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Enhancing energy expenditure (EE) is an attractive strategy to combat obesity and diabetes. Global deletion of \textit{lp6k1} protects mice from diet-induced obesity (DIO) and insulin resistance, but the tissue-specific mechanism by which IP6K1 regulates body weight is unknown. Here, we have demonstrated that IP6K1 regulates fat accumulation by modulating AMPK-mediated adipocyte energy metabolism. Cold exposure led to downregulation of \textit{lp6k1} in murine inguinal and retroperitoneal white adipose tissue (IWAT and RWAT) depots. Adipocyte-specific deletion of \textit{lp6k1} (AdKO) enhanced thermogenic EE, which protected mice from high-fat diet–induced weight gain at ambient temperature (23°C), but not at thermoneutral temperature (30°C). AdKO-induced increases in thermogenesis also protected mice from cold-induced decreases in body temperature. UCP1, PGC1α, and other markers of browning and thermogenesis were elevated in IWAT and RWAT of AdKO mice. Cold-induced activation of sympathetic signaling was unaltered, whereas AMPK was enhanced, in AdKO IWAT. Moreover, beige adipocytes from AdKO IWAT displayed enhanced browning, which was diminished by AMPK depletion.

Furthermore, we determined that IP6 and IP6K1 differentially regulate upstream kinase-mediated AMPK stimulatory phosphorylation in vitro. Finally, treating mildly obese mice with the IP6K inhibitor TNP enhanced thermogenesis and inhibited progression of DIO. Thus, IP6K1 regulates energy metabolism via a mechanism that could potentially be targeted in obesity.

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Adipocyte-specific deletion of *Ip6k1* reduces diet-induced obesity by enhancing AMPK-mediated thermogenesis

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Enhancing energy expenditure (EE) is an attractive strategy to combat obesity and diabetes. Global deletion of *Ip6k1* protects mice from diet-induced obesity (DIO) and insulin resistance, but the tissue-specific mechanism by which IP6K1 regulates body weight is unknown. Here, we have demonstrated that IP6K1 regulates fat accumulation by modulating AMPK-mediated adipocyte energy metabolism. Cold exposure led to downregulation of *Ip6k1* in murine inguinal and retroperitoneal white adipose tissue (IWAT and RWAT) depots. Adipocyte-specific deletion of *Ip6k1* (AdKO) enhanced thermogenic EE, which protected mice from high-fat diet-induced weight gain at ambient temperature (23°C), but not at thermoneutral temperature (30°C). AdKO-induced increases in thermogenesis also protected mice from cold-induced decreases in body temperature. UCPI, PGC1-α, and other markers of browning and thermogenesis were elevated in IWAT and RWAT of AdKO mice. Cold-induced activation of sympathetic signaling was unaltered, whereas AMPK was enhanced, in AdKO IWAT. Moreover, beige adipocytes from AdKO IWAT displayed enhanced browning, which was diminished by AMPK depletion. Furthermore, we determined that IP6 and IP6K1 differentially regulate upstream kinase-mediated AMPK stimulatory phosphorylation in vitro.

Finally, treating mildly obese mice with the IP6K inhibitor TNP enhanced thermogenesis and inhibited progression of DIO. Thus, IP6K1 regulates energy metabolism via a mechanism that could potentially be targeted in obesity.

**Conflict of Interest:** The authors have declared that no conflict of interest exists.

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were identified in the mid-1990s (39, 40). 5-IP7 (5-diphosphoinositol pentakisphosphate) is the major IP7 isoform in mammals (41), which is synthesized by a family of 3 IP6 kinases (IP6Ks) (42). At lower ATP/ADP ratio, IP6Ks dephosphorylate IP6 to a distinct form of IP5 [Ins(2,3,4,5,6)P5, IP5*] (43). IP6 regulates cellular processes by binding to its protein targets (44), whereas 5-IP7 modulates its targets by binding or by pyrophosphorylation (42, 45–48). IP6Ks also regulate cellular functions via distinct mechanisms that do not require catalytic activity (49–52). For instance, IP6K1 interacts with glycogen synthase kinase and perilipin to modulate their cellular functions (49, 52).

Functions of 3 mammalian IP6K isoforms (IP6K1–3) are being explored. Although IP6Ks display conserved active sites, they possess unique regulatory regions, which mediate specific protein-protein interactions (45, 53). Accordingly, IP6Ks exhibit distinct cellular functions (45, 54). IP6K1-generated 5-IP7 modulates cell metabolism, signaling, exocytosis, and migration (48, 55–58). IP6K1-generated 5-IP7 regulates insulin signaling by inhibiting the insulin-sensitizing protein kinase Akt both in vivo and in vitro (58–61), whereas it promotes insulin secretion from pancreatic β cells (57). IP6K1 also regulates global transcription by 5-IP7-mediated inhibition of the histone demethylase JMJD2C (62). In immortalized mouse embryonic fibroblast (MEF) cells, IP6K1-generated 5-IP7 reduces cellular ATP level and lowers the glycolytic/oxidative phosphorylation ratio (55).

The compound TNP [N2-(m-trifluorobenzyl),N6-(p-nitrobenzyl) purine] (63) inhibits IP6Ks (64), whereas it does not influence the other IP7-generating (1-IP7-generating) enzyme PP-IP5K (65) or 71 unrelated kinases (64). Thus, TNP is a specific IP6K inhibitor. Accordingly, TNP reduces intracellular 5-IP7 levels and activates Akt (45, 58, 59, 61, 64, 66). TNP-mediated inhibition of IP6Ks also leads to a corresponding increase in the intracellular concentration of the IP6K substrate IP6 (58, 67). Recent studies indicate that TNP-mediated IP6K inhibition has functional significance in vivo. For instance, 5-IP7 level is increased in aged mesenchymal stem cells (61). TNP reduces 5-IP7, which stimulates Akt-mediated survival of these cells (61). Moreover, TNP treatment leads to a substantial reduction in 5-IP7 levels in the hearts of diabetic db/db mice (66). Accordingly, TNP protects cardiac ischemia in these mice via enhancement of Akt-mediated survival and AMPK/PGC1α-mediated mitochondrial biogenesis (66).

In order to understand the physiological role of IP6K1, mice with this gene globally deleted were generated (68). Deletion of Ip6k1 significantly influences various aspects of mouse biology, and thus, targeting this isoform is expected to be beneficial in obesity, T2D (45, 58, 69, 70), innate immunity (59), thromboembolism (71), cancer metastasis (56), and psychiatric and neurodegenerative diseases (49). Ip6k1 KO mice are protected against high-fat diet-induced (HFD-induced) weight gain, insulin resistance, and fatty liver (58). HFD-fed Ip6k1 KOs maintain insulin sensitivity by sustaining Akt activity (58). Moreover, HFD-fed Ip6k1 KO mice expend more energy, whereas their energy intake is unaltered (58). Yet the precise tissue-specific mechanism by which IP6K1 regulates energy homeostasis is not known. To determine this, we generated an adipocyte-specific Ip6k1 KO mouse model (AdKO).

Using this model, we demonstrate that IP6K1 regulates global energy homeostasis by modulating AMPK-mediated adipose tissue browning and thermogenesis. In addition to its inhibitory actions on Akt, IP6K1 reduces AMPK activity in the adipose tissue, although via a distinct and novel mechanism.

**Results**

Adipocyte-specific Ip6k1-deleted mice accumulate less fat because of increased thermogenic EE under chow-fed conditions. Adipocyte-specific Ip6k1 KO (AdKO) mice are generated by crossing of Ip6k1-LoxP mice (68) with Adipoq-Cre mice following standard procedure (Figure 1A). IP6K1 protein level is dramatically reduced in diverse adipose tissue depots of AdKOs (Figure 1B). Ip6k2 or Ip6k3 does not compensate for Ip6k1 deletion in the AdKO adipose tissue (Supplemental Figure 1A; supplemental material available online with this article; doi:10.1172/JCI85510DS1). Chow diet–fed AdKOs (CD-AdKOs), at a young age (3–4 weeks), do not display any variation in body weight, which indicates that the knockouts do not exhibit developmental defects in weight gain (Supplemental Figure 1B). At 2 months, AdKOs display a marginal decrease in fat mass (Figure 1C and Supplemental Figure 1C), whereas lean and fluid masses are unchanged (Supplemental Figure 1, D and E). Accordingly, weights of various adipose depots and their adipocyte sizes are marginally less in these mice (Figure 1D and Supplemental Figure 1F). Weights of other tissues such as liver and heart are similar (Supplemental Figure 1G). Food intake is comparable in CD-fed LoxPs and AdKOs (data not shown). Conversely, CD-AdKOs exhibit enhanced VO2/EE at 23°C (Figure 1, E and F). We also measured these parameters at 5°C (24 hours) and acute fasting (6 hours) at 5°C (to enhance expenditure of stored energy to maintain body temperature). CD-AdKOs exhibit higher VO2/EE under these conditions (Figure 1, E and G). Respiratory exchange ratio (RER) is slightly higher in CD-AdKOs at 23°C but not at 5°C (Supplemental Figure 1H). Further analyses (72) reveal that CD-AdKOs consume more oxygen for carbohydrate oxidation at 23°C and 5°C whereas they oxidize more fat when exposed to cold plus fasting conditions (Figure 1, H and I). Activity profiles are unaltered (Supplemental Figure 1I). Enhanced thermogenic EE protects CD-AdKOs from both acute- and chronic-cold-induced decline in body temperature (Figure 1, J and K). Further analyses indicate that Ip6k1 is the major isoform in various adipose depots of WT C57BL/6 mice (Figure 1L). Moreover, Ip6k1 expression is induced in inguinal WAT (IWAT) and retroperitoneal WAT (RWAT) but not in BAT following acute cold exposure (Figure 1L). Similarly to acute, chronic cold exposure also reduces Ip6k1 expression in the IWAT, whereas Ip6k2 and Ip6k3 levels are unaltered (Figure 1M). These results suggest that Ip6k1, the major Ip6k isoform in the adipose tissue, inhibits thermogenic EE.

**Increased thermogenic EE decelerates HFD-induced weight gain in AdKO mice.** At 23°C, AdKOs are protected against HFD-induced weight gain, which is apparent from 2 weeks and becomes significant after 4 weeks (Figure 2A and Supplemental Figure 2A). After 14 weeks of HFD, AdKOs appear substantially smaller, because of less fat accumulation (Figure 2B). In HFD-AdKOs, total fat mass is reduced (Δ 4.52 g) to a greater extent than lean mass (Δ 2.46 g) (Figure 2C). Percentage fat mass (of total body weight) is also reduced in HFD-AdKOs, whereas percentage lean mass is increased (Supplemental Figure 2B). Accordingly, weights of diverse adipose depots are significantly less in HFD-AdKOs (Fig-
Figure 1. Adipocyte-specific Ip6k1-deleted mice accumulate less fat because of enhanced thermogenic EE under chow-fed conditions. (A) Generation of AdKO. Details of the knockout generation strategy are described in Methods. (B) Immunoblot analyses confirm deletion of IP6K1 protein from various adipose tissue depots. Detection of residual IP6K1 is due to the presence of macrophages and other immune cells in the adipose tissue. Liver samples from AdKO mice were run as a control. BAT and liver samples were run on a separate gel. Blot represents data from 3 independent experiments. (C) CD-AdKOs (8 weeks old) display significantly reduced percentage fat mass (n = 10 mice per group; t test). (D) Weights of diverse adipose tissue depots are marginally less in CD-AdKOs (n = 4 mice per group; t test). (E) VO2 is higher in CD-AdKOs at 23°C and following cold and cold + fasting exposures (n = 8 mice per group). (F and G) EE is higher in CD-AdKOs under the above-mentioned conditions (n = 8 mice per group; t test). (H) CD-AdKOs consume more oxygen to oxidize carbohydrate at 23°C and 5°C (n = 8 mice per group; t test). (I) Following acute fasting at 5°C, CD-AdKOs consume more oxygen for fat oxidation (n = 8 mice per group; t test). (J) CD-AdKOs maintain body temperature more efficiently than LoxPs following acute cold + fasting exposure (n = 8 mice per group; 2-way ANOVA). (K) CD-AdKOs are also partly protected against drop in body temperature induced by chronic cold (3 days) and acute cold + fasting (n = 8 mice per group; 2-way ANOVA). (L) Ip6k1 is the major Ip6k isoform in the adipose tissue, which is downregulated by acute cold exposure in IWAT and RWAT. Ip6k1 expression is lower in the BAT and is not further reduced by cold (n = 5 mice per group; 2-way ANOVA). (M) Chronic cold also specifically reduces Ip6k1 expression in IWAT (n = 6 mice per group; t test). Data in all panels are expressed as mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001, §§P < 0.0001.
HFD-induced weight gain. Moreover, AdKOs maintain metabolic homeostasis by expending various forms of energy, depending on diet and temperature conditions.

Adipocyte-specific Ip6k1 deletion protects mice from HFD-induced insulin resistance. Blood glucose levels are slightly lower in CD-AdKOs at 23°C (Supplemental Figure 3A). However, AdKOs (23°C) are significantly protected against HFD-induced insulin resistance. Blood glucose level in AdKOs fed HFD for 4 weeks is slightly albeit significantly less in comparison with LoxPs (data not shown), and this is clearly evident after 14 weeks of HFD feeding (Figure 3A). Improved glucose and insulin tolerance tests (GTT and ITT), in mice fed HFD for 14 weeks, demonstrate that HFD-
higher in HFD-AdKOs (Figure 3, H–J, and Supplemental Figure 3C). Thus, adipocyte-specific deletion of \( Ip6k1 \) protects mice from HFD-induced global insulin resistance at ambient temperature.

Pharmacologic inhibition of IP6K enhances thermogenic EE and blocks the progression of HFD-induced obesity. Thus far, our results demonstrate that adipocyte-specific \( Ip6k1 \) deletion enhances thermogenic EE and global insulin sensitivity, which protects mice from HFD-induced weight gain and insulin resistance. Moreover, the lean phenotype of AdKOs resembles global \( Ip6k1 \) KO mice (58). Therefore, we monitored impacts of pharmacologic inhibition of the inositol pyrophosphate pathway on progression of diet-induced obesity (DIO) in C57BL/6 mice (Figure 4A). Mildly obese (4 weeks HFD-fed) mice were injected with vehicle or TNP for 7 weeks (Figure 4, A and B). Total body weight and weight gain are less in TNP-treated mice (Figure 4B and Supplemental Figure 4A).}

**Figure 3. Adipocyte-specific \( Ip6k1 \) deletion protects mice from HFD-induced insulin resistance.** (A) Blood glucose level is lower in ad libitum HFD-AdKOs. Overnight fasting decreases blood glucose level in HFD-AdKOs but not in corresponding HFD-LoxPs (\( n = 8 \) mice per group; \( t \) test). (B) Improved glucose disposal (GTT) in HFD-AdKOs (\( n = 8 \)–10 mice per group; 2-way ANOVA). (C) Improved glucose disposal in HFD-AdKOs following insulin injection (ITT) (\( n = 8 \)–10 mice per group; 2-way ANOVA). (D) AUC analyses of B and C confirm enhanced glucose and insulin tolerances in HFD-AdKOs (\( n = 8 \)–10 mice per group; \( t \) test). (E) HFD-AdKOs exhibit reduced plasma insulin level (\( n = 8 \) mice per group; \( t \) test). (F) Akt (S473) stimulatory phosphorylation is substantially higher in the EWAT, liver, and gastrocnemius muscle of HFD-AdKOs (\( n = 4 \) mice per group). (G) ImageJ quantification of F reveals that Akt stimulatory phosphorylation (S473) is 4- to 6-fold higher in EWAT, gastrocnemius muscle, and liver of HFD-AdKOs (\( t \) test). p/T, phosphorylated Akt/total Akt. (H) Plasma levels of HMW and total but not LMW ADIPOQ are higher in HFD-AdKOs (\( n = 7 \) mice per group). (I) ImageJ quantification of H reveals that the plasma level of HMW-ADIPOQ is 2.5-fold higher in HFD-AdKOs (\( t \) test). (J) ImageJ quantification of H demonstrates a 2-fold increase in the plasma level of total ADIPOQ in HFD-AdKOs (\( t \) test). Data in all panels are expressed as mean ± SEM. *\( P < 0.05 \), **\( P < 0.01 \), §\( P < 0.0001 \).
ure 4A). During the treatment period, vehicle mice gained 5.8 g while TNP mice lost 2.5 g body weight (Supplemental Figure 4B).

Accordingly, TNP reduces weight of diverse adipose tissue depots (Figure 4C). TNP mice also display enhanced VO₂ consumption, especially during dark conditions (Figure 4D), whereas energy intake is unaltered (data not shown). Total activity is similar in vehicle- and TNP-treated mice (Supplemental Figure 4C). TNP enhances thermogenic EE, as it protects these mice from cold-induced decline in body temperature (Figure 4E). Moreover, TNP improves glucose disposals following exogenous insulin (ITT) and glucose (GTT) injections (Figure 4, F–H).

**Adipocyte-specific Ip6k1 deletion enhances browning.** Next, we determined the mechanism by which IP6K1 regulates thermogenic EE. First, we monitored browning in *Ip6k1*-deleted adipose tissue. Chronic cold exposure dramatically induces adipose tissue remodeling, which resembles browning, especially in AdKO IWAT and RWAT (Figure 5A). PGC1-α and UCP1 expression levels are higher in AdKOs at 23°C, and further enhanced following chronic cold exposure (Figure 5B). Accordingly, PGC1-α and UCP1 protein levels are also elevated in IWAT and RWAT but not in BAT of AdKOs (Figure 5, C and D). Other components of the thermogenic/EE machinery such as *Cidea*, the transcriptional coregulator *Pdm16*, the transcription factors PPARα/δ, and the mitochondrial markers *Cpt1* and *COX2* are also elevated in chronic-cold-exposed AdKO IWAT (Figure 5E).

Thereafter, we determined whether *Ip6k1* deletion enhances browning in a cell-autonomous manner. Stromal vascular fractions (SVFs) isolated from IWAT of LoxPs and AdKOs were differentiated to beige adipocytes. AdKO SVFs display higher beige adipogenesis (Figure 5F), which is further evidenced by enhanced expression of the thermogenic/EE machinery (Figure 5G). Accordingly, mitochondrial oxygen consumption rate (OCR) and proton leak are higher in AdKO beige adipocytes (Figure 5, H and I). Glycolysis is unaltered in AdKO adipocytes (Supplemental Figure 5A). In contrast, browning/thermogenic marker expression levels are largely unaltered in AdKO BAT adipocytes (Supplemental Figure 5B). Thus, *Ip6k1* deletion enhances browning of WAT.

**IP6K1 reduces AMPK-mediated adipocyte browning.** How might *Ip6k1* deletion enhance browning and thermogenesis? AdKO IWAT SVFs display enhanced browning in a cell-autonomous manner (Figure 5, F and G). Moreover, global PKA signaling, tyrosine hydroxylase protein level, and β-AR (*Adrb3*) expression level are unaltered in cold-exposed AdKO IWAT (Supplemental Figure 6, A and B). These results suggest that hyperactivation of the sympathetic/β-AR pathway in AdKOs is unlikely. Conversely, Akt stimulatory phosphorylation is higher in chronic-cold-induced AdKO IWAT (Supplemental Figure 6, C and D). Activation of the PI3K/Akt pathway enhances browning, UCP1 expression, and glucose uptake (74–76). Moreover, ADIPOQ (which is higher in AdKOs) enhances cold-induced browning of subcutaneous adipose tissue via Akt-mediated proliferation of M2 macrophages (76). *Adipoq* expression is higher in the chronic-cold-exposed AdKO IWAT, yet the macrophage population is unaltered in this condition (Supplemental Figure 6E). Thus, although the ADIPOQ/Akt
axis is more active in cold-exposed AdKOs, it may not have exerted its influence on macrophages at least under the conditions tested.

Further analyses reveal that the stimulatory phosphorylation (T172) of AMPK is slightly higher in chronic-cold-exposed AdKOs (Supplemental Figure 6F). AMPK is sensitive to acute and subtle changes in energy and stress. Thus, we presumed that chronic cold exposure may have saturated AMPK activity. Indeed, WT mice display substantially enhanced AMPK stimulatory phosphorylation following acute cold exposure (Figure 6A). Therefore, we tested AMPK activity in acute-cold-exposed LoxPs and AdKOs. Acute-cold-induced AMPK stimulatory phosphorylation is substantially higher in AdKOs (Figure 6B and Supplemental Figure 6G). AMPK
phosphorylation is also higher in 2 separate preparations of beige adipocytes differentiated from IWAT SVFs of AdKOs (Figure 6C).

Moreover, Ip6k1 deletion–mediated enhancement in adipose tissue browning is modulated by AMPK, as AMPKα depletion reduces PGC1-α and UCP1 protein levels in AdKO adipocytes (Figure 6, D and E).

Next, we tested whether pharmacologic inhibition of IP6Ks or activation of AMPK stimulates UCP1 expression in beige adipocytes. TNP (5 μM, 16 hours) substantially enhances UCP1 expression in LoxP adipocytes (Figure 6F). Conversely, TNP does not further stimulate UCP1 expression in AdKO adipocytes (Figure 6F). The AMPK activator AICAR significantly enhances UCP1 expression in LoxP adipocytes albeit its effects on AdKO adipocytes are minimal (Figure 6F). Moreover, TNP enhances AMPK phosphorylation and activity in LoxP beige adipocytes (Figure 6G). Therefore, we monitored whether TNP enhances AMPK activity in the adipose tissue of HFD-fed mice. Indeed, a single dose of TNP substantially enhances AMPK phosphorylation and its activity in the IWAT of HFD-fed mice (Figure 6, H and I).

Thus, IP6K1 regulates adipose tissue browning via modulation of the AMPK pathway.

**IP6 and IP6K1 differentially regulate AMPK stimulatory phosphorylation.** Next, we determined the mechanism by which IP6K1 regulates AMPK activity and signaling. Overexpression of Ip6k1 in HEK293 cells reduces phosphorylation and activity of AMPK on its target ACC (Figure 7A). Thereafter, we monitored effects of various inositol phosphates and pyrophosphate on LKB1-mediated AMPK phosphorylation in a purified in vitro system. We used a commercially available unphosphorylated (UP) AMPKα2/β1/γ1 complex as the LKB1 substrate (Supplemental Figure 7A). LKB1 requires coactivators MO25 and STRADα for its full activity, and thus, in the absence of these coactivators, LKB1 is partly active (36). We used both LKB1-only (without coactivators) and LKB1-complex (with coactivators) on AMPKα2/β1/γ1 phosphorylation. First, we used LKB1-only, which phosphorylates UP-AMPK (Supplemental Figure 7, B and C). Next, we added increasing concentrations of IP6 and 5-IP7 in the assay. IP6 but not 5-IP7 potently stimulates LKB1-only–mediated UP-AMPK phosphorylation (Figure 7B). IP6 displays an EC50 of about 150 nM (Figure 7C and Supplemental Figure 7D). Among various inositol polyphosphates, IP6 is the most potent in the assay, followed by IP5, whereas others are less effective or are ineffective (IP6 > IP5 > IP4 = IP5* > IP3 > IP2) (Figure 7D). The products (5-IP7 and IP5*) of IP6K1 do not display any stimulatory effect on AMPK phosphorylation (Figure 7D). Thus, it is conceivable that in the presence of IP6K1, IP6 is converted to either 5-IP7 or IP5*, which neutralizes IP6-mediated AMPK activation. To test this, we used purified catalytically active and inactive versions of recombinant human IP6K1 (49). Catalytically active IP6K1 potently decreases IP6’s stimulatory action, whereas the inactive IP6K1 is largely ineffective (Figure 7E). 5-IP7, at physiological concentration, also reverses IP6’s stimulatory action.
removal of IP6 by converting it to 5-IP7 or IP5*. Endogenous IP6K1 and AMPK interact, which supports this possibility (Supplemental Figure 7E). Overexpression of glutathione S-transferase (GST)–IP6K1 deletion mutants reveals that the IP6K1 146–205 fragment effect (Figure 7F). We presume that 5-IP7 (1–5 μM in cells) is less likely to inhibit IP6-mediated activation, as IP6 is more potent in the assay, and is abundant (~50 μM) in cells (45). Rather, it is conceivable that IP6K1 interferes with IP6’s action on AMPK via removal of IP6 by converting it to 5-IP7 or IP5*. Endogenous IP6K1 and AMPKα interact, which supports this possibility (Supplemental Figure 7E). Overexpression of glutathione S-transferase (GST)–IP6K1 deletion mutants reveals that the IP6K1 146–205 fragment

Figure 7. IP6 and IP6K1 differentially regulate AMPK stimulatory phosphorylation. (A) Immunoblot demonstrates that overexpression of GST-IP6K1 diminishes AMPK and ACC phosphorylation levels in HEK293 cells under basal (+), glucose-deprived (−), and glucose-reintroduced (+/−) conditions. Data represent results from at least 3 independent experiments. (B) IP6, at increasing concentrations, stimulates AMPKα2/β1/γ1 (AMPKα T172) phosphorylation mediated by LKB1-only (120 ng/reaction). Data represent results from at least 3 independent experiments. (C) Quantification of B reveals that IP6 leads to 6- to 7-fold enhancements in LKB1-mediated AMPK phosphorylation in vitro. Data represent results from at least 3 independent experiments (1-way ANOVA). (D) Among inositol phosphates, IP6 most potently stimulates LKB1-only–mediated AMPKα2/β1/γ1 phosphorylation (IP6 > IP4 > IP3 > IP5 > IP5* > 5-IP7). 5-IP7 displays partial stimulatory effect. IP6K1 products IP5* and 5-IP7 are ineffective. Data represent results obtained from 3 independent experiments. (E) IP6K1 inhibits IP6-mediated AMPKα2/β1/γ1 phosphorylation by LKB1-only, whereas the inactive IP6K1 mutant is much less efficient. Data represent results obtained from 3 independent experiments. (F) 5-IP7 reverses IP6-mediated AMPKα2/β1/γ1 phosphorylation by LKB1-only. AMPKα2/β1/γ1 phosphorylation by LKB1 only. Data represent results obtained from 3 independent experiments. (G) IP6 also enhances LKB1-complex–mediated AMPKα2/β1/γ1 phosphorylation. Other inositol polyphosphates are ineffective. Data represent results obtained from duplicate experiments. (H) IP6 enhances CaMKKβ-mediated AMPKα2/γ1 phosphorylation. Data represent results obtained from duplicate experiments. (I) IP6 does not enhance LKB1-mediated phosphorylation of heat-denatured AMPKα2/β1/γ1. Data represent results obtained from 3 independent experiments. (J) IP6-mediated enhancement of AMPK phosphorylation is largely similar when AMPKα is used alone or in a complex with AMPKα2/β1/γ1. Data represent results obtained from 3 independent experiments. (K) AMP stimulates LKB1-mediated AMPKα2/β1/γ1 phosphorylation at 50- to 100-μM concentrations. Data represent results obtained from duplicate experiments. (L) IP6 or 5-IP7 does not influence active AMPK’s activity on its target SAMStide (n = 3; 1-way ANOVA). Data are expressed as mean ± SEM. §P < 0.0001.
α conformation, which indicates that IP6 may influence AMPK 2/α1/α2 interaction (Figure 7I). IP6 activates LKB1-only–mediated phosphorylation of AMPK 2/β1/α1 when AMPK phosphorylation (Supplemental Figure 7L). Moreover, IP6 stimulates CaMKK β-mediated phosphorylation compared with its effects on the LKB1-only assay (Supplemental Figure 7). IP6's binding to its bona fide substrate IP6 and/or (b) by IP6K1-interference neutralizes IP6's stimulatory action with AMPK α1 conformation is disrupted by heat denaturation (Figure 7J). AMP, the known AMPK activator, is less effective than IP6 in stimulating LKB1-only–mediated AMPK phosphorylation (Figure 7K). Conversely, IP6 and 5-IP7, unlike AMP, do not stimulate phosphorylated/active AMPK's catalytic activity on SAMStide (Figure 7L and Supplemental Figure 7M). Unlike AMP, IP6 and IP6K1 do not directly influence active AMPK's activity on its targets. Thus, IP6 enhances AMPK phosphorylation, whereas IP6K1's interference neutralizes IP6's stimulatory action with AMPK α1's binding to its bona fide substrate IP6 and/or (b) by IP6K1-mediated conversion of IP6 to 5-IP7 or IP5*.

Thermoneutrally placed HFD-AdKO mice do not display leanness, though they exhibit insulin sensitivity. If enhanced thermogenic EE reduces fat accumulation in AdKO mice, placing them at thermoneutral (30°C) temperature would, at least in part, impair their lean phenotype. Indeed, in this condition, HFD-fed LoxPs and AdKOs display largely similar body weight gain after 8 weeks (Figure 8A). HFD-induced body weight gain in AdKOs at 23°C is substantially less than that in LoxPs, whereas it is similar at 30°C (Figure 8B). Accordingly, thermoneutral HFD-AdKOs do not display any alterations in body composition (Figure 8C). EE, which is higher in AdKOs at 23°C after 8 weeks of HFD (Supplemental Figure 2J), is unaltered at 30°C (Figure 8D). RER and total activity profiles are largely similar in HFD-fed LoxPs and AdKOs in this condition (Supplemental Figure 8, C and D). Like the LKB1-only assay, inositol polyphosphates other than IP6 are largely ineffective (Figure 7G). IP6K1 also inhibits IP6's effects on LKB1-complex–mediated AMPK phosphorylation to a similar extent compared with its effects on the LKB1-only assay (Supplemental Figure 7L). Moreover, IP6 stimulates CaMKKβ-mediated AMPK phosphorylation (Figure 7H). These results suggest that IP6 may not influence upstream kinases of AMPK. Conversely, IP6 fails to activate LKB1-only–mediated AMPK phosphorylation when AMPKα2/β1/γ1 conformation is disrupted by heat denaturation (Figure 7I). IP6 activates LKB1-only–mediated phosphorylation of the AMPKα2/β1/γ1 complex and AMPKα alone to a similar extent, which indicates that IP6 may influence AMPKα conformation (Figure 7J). AMP, the known AMPK activator, is less effective than IP6 in stimulating LKB1-only–mediated AMPK phosphorylation (Figure 7K). Conversely, IP6 and 5-IP7, unlike AMP, do not stimulate phosphorylated/active AMPK's catalytic activity on SAMStide (Figure 7L and Supplemental Figure 7M). Unlike AMP, IP6 and IP6K1 do not directly influence active AMPK's activity on its targets. Thus, IP6 enhances AMPK phosphorylation, whereas IP6K1's interference neutralizes IP6's stimulatory action with AMPK α1's binding to its bona fide substrate IP6 and/or (b) by IP6K1-mediated conversion of IP6 to 5-IP7 or IP5*.
largely similar body weight compared with LoxPs (Supplemental Figure 8E). Accordingly, adipose tissue browning and mitochondrial activity markers are similar in LoxPs and AdKOs under this condition (Figure 8E), which further suggests that leanness in AdKOs is due to increased thermogenesis. However, at 30°C, AdKOs fed HFD for 14 weeks dispose glucose more efficiently than LoxPs (Figure 8, F and G). Akt stimulatory phosphorylation is also higher in AdKO adipose tissue and liver under this condition (Figure 8H and Supplemmental Figure 8F). Thus, 14 weeks of thermoneutrality abolishes leanness/browning in HFD-AdKOs, whereas insulin sensitivity is retained. Although it is not entirely clear at this point, we presume that IP6K1-mediated distinct regulation of browning and insulin sensitivity partly uncouples the lean and insulin-sensitive phenotype at thermoneutral temperature.

Discussion

By generating an adipocyte-specific mouse model (AdKO), we demonstrate that IP6K1 regulates global energy homeostasis via modulation of adipose tissue energy metabolism. AdKOs display enhanced AMPK-mediated browning/thermogenesis, which protects them from HFD-induced weight gain. In addition, HFD-fed AdKOs exhibit higher activity of the insulin-sensitizing protein kinase Akt in the adipose tissue. Moreover, HFD-AdKOs show higher plasma levels of ADIPOQ. As a result, the knockouts are protected against HFD-induced weight loss and insulin resistance. Thus, adipocyte-specific IP6K1 is a major regulator of global energy metabolism. Striped and bold encircling arrows indicate reduced and increased effects of AMPK and Akt on tissue metabolism, in the presence and absence of IP6K1, respectively. Thin and thick straight arrows indicate less and more active pathways, respectively.
Our study also unravels a novel mechanism by which IP6K1 regulates adipocyte browning. IP6K1’s substrate IP6 stimulates LKB1-mediated AMPK phosphorylation, whereas IP6K1 and its product 5-IP7 diminish IP6’s stimulatory effect. Thus, in the absence of IP6K1, AMPK activity is higher in the adipose tissue. A number of studies appreciate inositol polyphosphates as regulators of protein kinases such as BTK, CK2, and Akt (46, 58, 79). In-depth biophysical, structural, and biochemical studies are required to further dissect the molecular mechanism by which IP6/IP6K1 regulates AMPK. 5-IP7 may also indirectly inhibit AMPK. For example, 5-IP7 promotes nuclear localization of LKB1 in cancer cells, which diminishes its activity toward the cytosolic targets (80). Cytosolic LKB1 phosphorylates AMPK (81). Accordingly, global Ip6k1 KO IWAT SVFs display higher cytosolic LKB1 (unpublished observations).

IP6/IP6K1-mediated regulation of AMPK may have broader implications, as aberration of the pathway is also observed in other diseases such as cancer, neurodegeneration, and aging. Accordingly, active research is ongoing to identify small-molecule activators of AMPK (82, 83). However, to our knowledge, physiological small-molecule activators of AMPK, other than AMP, are not known. Therefore, AMPK activation by the biomolecule IP6 is novel. IP6K1-AMPKα interaction interferes with IP6-mediated AMPK activation. Thus, disruption of the interaction seems to be critical in enhancing AMPK activity. Accordingly, TNP, which inhibits IP6K activity (64), increases IP6 levels (58, 67), and disrupts IP6K1-AMPK binding (Supplemental Figure 7H), dramatically enhances AMPK activity in adipocytes. Enzymes that generate and/or metabolize IP6 may also regulate AMPK. For example, the primary physiological role of inositol polyphosphate multikinase (IPMK) is to form the bulk of IP5, which is the precursor of IP6, and thus, Ipmk deletion reduces IP5 and IP6 levels (84, 85). IPMK-AMPK interaction is regulated by glucose availability (86). Moreover, metformin-induced AMPK stimulatory phosphorylation and activity are impaired in Ipmk-deleted MEFs, which are restored by overexpression of catalytically active IPMK (84). Thus, it is conceivable that the ratio of IP6 to 5-IP7 in a microenvironment may regulate cellular targets such as AMPK. The ratio of IP6 to 5-IP7 is altered by various intrinsic or extrinsic factors such as growth factor deprivation, altered ATP levels, chemotactic stimulation, and apoptotic stimuli (43, 45, 58, 59). In addition, various conditions that regulate expression levels of Ip6k1 (such as cold exposure) may also alter the ratio.

IP6K1-mediated regulation of EE exhibits cell/tissue specificity. For instance, global Ip6k1 KO mice (58) and AdKOs display enhanced EE. Moreover, Ip6k1-deleted beige adipocytes show increased mitochondrial OCR due to proton leak, whereas glycolysis is unchanged. TNP-treated diabetic cardiomyocytes also exhibit enhanced mitochondrial biogenesis and function (66). Conversely, immortalized MEFs from global Ip6k1 KO mice are reported to display higher glycolysis whereas oxygen consumption is compromised (55). AMPK stimulates glycolysis (28); thus it is feasible that increased AMPK in certain tissues may also enhance glycolytic energy metabolism. In conclusion, improved EE following adipocyte-specific Ip6k1 deletion or IP6K inhibition indicates that the pathway may have therapeutic potential in human obesity, T2D, and other metabolic diseases.

Methods

Quantitative RT-PCR probes. TaqMan probes were purchased from Life Technologies.

Antibodies. Phospho-Akt (S473; catalog 4060), total Akt (pan; catalog 2920), PKA phospho-substrate (catalog 9621), phospho-ACC (S79; catalog 3661), total ACC (catalog 3676), phospho-AMPK (T172; catalog 2535), total AMPK (catalog 5831), and ADIPOQ total monomer (catalog 2789) were from Cell Signaling Technology. ADIPOQ HMW/LMW (catalog 5901) was from Biovision. Tyrosine hydroxylase (catalog TI299) and UCP1 (catalog U6382) were from Sigma-Aldrich. β-Actin (catalog sc-47778) was from Santa Cruz Biotechnology. PGCl-α (catalog AB3242) was from EMD Millipore. IP6K1 (catalog GTX103949) was from Genetex.

Enzymes and proteins. LKB1 only (catalog L15-30G), unphosphorylated/inactive AMPK (α2/β1/γ1) (catalog P48-14H), active CaMKKβ (catalog C18-10G), and phosphorylated/active AMPK (catalog P48-10H) were from SignalChem. Unphosphorylated/inactive AMPKα2 only (catalog H00005563-P01) was from Abnova. LKB1-MO25/STRA7 (catalog 14-596) was from EMD Millipore. Insulin was from Novo Nordisk.

Kits. The Vectastain Elite ABC kit was from Vector Laboratories. The BCA protein assay kit was from Pierce Biotechnology. The insulin ELISA was from Crystal Chem Inc.

siRNA. AMPKα (catalog sc-45313) and control (catalog sc-37007) siRNAs were from Santa Cruz Biotechnology.

Protease plus phosphatase inhibitor tablets and glutathione arsagrose. Protease plus phosphatase inhibitor tablets and glutathione arsagrose were from Thermo Scientific. Unless otherwise stated, all other chemicals were purchased from Sigma-Aldrich.

Animals. Mice were housed in 12-hour light/12-hour dark cycles at 23°C and were fed standard chow diet (CD), unless otherwise stated. Two-month-old male mice fed a CD (2018SX; Harlan Laboratories) or a HFD (60% kcal HFD for indicated periods; S3282; Bio-Serv) were used in the studies. For tissue collection, mice were euthanized by carbon dioxide asphyxiation (ARC Facility).

Generation of AdKO mice. The Ip6k1f/f mouse was previously described (68) and was provided by Solomon Snyder (Johns Hopkins University, Baltimore, Maryland, USA). Details of the strategy to target the Ip6k1 locus are mentioned in a previous literature (68). Briefly, the mouse Ip6k1 gene has 6 exons, of which exon 1 is exclusively non-coding (68). The start codon is located in exon 2, and the stop codon is located in exon 6. Exon 6 encodes the C-terminal domain of Ip6k1 (amino acids 265–433), and the 3′-UTR of the mRNA. Ip6k1 targeting was achieved by deletion of the coding region of exon 6, including the exon 6 splice site. The resulting transcript lacks exon 6 (including the 3′-UTR) and therefore is unstable, resulting in a complete knockout.

To generate adipocyte-specific Ip6k1 KO mice, the following mice were purchased from The Jackson Laboratory: FLP (neomycin) deleter (stock 009086); Adipoq-Cre (stock 010803). FRT sites flanking the neomycin resistance gene facilitate its removal by FLP recombinase, and loxp sites facilitate removal of the targeted exon 6 by Cre recombinase. Ip6k1f/f mice were mated with mice expressing FLP recombinase (JAX stock 009086) to excise the neomycin resistance gene to generate Ip6k1f/f/f/f mice. Genotyping was performed by PCR using primer sequences from exon 6 flanking the neomycin resistance gene and FRT sites. Homozygous Ip6k1f/f/f/f mice were crossed with the transgenic mice expressing Cre recombinase under the control of the ADIPOQ promoter (JAX stock 010803). Offspring inheriting both the
targeted allele and the Cre transgene (Adipoq-Cre-Ip6k1flipped/flipped) were crossed with Ip6k1flipped/flipped mice to yield Ip6k1flipped/flipped Cre adipocyte-specific knockout mice (AdKOs). AdKOs were verified by PCR with Cre primer sequences and primers flanking LoxP sites and exon 6 using DNA extracted from their tails. Male mice were used along with their LoxP (Ip6k1flipped/flipped) littermate controls unless otherwise indicated.

**Exposure to cold temperature.** Chronic exposure was performed following standard procedure (87). Briefly, 2-month-old LoxPs and AdKOs were placed in a cold room (5°C). After 6 days of cold exposure, mice were sacrificed and tissues were collected for histology and protein analysis.

To induce expenditure of stored energy under cold conditions to maintain body temperature, mice were simultaneously fasted and exposed to cold for indicated time periods (88). Rectal temperature was measured at indicated time points using a mouse-specific rectal probe (RET-3) and a TH-5 thermometer (Physitemp).

**Exposure to thermoneutral temperature.** Animals were exposed to thermoneutral temperature following standard procedure (89). Briefly, weaned mice (22 days old) were introduced to a 30°C room. After 1 week of acclimatization, mice were fed a HFD for the indicated period of time.

**Food intake studies.** Food intake was monitored in HFD-fed LoxPs and AdKOs (at the onset of HFD) using BioDAQ instrumentation (Research diet) following the vendor’s recommendations. Mice were acclimatized for 3 days. Afterward, food consumption of each mouse was recorded for 5 consecutive days. Each meal is defined as food intake of 0.1 g or more within 900 seconds, which is the gap between feeding bouts. Data represent the average daily intake. Average values are compared by 2-tailed Student’s t test to determine significance.

**Comprehensive Laboratory Monitoring System.** Mice were placed individually in metabolic cages with a precise thermostatic control in a Comprehensive Laboratory Monitoring System (CLAMS; Columbus Instruments) and were acclimatized at ambient temperature for 2 days. Afterwards, VO₂, VCO₂, and spontaneous locomotor activity were measured for indicated time periods. Cold-induced oxygen consumption was assessed by changing the temperature from ambient to 5°C for 24 hours followed by 6 hours of fasting. Respiratory exchange ratio (RER) and energy expenditure (EE) were calculated using the following equations: RER = VCO₂/VO₂; EE (kcal/h) = (3.815 + 1.232 × RER) × VO₂. Values were normalized by lean body mass.

**Calculation of VO₂ used for carbohydrate or fat oxidation.** Based on mean RER values, the exact percentage of VO₂ consumed for carbohydrate and fat oxidation can be determined (72). For example, RER 1.00 means 100% carbohydrate, whereas RER 0.7 indicates 100% fat consumption. A complete chart is presented in the literature (72).

Accordingly, we calculated VO₂ consumed by carbohydrate and fat for AdKOs and LoxPs under various conditions. Significance was calculated by 2-tailed Student’s t test.

**NMR.** Fat, lean, and fluid mass of CD- and HFD-fed mice was measured using the Minispec LF-NMR (Bruker Optics) analyzer.

**Blood glucose measurement.** Blood glucose in ad libitum and overnight-fastened HFD-fed mice was measured using a glucometer (Bayer) with glucose testing strips as described previously (58). Significance is calculated by 2-way ANOVA.

**Glucose tolerance test.** GTT was performed after 14 weeks in HFD-fed AdKOs under various temperature conditions unless otherwise stated. Glucose (2 g/kg body weight; i.p.) was injected into overnight-fasted mice. After injection, blood was taken by puncturing of the tail vein, and glucose levels were measured using a glucometer at indicated time periods. Blood glucose was also measured before the injection (time point 0).

**Insulin tolerance test.** After 14 weeks of HFD at 25°C, ITT was performed in AdKOs and LoxPs. After 6 hours of fasting in the morning, human recombinant insulin (0.75 U/kg body weight; i.p.) was injected. Blood glucose levels were measured as described above.

**TNP injection in mice.** WT C57BL/6 mice were fed a HFD for 4 weeks. Thereafter, vehicle or TNP (10 mg/kg body weight; daily; i.p.) (n = 5 per group) was injected for 7 weeks. Laboratory-synthesized TNP was dissolved in DMSO/Tween-80/water (1:1:8). Weekly body weight was measured. After 4 weeks of injection, oxygen consumption was measured as described above. GTT and ITT were performed on the fifth and sixth weeks of TNP treatment, respectively. In an independent experiment (n = 3 per group), mice were fed a HFD and treated with TNP as described above. After 7 weeks, they were cold-exposed for 4 days. To monitor AMPK activation, a single dose (20 mg/kg body weight) of TNP was injected in DIO mice. After the treatment, mice were fasted for 5 hours, after which they were sacrificed for tissue collection.

**Plasma insulin measurement.** Blood was collected from LoxPs and AdKOs after 14 weeks of HFD feeding in fed and 6-hour-fast conditions inside a K,EDTA-containing tube, and plasma was isolated by centrifuge at 4°C for 15 minutes at 2,000 g. Insulin concentrations were determined using an ultrasensitive mouse ELISA kit.

**Histological and immunohistochemical analyses of adipose tissue.** Tissues isolated from various adipose depots were fixed overnight in 10% neutral-buffered formalin. Samples were processed in a dehydrating ethanol gradient, followed by xylene incubation and paraffin embedding. Serial 8-μm sections were used for H&E staining, or for UCP1 immunohistochemical analysis. For immunohistochemistry, primary antibody was detected by Vectastain Elite ABC kit and diaminobenzidine reagent using the manufacturer’s instructions.

**Analyses of plasma ADIPOQ.** ADIPOQ complexes in the plasma were monitored following standard protocol (90). Briefly, equal volume of plasma was mixed with Laemmli buffer without β-mercaptoethanol, and proteins were resolved in 4%-15% SDS-PAGE. Primary antibody against ADIPOQ complex (Biovison) was used to detect the high-molecular weight (HMW) and low-molecular weight (LMW) complexes. To quantify total ADIPOQ, Laemmli buffer containing β-mercaptoethanol was used, and samples were heated at 95°C for 10 minutes. Primary antibody against monomeric ADIPOQ (Cell Signaling Technology) was used. The nitrocellulose membranes were ponczua-stained to confirm sample loading.

**RNA isolation and quantitative RT-PCR.** RNA isolation was done using the traditional Trizol method. Quantitative gene expression was performed by 2-step quantitative RT-PCR (qRT-PCR) using Applied Biosystems kit and TaqMan probes. mRNA expression levels were determined using the ΔΔCT method and normalized to the housekeeping gene β-actin.

**SVF culture and beige adipocyte differentiation.** Subcutaneous ingui

nal fat from 6- to 8-week-old mice (n = 6 per group) was used for isolating stromal vascular cells as previously described (91). Cells were cultured in DMEM/F12 containing 10% FBS and penicillin/streptomycin. Ninety-five to ninety-seven percent confluent cells were differentiated with a medium containing 5 μg/ml insulin, 125 μM indomethacin, 2 μg/ml dexamethasone, 0.5 mM 3-isobutyl-1-methylxanthine, 0.5 μM rosiglitazone, and 1 nM 3,3',5-triiodo-l-thyronine (T₃). After 2 days, medium was replaced with maintenance medium containing 5 μg/ml insulin and 2 μg/ml dexamethasone. After 6 days of culture, cells were fixed and stained for a single day using a 3,3′,5-triiodo-l-thyronine (T₃). After 2 days, medium was replaced with maintenance medium containing 5 μg/ml insulin and
1 nM T3. Medium was refreshed every 2–3 days until the cells were fully differentiated at day 7. For AICAR and TNP treatments, differentiated beige adipocytes were incubated overnight with 1 mM AICAR or 5 μM TNP. Cells were washed with PBS and harvested for qRT-PCR analyses.

Mitochondrial respiration analysis. For measuring mitochondrial respiration, SVF cells were seeded into 96 XF plates and differentiated to beige adipocytes. Oxygen consumption rate (OCR) and extracellular acidification were measured by Seahorse Extracellular Flux Analyzer (Seahorse Bioscience) in XF assay medium supplemented with 5 mM pyruvate and 2.5 mM glucose. After baseline measurements, 3 sequential injections were applied to the cells as indicated: oligomycin (1 μg/ml), FCCP (1 μM), and antimycin A (0.8 μM) combined with rotenone (3 μM). Basal, uncoupled, and maximal OCRs were calculated by averaging of values from each phase.

Signaling studies. SVF cells isolated from IAWT were differentiated to beige adipocytes as described above, and the cells were lysed by addition of a lysis buffer (20 mM Tris pH 7.4; 150 mM NaCl; 1% Triton X-100) containing a protease-phosphatase inhibitor tablet. Total protein was quantified using a BCA protein assay kit (Pierce Biotechnology). Equal amounts of total protein were loaded on SDS-PAGE. Various proteins and their phosphorylation levels were monitored by immunoblotting following standard procedures (58). Beige adipocytes or 3T3L1 preadipocytes were treated with indicated concentrations of TNP under basal and/or 2-hour-glucose-starved conditions. HEK293 cells were transfected using Polyfect transfection reagent (Qiagen) following standard procedure (93), all the other inositol phosphates other than IP5* were purchased from Cayman Chemical. IP5* was from Enzo Life Science.

To determine IP6 and IP6K1’s effects on AMPK phosphorylation mediated by LKB1-complex (with MO25 and STRADα), LKB1 concentration for the assay was determined to be 10 ng/reaction, as opposed to 120 ng/reaction of LKB1-only. Everything else was kept similar to the above assay.

To compare IP6’s effects on LKB1-mediated AMPK (α2/β1/γ) and AMPK2, equimolar concentrations of AMPK (25 nM) were used as substrates. In the assay of CaMKK2-mediated AMPK activity, active CaMKKβ (10 ng/reaction) was used instead of LKB1, and the rest was similar as above. Except for 5-IP7, which was chemically synthesized following standard procedure (93), all the other inositol phosphates were purchased.

AMPK activity assay in vitro. AMPK activity was determined using the SAMS peptide (SAMStide; Anaspec) as a substrate. The assay buffer consisted of 25 mM Tris (pH 7.2), 5 mM MgCl2, 2.5 mM DTT, 200 ng/l reaction mixtures onto individual precut strips of 33P-labeled IP6K1. The extent of AMPK phosphorylation (T172) was detected by immunoblotting following standard procedures if they showed any signs of sickness. Numbers of mice (n) used in experiments are indicated in the legends. Immunoblots were quantified using ImageJ software (NIH). For multiple comparisons, 2-way ANOVA with Holm–Šidák multiple comparison test was used. For 2 independent data sets, 2-tailed Student’s t test was used. Data are presented as mean ± SEM (P < 0.0001, *P < 0.001, **P < 0.01, and *P < 0.05). Statistical significance was calculated in GraphPad Prism, version 6. The data show normal variance.

Statistics. Trial/prior experiments were used to determine sample size with ample statistical power. Animals were excluded from experiments if they showed any signs of sickness. Numbers of mice (n) used in experiments are indicated in the legends. Immunoblots were quantified using ImageJ software (NIH). For multiple comparisons, 2-way ANOVA with Holm–Šidák multiple comparison test was used. For 2 independent data sets, 2-tailed Student’s t test was used. Data are presented as mean ± SEM (P < 0.0001, *P < 0.001, **P < 0.01, and *P < 0.05). Statistical significance was calculated in GraphPad Prism, version 6. The data show normal variance.

See complete unedited blots in the supplemental material.

Study approval. Animal care and experimentation were approved by the Scripps Florida Institutional Animal Care and Use Committee.

Author contributions. AC conceived and supervised the project; QZ generated AdKO mice; QZ and AR maintained mouse colonies; QZ performed the majority of the experiments; S. Ghoshal performed pharmacologic experimen-
mements; AR and S. Gao provided technical support; AA and TMK synthesized TNP; JCB supplied chemically synthesized 5-IP7; AC and QZ analyzed and interpreted data; AC wrote the manuscript.

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