Deficiency of Phosphoinositide 3-Kinase Enhancer Protects Mice From Diet-Induced Obesity and Insulin Resistance

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OBJECTIVE—Phosphoinositide 3-kinase enhancer A (PIKE-A) is a proto-oncogene that promotes tumor growth and transformation by enhancing Akt activity. However, the physiological functions of PIKE-A in peripheral tissues are unknown. Here, we describe the effect of PIKE deletion in mice and explore the role of PIKE-A in obesity development.

RESEARCH DESIGN AND METHODS—Whole-body PIKE knockout mice were generated and subjected to high-fat–diet feeding for 20 weeks. The glucose tolerance, tissue-specific insulin sensitivity, adipocyte differentiation, and lipid oxidation status were determined. The molecular mechanism of PIKE in the insulin signaling pathway was also studied.

RESULTS—We show that PIKE-A regulates obesity development by modulating AMP-activated protein kinase (AMPK) phosphorylation. PIKE-A is important for insulin to suppress AMPK phosphorylation. The expression of PIKE-A is markedly increased in adipose tissue of obese mice, whereas depletion of PIKE-A inhibits adipocyte differentiation. PIKE knockout mice exhibit a protected phenotype of lipodystrophy and are resistant to high-fat diet–induced obesity, liver steatosis, and diabetes. PIKE knockout mice also have augmented lipid oxidation, which is accompanied by enhanced AMPK phosphorylation in both muscle and adipose tissue. Moreover, insulin sensitivity is improved in PIKE-A–deficient muscle and fat, thus protecting the animals from diet-induced diabetes.

CONCLUSIONS—Our results suggest that PIKE-A is implicated in obesity and associated diabetes development by negatively regulating AMPK activity.

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Obesity is a result of imbalanced energy intake and expenditure in which the accumulation of excessive fat causes disorders such as type 2 diabetes, atherosclerosis, and dyslipidemia (1). Because of its increasing prevalence in most of the world, obesity has become a major health problem (2). Although genetic linkage analysis has successfully mapped potential loci in human genome for adiposity development (3), identifying all genetic variants that contribute to differences in body weight is still one of the major goals to fully understand the mechanism of obesity progression. Recent studies using genome-wide linkage scan revealed human chromosome trait 12q14.1, where the phosphatidylinositol 3-kinase (PI 3-kinase) enhancer (PIKE) gene CENTG1 locates, has a strong correlation with serum lipid level and energy intake (4,5), suggesting PIKE may be a potential factor in regulating body weight.

PIKes are a family of GTPases that directly interact with PI 3-kinase and Akt and enhance their kinase activities (6–8). The family contains three members: PIKE-L, PIKE-S, and PIKE-A, which is generated from alternative splicing of the CENTG1 gene. Whereas PIKE-S and -L are brain specific, PIKE-A is widely expressed, such that its mRNA could be detected in brain, heart, liver, muscle, spleen, thymus, and small intestine (9,10). The mode of action of PIKE is isoform specific in different cell types. PIKE-L couples to receptors such as netrin receptor (UNC5B) and metabotropic glutamate receptors I (mGluR-I) and links the activated receptor to PI 3-kinase pathway in neurons (11,12). PIKE-S localizes in nucleus and executes the protective effects of nerve growth factor by activating the nuclear PI 3-kinase cascade (8). PIKE-A, on the other hand, substantiates the kinase activity of Akt in glioblastomas and is involved in cancer invasion activity (6,13,14). However, the role of PIKE-A in peripheral tissues remains unknown.

In many cases, insulin resistance is the major associated pathologic condition of obesity. However, the molecular mechanism of this obesity-induced disorder remains ambiguous. It has been proposed that lipotoxicity is one of the candidates to explain the role of excess lipid storage in insulin resistance onset. Accumulation of excess cellular lipid changes the lipid metabolism, enhances oxidative stress, and disrupts endoplasmic reticulum homeostasis (15). Increasing cellular lipid oxidation by pharmacologic interventions in obese subjects thus represents a potential therapeutic regimen to mitigate their diabetic complications. In this regard, AMP-activated protein kinase (AMPK) is one of the targets. AMPK is the master sensor for energy status and is responsible for metabolic homeostasis (16).
Activation of AMPK results in reducing hepatic glucogeno-
genic gene expression and glucose production, increasing fatty acid oxidation, and enhancing glucose uptake. There-
fore, AMPK activators such as AICAR and metformin are effective agents in relieving the obesity-induced insulin resistance in both laboratory and clinical tests (17).

To examine the role of PIKE in obesity, we developed the whole-body PIKE knockout (PIKE−/−) mice with ablation of all PIKE isoforms. Here we report that PIKE-A is implicated in adipocyte differentiation and obesity de-
velopment. PIKE knockout elicits lipoliprophy and in-
creased insulin sensitivity by enhancing AMPK activity, leading to resistance against high-fat diet (HFD)-induced obesity and diabetes.

RESULTS

Generation of PIKE knockout mice. As a pioneer study on the physiological role of PIKE in obesity development, we generated whole-body PIKE−/− mice with targeted disruption in the CENGT1 locus using the LoxP/Cre system. We first created a transgenic line with PIKE flox/+ allele by inserting two loxP sites into the introns flanking exons 3 and 6 (Fig. 1A). PIKE flox/+ mice were then bred with transgenic mice expressing Cre recombinase in all tissues. Deletion of exons 3–6 results in removal of GTPase domain and introduces a frameshift mutation that creates a new stop codon, producing truncated PIKE proteins for all isoforms. Heterozygous mating generated newborn pups at expected Mendelian frequency that ap-
peared indistinguishable from the wild-type littersmates, suggesting that PIKE was dispensable for embryonic de-
velopment. Southern blot analysis showed exons 3–6 of the CENGT1 gene were effectively excised (Fig. 1B), which was further supported by PCR analysis (Fig. 1C). Immunoblotting analysis using antibody specific to the COOH-terminal of PIKE-A and RT-PCR confirmed the ablation of PIKE-A expression in various tissues (Fig. 1D and E). PIKE−/− mice are viable and fertile. However, a significant reduction of white adipose tissues (WATs) was detected in the PIKE−/− mice, whereas no noticeable difference was found in other peripheral tissues (Fig. 1E).

PIKE−/− mice are resistant to diet-induced obesity. When fed a chow diet, the body weight of female PIKE−/− mice was slightly, but significantly, lower at 8 weeks old compared with wild-type mice (17.29 ± 0.27 vs. 16.33 ± 0.29 g, P < 0.05, n = 7, Student t test). The difference was more prominent in mice fed with HFD (55% of calories derived from fat). After HFD feeding for 14 weeks, obesity developed in wild-type but not in PIKE−/− animals (Fig. 2A). Daily food intake of PIKE−/− mice fed a chow diet was normal, but the amount of food intake in PIKE−/− mice was substantially less than that in the control fed HFD (Fig. 2B). Increased body weight was associated with a drastic gain of inguinal WAT weight in wild-type but not in PIKE−/− mice (312% in wild-type vs. 46.5% in knockout) (Fig. 2C). The adipocytes in PIKE−/− mice were also smaller in both feeding conditions (Fig. 2D and E). Moreover, circulating leptin and TNF-α concentrations were lower in PIKE−/− mice (Fig. 2F and G). Expression of PIKE-A was greatly enhanced in the WAT and muscle of mice fed with HFD and the genetically obese (ob/ob) mice (Fig. 2H, first and fifth panels). In contrast, no noticeable alternation of hepatic PIKE expression was detected among all the tested groups (Fig. 2H, third panel), sug-

RESEARCH DESIGN AND METHODS

Generation of knockout animals and genotyping. Heterozygous PIKE−/+ C57BL/6 mice with a targeted deletion of exons 3–6 of CENGT1 were generated under contract by Ozone (Bentley DC, Australia). Genotyping was performed by PCR using genomic DNA isolated from the tail tip. PCR was performed using a combination of primers D (5′-ACAGGACTGCTGCACTCAC-TACGTC-3′), H (5′-CTGGCACCACCTACAGGAGTAG3′), A (5′-TCCAGTGGAC-GGAAGCTTCTG-3′), and C (5′-CCAGAGGCATCTCATGCTTAG3′). Immunoprecipitation and Western blotting. Tissue extracts were prepared by homogenizing the tissues in buffer as reported (18). Immunoprecipi-
tation was performed as described (18). Antibodies used in the Western blot analysis were obtained from Santa Cruz Biotechnology (insulin receptor, Akt) and Cell Signaling Technology (anti-phospho-Thr308 of Akt, anti-phospho- Thr37/40 of AMPK, anti-phosphor-Ser102 of acetyl-CoA carboxylase [ACC], anti-
AMPKα, and anti-ACC).

Southern blot analysis. Southern blot analysis using mouse tail genomic DNA was performed as reported (19).

Analytic procedures. All animal experiments were performed according to the care of experimental animal guidelines from Emory University. Twelve-
week-old female mice were fed with chow or HFD (Research Diets) for 20 weeks. Blood glucose level was measured by ACCU-CHECK Advantage Blood Glucose Meter (F. Hoffmann-La Roche, Basel, Switzerland). Serum insulin was measured by ELISA (Crystal Chem). Serum triglyceride level was measured by Serum Triglyceride Determination Kit (Sigma-Aldrich). Serum tumor necrosis factor-α (TNF-α) was measured by ELISA (BD Biosciences). Glucose tolerance test (GTT) was performed on mice after peritoneal injection of d-glucose (2 g/kg body wt).

In vivo insulin stimulation. Animals (16 h fasting) were anesthetized by intraperitoneal administration of sodium pentobarbital (50 mg/kg body wt). Saline or 5 units human insulin (Eli Lilly) was injected through inferior vena cava. After 5 min, liver, hind limb muscles, and inguinal fat were removed and immediately frozen in liquid nitrogen.

PI 3-kinase assay. In vitro PI 3-kinase assay was performed using anti-p110α (Santa Cruz Biotechnology) as described previously (8).

RT-PCR. Total RNA from various tissues was prepared by Trizol Isolation Reagent (Invitrogen). First-strand cDNA from total RNA was synthesized using Superscript III reverse transcriptase (Invitrogen) and Oligo-dT17 as primer. Amplification of preadipocyte factor 1 (PREF-1), adipocyte protein 2 (aP2), pexinosome proliferator-activated receptor-γ (PPARy), and C BPα was per-
formed using primers mPrel-F (5′-GCACCAACCTGTGACCC-3′), mPrel-R (5′-CAGGACGCTGTGGACCC-3′), map2F (5′-CAAAATGTGATGCTTTCTGTTG-3′), map2-R (5′-ATGCT GTATTGGTGGAAACT-3′), mBPα-F (5′-CTGGAAGCTCGACATATTATGTGTA-3′), and mBPα-R (5′-AAGTCTTACGGGAGAAC-3′). Expression of PIKE-A was determined using primers D and H as described above. Glyceroldehyde-3-phosphate dehydrogenase (GAPDH) was also amplified as internal standard using primers 5′-GGCTTCTGACCTGCGCTATTT-3′ (forward) and 5′-GCGCTTACGCTGGGTCATTT-3′ (reverse).

In vitro H2-deoxoglucose uptake. H2-deoxoglucose uptake in soleus muscle and inguinal fat pad was performed in the presence or absence of human insulin (Eli Lilly) as reported (20).

Fatty acid oxidation assay. Fatty acid oxidation was measured by deter-
mining the production of 14C from [9,10-3H]-palmitate as reported (21).

Hyperinsulinemic-euglycemic clamp and metabolic cage studies. Metabo-
ic cage studies and in vivo glucose metabolisms including glucose infusion rate, glucose turnover rate, and glycogen synthesis were determined by hyperinsulinemic-euglycemic clamp as reported (22,23).

Statistical analysis. Results were considered significant when P < 0.05. Statistical analysis was performed using either Student t test, one-way ANOVA, or two-way ANOVA followed by Tukey multiple comparison test or Bonferroni post-tests using the computer program GraphPad Prism (Graph-

Pad Software).

Detailed experimental procedures are in the supplementary methods (http://diabetes.diabetesjournals.org/cgi/content/full/db09-1404/DC1).
FIG. 1. Targeted disruption of **PIKE**. **A**: Schematic representation of mouse **PIKE** (top), the targeting vector (middle), and the targeted gene region (bottom). The locations of loxP sites were marked as solid triangles and of FRT sites, as solid bars. **B**: Southern blot analysis of progeny produced from heterozygote mating. Genomic DNA was isolated from mouse tail and was digested with *Nhe* I and probed with fragment A as indicated. The 8.5-kb band represents the wild-type allele and the 6-kb fragment corresponds to the knockout allele. **C**: PCR screening of mice from heterozygote mating. Genomic DNA isolated from wild-type (+/H11545/H11545), heterozygous (+/H11545/H11545), and knockout (−/H11546/H11546) mice tail was used in PCR screening. The locations of primers used in the reactions were indicated in **A**. **D**: RT-PCR screening of **PIKE** expression in different tissues. Complementary DNA was synthesized from RNA extracted from various tissues as indicated. Primers D and H as shown in **A** were used in PCR (upper panel). Expression of **GAPDH** was examined as the internal control (lower panel). **E**: Western blot analysis of **PIKE-A**. Proteins extracts of different tissues from wild-type (+/+) and knockout (−/−) mice (3 months old) were prepared, and the expression of **PIKE-A** was detected using specific antibody against the COOH-terminal of human **PIKE-A** (top panel). The amount of tubulin in each sample was examined to demonstrate equal loading (bottom panel). Representative result of three mice from each genotype was shown. **F**: Weight of heart, spleen, pancreas, kidney, and inguinal WAT in 3-month-old mice. The weight was normalized with the total body weight and was expressed as means ± SEM (*n* = 5). Significant reduction of WAT weight was observed in **PIKE**−/− mice (****P < 0.001, Student *t* test).
PIKE knockout mice are resistant to diet-induced obesity. A: Growth curve of 3-month-old wild-type (+/+) and PIKE knockout (-/-) mice fed with HFD. Body weight was measured weekly and expressed as mean ± SEM (n = 7–10; **P < 0.01, ***P < 0.001, two-way ANOVA). B: Food intake by wild-type (+/+) and knockout (-/-) mice fed with chow or HFD was measured in a 3-day period. Results were expressed as mean ± SEM (***P < 0.001 vs. the same genotype; b: P < 0.001 vs. the same diet treatment; one-way ANOVA). C: Weight of inguinal WAT from wild-type (+/+) and PIKE knockout (-/-) mice (8–9 months old, n = 5) that have been fed with chow diet or HFD for 20 weeks. Data were expressed as mean ± SEM (**P < 0.001 vs. the same genotype; b: P < 0.01, c: P < 0.001 vs. the same diet treatment; one-way ANOVA). D: Pictures of hematoxylin-eosin (H&E) staining of inguinal WAT cell area from wild-type (+/+) and PIKE-null (-/-) animals (8–9 months old) that have been fed with chow diet or HFD for 20 weeks (n = 4). Results were expressed as mean ± SEM (**P < 0.001 vs. the same genotype; b: P < 0.01 vs. the same diet treatment; one-way ANOVA). E: Circulating leptin concentration of wild-type (+/+) and PIKE knockout (-/-) mice (8–9 months old) that have been fed with chow or HFD for 20 weeks. Results were expressed as mean ± SEM (n = 4; P < 0.05, ***P < 0.001 vs. the same genotype; a: P < 0.05, b: P < 0.01 vs. the same diet treatment; one-way ANOVA). F: Circulating leptin concentration of wild-type (+/+) and PIKE knockout (-/-) mice (8–9 months old) that have been fed with chow or HFD for 20 weeks. Results were expressed as mean ± SEM (n = 4; P < 0.05, ***P < 0.001 vs. the same genotype; a: P < 0.05, b: P < 0.01 vs. the same diet treatment; one-way ANOVA). G: Circulating TNF-α concentration of wild-type (+/+) and PIKE knockout (-/-) mice (8–9 months old) that have been fed with chow or HFD for 20 weeks. Results were expressed as mean ± SEM (n = 4; P < 0.05, ***P < 0.001 vs. the same genotype; a: P < 0.05, b: P < 0.01 vs. the same diet treatment; one-way ANOVA). H: Elevated PIKE-A expression in the WAT and muscle of diet-induced or genetically obese mice. RNA from WAT, liver, and muscle of ob/ob mice or normal mice (8–9 months old) that have been fed with chow diet or HFD was extracted and reverse transcribed. (A high-quality color representation of this figure is available in the online issue.)

Pre-F-1 was also detected in both genotypes treated with HFD, suggesting the proliferation of preadipocyte was similar in both genotypes (Fig. 3A). However, subsequent adipocyte differentiation was impaired in PIKE-/-/ mice as the expression of aP2, PPARγ, and C/EBPα was greatly reduced.

Next, we sought to determine whether the deletion of PIKE-A per se is sufficient to prevent adipogenesis in vitro. Whereas mouse embryonic fibroblasts (MEFs) from wild-type mice differentiated into adipocytes, as evident by the accumulation of lipid droplets within the cells, PIKE-/-/ MEFs failed to fully differentiate under the same condition (Fig. 3B). A significantly lower amount of oil red O staining was found in PIKE-/-/ MEFs after induction (Fig. 3C). Moreover, expression of mature adipocyte markers aP2, C/EBPα, and PPARγ was lower in PIKE-/-/ MEFs after differentiation (Fig. 3D), which was consistent with the findings in WAT. These results suggest lipatrophy is a direct consequence of PIKE-A deletion in WAT, which may explain the reduced adiposity of mutant mice.

PIKE knockout mice are protected from diet-induced hyperglycemia by enhanced systemic insulin sensitivity. Adipocyte dysfunction is one of the major factors that causes insulin resistance (26); therefore, we examined the effects of diet-induced diabetes in PIKE-/-/ mice. In HFD treatment, hyperglycemia was observed in both genotypes when the animals were fed. Hyperglycemia was sustained in fasted wild-type animals but not in PIKE-/-/ mice (Fig. 4A). Moreover, PIKE-/-/ mice on HFD showed improved glucose tolerance during the GTT (Fig. 4B). A lower amount of insulin was also secreted in PIKE-/-/ mice treated with HFD during the GTT (supplementary Fig. 1C). In parallel, less circulating insulin was detected in PIKE-/-/ mice in both feeding conditions (Fig. 4C), suggesting a higher insulin sensitivity. This notion was further supported by higher glucose infusion rate (Fig. 4D), whole-body glucose turnover (supplementary Fig. 1B), and glycogen synthesis (supplementary Fig. 1C) in PIKE-/-/ mice during the hyperinsulinemic-euglycemic clamp studies. Hepatic insulin resistance was also alleviated in PIKE-/-/ mice fed with HFD (supplementary Fig. 1D), which provides further explanation to the relieved diabetic phenotype in PIKE-/-/ mice because hepatic insulin resistance is associated with diabetes (27). We also examined the insulin-stimulated signaling in tissues responsible for glucose utilization to reveal the molecular basis of the enhanced insulin sensitivity in PIKE-/-/ mice. In mice fed a normal chow diet, comparable tyrosine phosphorylation of insulin receptor occurred in WAT and muscle of both genotypes after in vivo insulin injection (Fig. 4E, first panel). However, insulin provoked higher insulin substrate-1 (IRS-1) phosphorylation, PI 3-kinase activity, and Akt phosphorylation in PIKE-/-/ WAT and muscle (Fig. 4E, second panel). This suggests that PIKE-A is a crucial factor in maintaining insulin sensitivity.

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PIKE-A is an Akt upstream effector, which binds Akt and enhances its kinase activity in glioblastomas (6,13). It is thus anticipated that \( \text{PIKE}^{-/-} \) mice would display diabetic phenotypes as deletion of Akt2 in mice showed impaired glucose tolerance (28). To our surprise, blood glucose level is normal in \( \text{PIKE}^{-/-} \) mice. Because \( \text{Akt1}^{-/-} \) or \( \text{Akt3}^{-/-} \) mice have no obvious defect in glucose homeostasis, the normoglycemic condition in \( \text{PIKE}^{-/-} \) mice could be explained if PIKE-A associates selectively with Akt1 and Akt3 rather than Akt2. As predicted, PIKE-A preferentially bound both Akt1 and Akt3 (supplementary Fig. 2A), suggesting that only Akt1 and Akt3 activities may be altered in \( \text{PIKE}^{-/-} \) tissues. Concurrent with this notion, the brain mass of \( \text{PIKE}^{-/-} \) mice was smaller than the control mice (supplementary Fig. 2B), a phenotype that is specifically observed in Akt3-null animals (29).

**Lipid oxidation is enhanced in \( \text{PIKE}^{-/-} \) mice.** Animal models with lipoatrophy often associate with hyperlipidemia and ectopic lipid accumulation (30). However, significant changes in neither circulating triglyceride (Fig. 5A) nor ectopic lipid depositions in liver (Fig. 5B) were seen in the \( \text{PIKE}^{-/-} \) mice, suggesting the excessively absorbed lipid during HFD feeding in \( \text{PIKE}^{-/-} \) animals may be metabolized rather than deposited as storage. To test this possibility, we first monitored the frequency of animal movements using open-field locomotor assay (31). Whereas the activity in wild-type mice decreased when they adapted to the test cage, physical movement of

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**FIG. 3.** PIKE is essential for adipocyte differentiation A: Impaired adipose gene expression in PIKE-null WAT. RNA from inguinal WAT of wild-type (+/+ ) and knockout (−/− ) mice (8–9 months old) that have been fed chow or HFD for 20 weeks was collected and used in RT-PCR. Expression of preadipocyte markers Pref-1, mature adipocyte marker aP2, transcription factors PPAR\( \gamma \) and C/EBP\( \alpha \) was normalized to GAPDH. Results were expressed as fold induction against the corresponding expression level in wild-type animals fed with chow diet (\( n = 3; * P < 0.05; ** P < 0.01; *** P < 0.001 \), Student \( t \) test). B: Oil red O staining of MEFs isolated from wild-type (+/+ ) and knockout (−/− ) mice before (day 0) and after (day 8) induced adipocyte differentiation. Scale bar represents 50 μm. Representative result of three independent experiments is shown. C: Quantification of accumulated lipid in MEFs before (day 0) and after (day 8) isobutylmethylxanthine-dexamethasone insulin (MDI) induction. Oil red O in the MEFs was extracted by isopropanol and measured in optical density 500 nm (\( * * P < 0.01; *** P < 0.001 \) vs. noninduced control of the same genotype; \( t \) test). D: Impaired adipose gene expression in PIKE-null MEFs. RNA from MEFs of wild-type (+/+ ) and knockout (−/− ) mice before (day 0) and after (day 8) induced adipocyte differentiation. Expression of mature adipocyte marker aP2 and transcription factors PPAR\( \gamma \) and C/EBP\( \alpha \) was normalized to GAPDH. Results were expressed as fold induction against the corresponding expression level in day-0 wild-type MEFs (\( n = 3; * P < 0.05; *** P < 0.001 \), Student \( t \) test). (A high-quality digital representation of this figure is available in the online issue.)
PIKE knockout mice (−/−) mice exhibited significantly lower glucose levels when fed a high-fat diet (HFD) for 20 weeks, compared to wild-type (+/+). These results were confirmed through a glucose tolerance test, where the knockout mice showed improved glucose metabolism. The AMPK expression in brown adipose tissue (BAT) and white adipose tissue (WAT) was also assessed, revealing that AMPK expression was reduced in both tissue types after HFD feeding in wild-type mice, whereas it remained unchanged in the knockout mice. This suggests that AMPK plays a crucial role in glucose metabolism and lipid oxidation, which is further supported by the high lipid oxidation observed in the knockout mice. The phosphorylation of ACC was reduced after HFD feeding in wild-type mice, but not in the knockout mice. This indicates that AMPK activation is essential for normal lipid metabolism. The results also suggest that PI3K and Akt signaling pathways are involved in the regulation of lipid metabolism and glucose homeostasis. The study provides valuable insights into the role of AMPK in obesity and diabetes, and the potential therapeutic targets for treating these conditions.
Liver that have been fed with chow or HFD for 20 weeks. Extracts of inguinal WAT, BAT, and muscle were prepared and immunoblotted with genotype, one-way ANOVA, for 20 weeks (Fig. 5). Analysis in Fig. 5 (E) mice. In agreement with the immunoblotting analysis (Fig. 5E), elevated fatty acid oxidation in cultured (upper panel) and oil red O (lower panel) staining of liver sections collected from wild-type (+/+) and knockout (−/−) mice (8–9 months old) that have been fed with HFD for 20 weeks. Scale bar represents 50 μm. Representative result of three mice from each genotype is shown. C: Spontaneous activity of wild-type (+/+) and PIKE knockout (−/−) mice fed with chow or HFD for 14 weeks. Results were expressed as mean ± SEM (n = 7; *P < 0.05, ***P < 0.001 vs. the same diet). D: Oxygen consumption (top panel), CO2 release (middle panel), and respiratory exchange ratio (bottom panel) in wild-type (solid bar) and knockout (open bar) mice fed with chow and HFD for 20 weeks (n = 4). Results were expressed as mean ± SEM (*P < 0.01, ***P < 0.001 vs. the same diet; a: P < 0.05, c: P < 0.001 vs. the same genotype, one-way ANOVA, n = 4). E: Analysis of ACC and AMPK phosphorylation in wild-type (+/+) and knockout (−/−) mice (8–9 months old) that have been fed with chow or HFD for 20 weeks. Extracts of inguinal WAT, BAT, and muscle were prepared and immunoblotted with phospho-Thr172-AMPK, phospho-Ser173-ACC, total AMPK, and ACC antibodies. F: Elevated fatty acid oxidation in PIKE−/− muscle cells. Rate of [H]-palmitate oxidation was measured in cultured skeletal muscle cells and hepatocytes isolated from wild-type (+/+) and PIKE knockout (−/−) mice. Results were expressed as mean ± SEM (*P < 0.05, one-way ANOVA, n = 5). (A high-quality digital representation of this figure is available in the online issue.)

In contrast, no significant changes in hypothalamic AMPK and ACC phosphorylation were found in PIKE−/− animals in both feeding conditions (supplementary Fig. 3A). Moreover, AMPK and ACC phosphorylation was not enhanced in PIKE−/− MEFs (supplementary Fig. 3B). These results suggest a tissue-specific effect of PIKE-A in modulating AMPK and ACC activity. We have also performed the fatty acid oxidation assay in cultured PIKE−/− muscle cells and hepatocytes. In agreement with the immunoblotting analysis in Fig. 5E, an elevated palmitate oxidation rate was detected in PIKE−/− muscle cells but not hepatocytes (Fig. 5F).

Therefore, the high physical activity of PIKE−/− mice and enhanced lipid oxidation in BAT, WAT, and muscle may account for their lean phenotype during HFD feeding. The higher AMPK phosphorylation in PIKE−/− WAT also provides a possible explanation for the defective adipogenesis observed, because prolonged AICAR-induced AMPK activation inhibits adipocyte differentiation by diminishing PPARγ and C/EBPα expressions (33,34). Furthermore, agonist-activated AMPK potentiates the insulin-stimulated glucose uptake by activating IRS-1 (35,36), which may explain the enhanced PI 3-kinase and Akt activities in the muscle and WAT of PIKE−/− mice.
**FIG. 6. PIKE interacts with insulin receptor and is essential for insulin-suppressed AMPK phosphorylation.**

**A:** Diagrammatic representation of various myc-tagged PIKE-A truncates. PIKE-A is a GTPase containing a short NH$_2$-terminal, a GTPase domain (GTPase) for hydrolysis of GTP, a PH domain for phosphoinositol lipid interaction, and a COOH-terminal region (GAP-ANK) with sequence homology to ARF/GAP protein and ankyrin repeats.

**B:** Mapping of the insulin receptor interaction domain in PIKE-A. Various myc-tagged PIKE-A truncation mutants as shown in **A** and His-tagged insulin receptor (His-IR) were cotransfected into HEK293 cells. The myc-tagged proteins were immunoprecipitated and the associated insulin receptor was detected using anti–insulin receptor antibody (top panel). Expression of His-IR (middle panel) and various myc-tagged proteins (lower panel) was also detected.

**C:** Fyn phosphorylation of PIKE-A is important for insulin receptor interaction. HEK293 PIKE AND OBESITY 890 DIABETES, VOL. 59, APRIL 2010 diabetes.diabetesjournals.org
PIKE-A is essential for insulin-suppressed AMPK phosphorylation. Next, we sought to clarify the role of PIKE-A in modulating AMPK phosphorylation. Fyn knockout (Fyn−/−) mice are lipodystrophic with enhanced AMPK activity in muscle and WAT (37). These metabolic characteristics highly resemble the phenotypes of PIKE−/− mice. Given that PIKE-A is a substrate of Fyn (38) and Fyn interacts with IRS-1 in an insulin-dependent manner (39), we hypothesized that PIKE-A may form a complex with Fyn and insulin receptor upon insulin stimulation, which is essential for insulin to suppress AMPK activity (40,41). In HEK293 cells, PIKE-A associated with insulin receptor through its NH2-terminal (1–72 amino acids) (Fig. 6A and B), in which their interaction could be enhanced by insulin stimulation (Fig. 6C, first panel). However, this interaction was abolished when the Fyn phosphorylation site (Tyr682 and Tyr774) in PIKE-A (PIKE-A YY) was mutated (Fig. 6C, first panel), suggesting Fyn phosphorylation is critical to the formation of PIKE-A/insulin receptor complex. The kinetics of insulin receptor/PIKE-A complex formation inversely correlated with the phosphorylation of AMPK (Fig. 6D, first and sixth panels). Remarkably, neither Thr172 nor Ser485/Ser491 phosphorylation (42) was altered by insulin in GST-PIKE-A YY-transfected cells (Fig. 4D, sixth and seventh panels), suggesting that PIKE-A binding to insulin receptor is critical to mediate the inhibitory action of insulin on AMPK phosphorylation. On the other hand, Akt phosphorylation was not affected in either wild-type PIKE-A or PIKE-A YY cells in response to insulin (Fig. 6D, fourth panel). The formation of PIKE/insulin receptor/Fyn complex was further demonstrated in muscle tissue. In vivo insulin injection in wild-type mice enhanced the formation of PIKE-A/insulin receptor complex, which was substantially reduced in Fyn−/− tissue (Fig. 4G, first panel). This complex was not detected in PIKE-null tissues (Fig. 6E, first panel). Our immunoprecipitation results also confirmed that the association of Fyn and insulin receptor in muscle is insulin dependent (Fig. 6E, second panel). Furthermore, the formation of Fyn/insulin receptor complex was not affected in PIKE-null tissues, suggesting that PIKE-A is not essential for their interaction (Fig. 6E, second panel). Thus, the interaction between PIKE-A and insulin receptor is important for insulin to suppress AMPK phosphorylation, which provides a possible explanation to the enhanced AMPK phosphorylation in PIKE−/− WAT and muscle.

**DISCUSSION**

One of the major findings in the current report is that PIKE-A is critical for adipocyte differentiation. Several lines of evidence support the role of PIKE-A in terminal adipocyte differentiation instead of preadipocyte formation. First, the mature adipocyte marker aP2 is significantly decreased during in vitro adipocyte differentiation in PIKE−/− MEFs, indicating PIKE-A is important for adipocyte differentiation (Fig. 3B and C). Second, PIKE-A expression is increased in fat tissue development of HFD-fed and ob/ob mice, which highlights its function in the process (Fig. 2H). Lastly, HFD induced comparable preadipocyte marker Pref-1 expression in both wild-type and PIKE−/− mice, indicating that formation of new adipocytes is normal in PIKE-null adipose tissue (Fig. 3A). Interestingly, we found a small portion of PIKE−/− MEFs was able to differentiate into mature adipocytes (Fig. 3B), and quantitative analysis revealed a small but statistically significant increment of lipid accumulation in PIKE−/− MEFs (Fig. 3C). This result indicates that a PIKE-A–independent mechanism is responsible for some adipocyte differentiation, which also accounts for the existence but not completely the absence of adipose tissue in PIKE−/− mice.

Ectopic lipid storage due to adipocyte differentiation defect is associated with hyperlipidemia and liver steatosis (43). However, we could not detect these pathologic conditions in PIKE−/− mice (Fig. 5A and B). It is thus reasonable to predict that the lipid spillover from adipocyte is metabolized in mutant animals. Our results that PIKE-null fat and muscle have significantly elevated AMPK and ACC phosphorylation suggest an elevated β-oxidation in these tissues (Fig. 5E), which is further supported by the enhanced fatty acid oxidation rate in the in vitro assay (Fig. 5F) and the low respiratory exchange ratio values (Fig. 5D). AMPK has been viewed as a fuel sensor for glucose and lipid metabolism. Once activated, AMPK initiates a concomitant inhibition of energy-consuming biosynthetic pathways and activation of ATP-producing pathways such as fatty oxidation in mitochondria (44). As a result, most of the lipids absorbed in PIKE−/− mice are oxidized as the energy source, which accounts for the lean phenotype during the HFD treatment.

The upregulated phosphorylation of AMPK and its downstream substrate ACC in PIKE-null muscle and adipose tissues indicates that PIKE-A negatively regulates the activities of these enzymes. This notion is further supported by the fact that PIKE-A is critical for insulin to inhibit AMPK phosphorylation in 293 cells (Fig. 6D). This upregulation of AMPK activity in PIKE−/− muscle and fat also provides a possible mechanism accounting for the elevated systemic insulin sensitivity, as AMPK and insulin signaling are intimately connected. Agonist-induced AMPK activation increases the glucose uptake in muscle (45). It also potentiates the insulin-stimulated glucose uptake by activating IRS-1 (35,36). A similar observation was made in...
adipose tissue that treatment of adipocytes with AMPK agonist AICAR enhanced basal glucose uptake by increasing GLUT4 translocation (46). Long-term activation of AMPK in mice increases the systemic insulin sensitivity and protects animals from HFD-induced obesity and diabetes (34,47), which is in agreement with our observations in PIKE−/− mice. It is noteworthy that the alleviated insulin resistance in PIKE−/− mice after HFD treatment may be a result of reduced inflammation. Because reduced circulating TNF-α could improve insulin sensitivity and increase AMPK activity (48,49), the low blood TNF-α in PIKE−/− mice (Fig. 2G) may also contribute significantly to improve the diet-induced insulin resistance.

Because the whole-body–knockout mice were used in the present study, we cannot exclude the possibility that deletion of PIKE-A in the brain causes a central effect to modify whole-body activity and metabolism. Because brain is the major site to control appetite and body weight (50), where PIKE is highly expressed (Fig. 1D), it is reasonable to suspect that reduced food intake (Fig. 2H) and elevated physical activity (Fig. 5C) in PIKE−/− mice are the primary causes of lean phenotype during HFD treatment. However, our data strongly support that peripheral ablation of PIKE-A does play a role in preventing obesity development. First, the feeding behavior is comparable between wild-type and PIKE−/− mice fed a chow diet, when lipatrophy is already obvious. Second, induced differentiation in PIKE−/− MEFs is greatly impaired (Fig. 3B), suggesting ablation of PIKE per se in MEFs is adequate to suppress adipogenesis. Third, PIKE-A interacts with the insulin receptor in a Fyn-dependent manner, which is essential for insulin-induced AMPK phosphorylation in muscle (Fig. 6D). Deletion of PIKE in muscle, therefore, would enhance the AMPK phosphorylation and lipid oxidation (Fig. 5E and F). Lastly, isolated PIKE−/− WAT and muscle, in which the metabolic influence by the brain is eliminated, have higher 3H-2-deoxyglucose uptake when stimulated by insulin (Fig. 4F).

Our data also suggest that the function of PIKE-A is not restricted to enhance Akt activity alone. We have demonstrated that PIKE-A physically interacts with the insulin receptor, which is important for insulin to suppress AMPK phosphorylation. Our data also provide a novel mechanistic insight into the phenotypes observed in Fyn−/− mice (37), as PIKE-A/insulin receptor association is Fyn dependent. Conceivably, PIKE-A is a downstream target of Fyn that inhibits the activity of AMPK during obesity development. Thus, PIKE-A may represent an additional regulatory point, in addition to Akt, for insulin to suppress AMPK phosphorylation.

In all, our results uncover the novel physiological functions of PIKE-A, which plays important roles in obesity development and the accompanied insulin resistance by regulating AMPK activities negatively. Consequently, less fat is deposited and the associated insulin resistance is ameliorated. Therefore, PIKE-A may represent a potential therapeutic target for obesity and the adjunct insulin resistance.

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