet-induced obese mice. (D) Serum leptin levels in 12-week-old male mice fed normal chow.
argue against degradation of PI3K signaling in POMC neurons being solely responsible for obesity-related leptin resistance.

A model of intracellular PI3K function in POMC neurons. Previous studies have shown that deletion of p85α, its splice variants, and p85β results in severe disruption of PI3K signaling, while maintenance of one of these functional regulatory subunits of PI3K permits survival (58). For instance, mice with a liver-specific deletion of the p85α regulatory subunit of PI3K exhibit an improvement of hepatic and peripheral insulin sensitivity (59). These findings are explained by the fact that in many cell types, p85 is stoichiometrically in excess of p110, and free p85 can act as a dominant negative to inhibit PI3K signaling (16). In addition, free p85 can form a cytoplasmic sequestration complex with phosphorylated IRS-1, therefore limiting the ability of IRS-1 to activate PI3K at the membrane (60). It is therefore not surprising that our electrophysiological findings demonstrated that POMC neurons continued to respond to leptin when the p85β regulatory isoform was present. For similar reasons, p85β KO mice have previously been shown to have a paradoxical increase in insulin sensitivity (33). This underlying phenotype may contribute to the lack of an observable increase in long-term body weight in our Pik3r1 POMCKO Pik3r2−/− mice. Finally, it should be noted that the gene encoding the p55γ (p55γPik3r1) isoform of p85 was unaltered in this study. In contrast to the isoforms discussed above, p55γ expression barely exceeds background in the murine hypothalamus (28) and was considered unlikely to be involved in POMC PI3K signaling. In addition, the pharmacological inhibitors of PI3K signaling used in our electrophysiology experiments inhibit all classes of PI3Ks and resulted in the same effect as deletion of p85α, its splice variants, and p85β. However, we cannot exclude the possibility that the p55γ isoform plays some role in the nonacute actions of leptin.

Plum and colleagues recently concluded from the use of Pomc-cre/Ptenlox/lox mice that leptin depolarizes and increases the firing rate of POMC neurons largely independent of PI3K signaling (61). It should be noted, however, that their model represents a constitutive elevation of PIP3, while Pik3r1 POMCKO Pik3r2−/− mice presumably have very low PIP3 concentrations in POMC neurons. In addition, a direct evaluation of leptin’s effects on PI3K signaling was not performed in the former study. We have directly assessed the effect of suppressing PI3K in POMC neurons in the context of direct leptin action and found that PI3K signaling was required for POMC neuronal excitation in response to leptin.

The question as to how leptin and insulin can excite and inhibit POMC cells, respectively, via an identical intracellular signaling cascade is an intriguing one. One possible explanation arises from the fact that leptin and insulin have each been shown to alter neuronal activity in only a percentage of POMC cells. Previous reports suggest that leptin activates 30%–90% of POMC cells (5, 43, 62), while insulin inhibits 50%–70% of the POMC cell population (43, 62). It has not been resolved whether these subpopulations correspond to identical or disparate groups of neurons. It is possible that leptin-responsive POMC neurons may not be insulin sensitive, and vice versa. Additional studies are needed to understand the heterogeneity of POMC neuronal populations.

The data presented here supports a model in which leptin acute-ly activates ARC POMC neurons via a PI3K-dependent mechanism (Figure 9). Furthermore, PI3K appears necessary for the acute reduction in food intake and body weight by leptin, but not sufficient to alter long-term food intake or energy balance. These data suggest that although PI3K is a required downstream target for acute leptin regulation, other signaling cascades may be essential for leptin regulation of long-term energy balance.
maintained by mating Pik3r1<sup>loxp<sub>loxP</sub>/Pik3r2<sup>−/−</sup> Rosa mice with Pik3r1<sup>POMCKO</sup>Pik3r2<sup>−/−</sup> Rosa mice. Because Pik3r2<sup>−/−</sup> mice ubiquitously lacking p85α have a mild increase in insulin sensitivity (35), wild-type mice would be an inappropriate control for physiological studies, which compare phenotypes involving more than single POMC neurons. Thus, littermate controls with the same genetic background as experimental animals except for POMC-cre expression (Pik3r1<sup>loxp<sub>loxP</sub>/Pik3r2<sup>−/−</sup> Rosa mice) were used for all experiments. Any mouse that tested positive for deletion of the Pik3r1 gene in tail tissue was excluded from all studies. Genotyping was performed according to protocols described previously (6, 17, 34, 35, 64).

**Characterization of Pik3r1 gene deletion in the hypothalamus.** Adult mice were killed by carbon dioxide inhalation and immediately decapitated, and the brain was rapidly removed. The pituitary and olfactory bulbs were excised and snap frozen, and then sagittal brain sections of 1 mm thickness were cut via razor using a chilled Micro Brain Matrix (Braintree Scientific Inc.). ARC, dorsomedial hypothalamic nucleus, paraventricular nucleus, and NTS were microdissected, snap frozen, and stored at −80°C. DNA was extracted using the Qiagen DNeasy kit using Polyf sodium salt (Roche) as a carrier. PCR was performed using Bio-Rad iTagq according to the manufacturer’s instructions.

**RNA isolation and cDNA preparation.** RNA was extracted using an RNeasy kit (Qiagen) according to the manufacturer’s directions. Genomic DNA contamination was eliminated by DNase treatment. Preparation of cDNA for quantitative PCR assays was performed according to published protocols (64) with the following changes: 2.4 μg of RNA was first treated with 2U DNase I and 4.2 mM MgCl<sub>2</sub> in a final volume of 40 μl. The reverse transcription reaction was carried out in 100 μl final volume. Following cDNA synthesis, DEPC-H<sub>2</sub>O was added to increase the sample volume to 300 μl.

**Quantitative PCR.** Tissue mRNA levels were measured with an ABI 7900HT Sequence Detection System. Analysis of neupeptide expression was performed using the TaqMan-based efficiency-corrected ΔCt assay with 10 ng cDNA per reaction for 50 cycles (15). miRNAs with cycle times of 34 or greater were determined to be below detection. Primer concentrations were 75 nM for cyclin. We used predeveloped assays purchased from Applied Biosystems for NPY, AgRP, and POMC. Quantitative PCR data were analyzed using ABI instrument software SDS2.1. Baseline values of amplification plots were set automatically, and threshold values were kept constant to obtain normalized cycle times and linear regression data. PCR efficiencies were calculated from the slope of the resulting standard curves. Normalized mRNA levels in arbitrary units were obtained by dividing the averaged, efficiency-corrected values for sample mRNA expression by that for cyclinH1 RNA expression for each sample. The resulting values were expressed as fold change above control levels.

**Hormone measurements.** For leptin assays, tail vein blood was obtained from ad libitum chow-fed mice at 12 weeks of age. Serum was collected by centrifugation and assayed by ELISA (Crystal Chem Inc.) using a commercially available kit.

Resting plasma corticosterone levels were obtained between 2:00 p.m. and 4:00 p.m. from 8 mice of each group isolated for 3 days. The stress procedure
research article

was performed during the same time frame following the protocol described by Popova and colleagues (41). Briefly, psychosocial stress was induced in 8 mice of each strain by aggregation for 30 min in groups of 4 animals after 3-day isolation in individual cages. Blood samples were taken after immediate decapitation of animals (within 30 seconds of handling). Blood was collected from the trunk in heparin-free tubes. The corticosterone concentration was measured from serum by ELISA (AssayDesigns Correlate-EIA Corticosterone kit) according to the manufacturer’s instructions.

Electrophysiological studies. For these studies examining the behavior of single POMC neurons, 3 types of animals were used: POMC-GFP (5) mice expressing EGFP under the transcriptional control of the POMC gene (referred to as WT), Pkb million POMC KO mice (lacking p85β in POMC neurons) carrying a floxed Rosa-GFP allele, and Pkb million POMC KO Pkb two mice (also lacking p85β) carrying a floxed Rosa-GFP allele. Three- to 8-week old mice were deeply anesthetized with iso-flurane and transected perfused with a modified ice-cold ACSF (described below), in which an eosinomol amount of sucrose was substituted for NaCl. The mice were then decapitated, and the entire brain was removed as previously described (5, 12, 31). After removal, brains were immediately submerged in ice-cold, carbogen-saturated (95% O₂ and 5% CO₂) ACSF (126 mM NaCl, 2.5 mM KCl, 1.2 mM MgCl₂, 2.4 mM CaCl₂, 1.2 mM NaH₂PO₄, 21.4 mM NaHCO₃, and 11 mM glucose). High K⁺ buffer was made by replacing 56 mM NaCl by KCl. A brain block containing the hypothalamus was made. Coronal sections (200–250 μM) were cut with a Leica VT1000S Vibratome and then incubated in oxygenated ACSF at room temperature for a least 1 h before recording.

Slices were transferred to the recording chamber and allowed to equilibrate for 10–20 min prior to recording. The slices were bathed in oxygenated ACSF (32°C–34°C) at a flow rate of approximately 2 ml/min. Epifluorescence was briefly used to target fluorescent cells, at which time the light source was switched to infrared differential interference contrast imaging to obtain the whole-cell recording (Zeiss Axioskop FS2 Plus equipped with a fixed stage and a Hamamatsu C2741-60 charged-coupled device camera). Electrophysiological signals were recorded using an Axopatch 700B amplifier (Axon Instruments), low-pass filtered at 2–5 kHz, digitized at 88 kHz (Neuro-corder; Cygnus Technology), stored on videotape, and analyzed offline on a PC with pCLAMP programs (Axon Instruments) or Mini-analysis (Synaptosoft). Recording electrodes had resistances of 2.5–5 MΩ when filled with the N-glucosinate internal solution. Input resistance was assessed by measuring voltage deflection at the end of the response to a hyperpolarizing rectangular current pulse (400–500 ms of −20 pA).

Pipette solution for whole-cell recording contained 110–128 mM K-glucosinate, 10 mM KCl, 10 mM HEPES, 1 mM EGTA, 0.3 mM CaCl₂, 1 mM MgCl₂, 2–5 mM (Mg)·ATP, and 0.3 mM (Na)·GTP, pH 7.3. For cell-attached loose patch recordings, pipette solution contained 150 mM NaCl, 3.5 mM KCl, 10 mM HEPES, 10 mM glucose, 2.5 mM CaCl₂, and 1.3 mM MgCl₂, pH 7.2–7.4. Wortmannin (10–100 nM; Alomone Laboratories), LY294002 (10 μM; Calbiochem), PD98059 (25–50 μM; Calbiochem), insulin (50 nM, Humulin-R 100 U/ml), and leptin (100 nM; provided by A.F. Parlow, Harbor-UCLA Medical Center, Torrance, California, USA; through the National Hormone and Peptide Program) were added to the ACSF for specific experiments. Wortmannin was dissolved in DMSO and added to ACSF to obtain a final DMSO concentration of less than 0.1%. LY294002 was dissolved in 100% ethanol, with the final ethanol concentration in ACSF less than 0.5%. Solution containing leptin was typically perfused for 2–4 min. The leptin-induced effect was required to be associated temporally with leptin application, and the response had to be stable within a few minutes. For each recording, the averaged firing rate from every 20-s epoch was taken as 1 data point. A total of 12–18 data points that represent stable activity levels were taken to calculate mean and SD before and after application of leptin, respectively. A neuron was considered activated or inhibited if the change in firing rate induced by leptin was greater than ±3 SD prior to leptin addition. A neuron was considered depolarized or hyperpolarized if a change in membrane potential was at least 2 mV in amplitude.

ICV leptin administration. Stainless steel cannulae were inserted into the lateral ventricles (0.5 mm caudal and 1 mm lateral from bregma; depth, 2.5 mm) of anesthetized mice (100 mg/kg ketamine hydrochloride and 6 mg/kg Xylazine i.p.). Mice were allowed to recover for at least 1 week before i.c.v. injection and were not used until they regained their presurgery weights. Cannulation placement was confirmed by demonstration of increased thirst after administration of angiotensin (10 ng). A bolus injection of leptin (6 μg/mouse) or saline was administered i.c.v., and weight and food intake was measured at 0 and 45 min and 3 and 24 h.

Feeding studies and body composition. High-fat diet–induced obese mice were housed individually for all food intake measurements. Body weight and chow consumption was measured once every 7 days. High-fat, high-sucrose diet (D-12331, 58% kcal from fat, 26% from sucrose, 5.557 kcal/g; Research Diets) was replaced weekly in order to reduce spillage. At the conclusion of the diet-induced obesity study, mice were ketamine anesthetized and subjected to dual-energy X-ray absorptiometry (MEC Lunar Corp.) for analysis of fat and lean body composition.

Oxygen consumption. O₂ consumption was measured for 48 h using an 8-chamber open-circuit Oxymax system that is a component of the Comprehensive Laboratory Animal Monitoring System (CLAMS; Columbus Instruments). Mice were housed individually in specially built Plexiglas cages (5 in. high, 4.5 in. wide, 8.5 in. deep) maintained at room temperature (22°C) under an alternating 12-h light/12-h dark cycle. Sample air was sequentially passed through O₂ and CO₂ sensors (Columbus Instruments) for determination of O₂ and CO₂ content. Food and water were provided ad libitum. Mice were acclimatized to monitoring cages for 48 h before data collection.

Statistics. Results are expressed as mean and SEM. Comparisons between 2 groups were made by unpaired 2-tailed Student’s t test or 1-way ANOVA with repeated measures, as appropriate. A P value less than 0.05 was considered to be statistically significant.

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Address correspondence to: Joel K. Elmquist, University of Texas Southwestern Medical Center, 5323 Harry Hines Boulevard, Dallas, Texas 75390, USA. Phone: (214) 648-2911; Fax: (214) 648-5612; E-mail: joel.elmquist@utsouthwestern.edu.

Lewis C. Cantley, Bradford B. Lowell, and Joel K. Elmquist are co–senior authors.

Jennifer W. Hill and Kevin W. Williams are co–first authors.


