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Sheng Zhong Duan, … , David S. Milstone, Richard M. Mortensen


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Hypotension, lipodystrophy, and insulin resistance in generalized PPARγ-deficient mice rescued from embryonic lethality

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We rescued the embryonic lethality of global PPARγ knockout by breeding Mox2-Cre (MORE) mice with floxed PPARγ mice to inactivate PPARγ in the embryo but not in trophoblasts and created a generalized PPARγ knockout mouse model, MORE-PPARγ knockout (MORE-PGKO) mice. PPARγ inactivation caused severe lipodystrophy and insulin resistance; surprisingly, it also caused hypotension. Paradoxically, PPARγ agonists had the same effect. We showed that another mouse model of lipodystrophy was hypertensive, ruling out the lipodystrophy as a cause. Further, high salt loading did not correct the hypotension in MORE-PGKO mice. In vitro studies showed that the vasculature from MORE-PGKO mice was more sensitive to endothelial-dependent relaxation caused by muscarinic stimulation, but was not associated with changes in eNOS expression or phosphorylation. In addition, vascular smooth muscle had impaired contraction in response to α-adrenergic agents. The renin-angiotensin-aldosterone system was mildly activated, consistent with increased vascular capacitance or decreased volume. These effects are likely mechanisms contributing to the hypotension. Our results demonstrated that PPARγ is required to maintain normal adiposity and insulin sensitivity in adult mice. Surprisingly, genetic loss of PPARγ function, like activation by agonists, lowered blood pressure, likely through a mechanism involving increased vascular relaxation.

Introduction
The metabolic syndrome (MetS) is a cluster of metabolic abnormalities (including obesity, insulin resistance, and dyslipidemia) associated with type 2 diabetes, hypertension, and atherosclerosis. It is considered one of the most serious medical conditions, with increasing threats to public health in modern societies (1, 2). The nuclear transcription factor PPARγ is a candidate for mediating the interface of metabolic and cardiovascular diseases (3). PPARγ can affect most, if not all, aspects of the MetS. It is highly expressed in adipocytes and important in controlling adipogenesis as well as gene expression in these cells (4, 5). Results of genetic deficiency studies have shown that PPARγ is required for adipogenesis (6–8). The PPARγ-agonist thiazolidinediones (TZDs) increase insulin sensitivity, lower blood glucose, decrease circulating free fatty acids and triglycerides, lower blood pressure, and reduce atherosclerosis in insulin-resistant patients and animal models (1, 3, 9). However, the role of PPARγ in these TZD-induced effects is not completely understood. Dominant-negative mutations of PPARγ (Pro467Leu and Val290Met) in humans are associated with insulin resistance, diabetes, and hypertension (10), whereas a similar mutation (Pro467Leu) in mice causes hypertension and fat redistribution but not insulin resistance or diabetes (11). The PPARγ Pro12Ala polymorphism in humans is associated with type 2 diabetes, insulin resistance, and obesity (12), while the Pro12Ala mutation of PPARγ2 in mice causes lipodystrophy and mild glucose intolerance (13).

In order to explore the physiologic importance of PPARγ in the MetS, recent studies using conditional knockout strategies in mice have revealed the roles of PPARγ in individual tissues. Adipocyte PPARγ is required for normal adiposity (14–16) and insulin sensitivity in fat and liver, but not in muscle (15). Myocyte PPARγ is required to maintain whole-body insulin sensitivity (17) or liver insulin sensitivity only (18). Hepatic PPARγ is required to maintain insulin sensitivity, particularly in older animals or in animals with genetically diabetic backgrounds (19, 20). Endothelial PPARγ is important in regulating diet-induced hypertension (21).

However, the effects of global deletion — including the multifaceted interaction PPARγ may have in multiple tissues — have not yet been addressed because of the lethality of the germline knockout of PPARγ (7, 8). The MetS is a complex disease involving interactions among different tissues. In order to better understand the function of PPARγ in this disease, a generalized knock-
out model is necessary. The whole-body knockout of PPARγ2 (22) is mostly restricted to adipocytes (5). Here we describe a generalized PPARγ-knockout (both PPARγ1 and PPARγ2) mouse model, Mox2-Cre–PPARγ knockout (MORE-PGKO), which rescues embryonic lethality by preserving PPARγ expression in the trophoblasts. Our data showed that global PPARγ deficiency caused lipodystrophy and insulin resistance, which was surprisingly accompanied by lower blood pressure. In contrast, A-ZIP mice, which expressed a dominant negative protein A-ZIP/F that caused lipodystrophy (23), were hypertensive, establishing them as a model of lipodystrophic hypertension comparable to that seen in humans. Thus, both genetic loss of PPARγ function and activation by pharmacologic agonist had similar effects on blood pressure.

Results

The MORE-PGKO mouse model. MORE-PGKO mice were born in the expected Mendelian ratio, but only approximately 10% survived to adulthood. In MORE mice, Cre recombinase is uniformly expressed in epiblast-derived tissue, but not other tissues (24), thus sparing trophoblast. Southern blot analysis revealed an 8-kb restriction fragment representing the PPARγ-null allele, which indicated successful Cre-mediated recombination (6), in multiple tissues of MORE-PGKO mice (Figure 1). Tissues from littermate control (LC) mice revealed 14- and 10-kb restriction fragments representing WT and floxed PPARγ alleles, respectively (6), which indicated lack of recombination. Among all tissues tested in MORE-PGKO mice, approximately 94.5% ± 4.3% and 95.4% ± 6% of the floxed PPARγ allele was recombined to the null allele in females and males, respectively.

Generalized PPARγ deficiency causes lipodystrophy and organomegaly. PPARγ has previously been shown to regulate adipocyte differentiation in vitro (6, 7), and mice with adipocyte-specific PPARγ deficiency have less adipose tissue than do WT controls, although a substantial amount of adipose tissue remains (14, 15). Furthermore, PPARγ is required for adipocyte survival in adult mice (16). MORE-PGKO mice, in contrast, showed more severe lipodystrophy (Figure 2). Dual-energy x-ray absorptiometry (DEXA) scan showed that MORE-PGKO mice had dramatically less fat mass than did LC mice at 6 months of age (Figure 2A). Both male and female MORE-PGKO mice lacked interscapular brown adipose tissue as well as mesenteric, perirenal, and subcutaneous fat (Figure 2, B–D, and data not shown). Epididymal fat and perivarien fat was also absent in male and female MORE-PGKO mice, respectively (Figure 2, B and C). However, female MORE-PGKO mice had a small amount of perituene fat, approximately 9% that of LC mice (Figure 2, B and C). Quantitative RT-PCR showed that PPARγ was expressed, albeit at very low level, in the perituerine fat of female MORE-PGKO mice (Figure 2E), indicating at least partial escape from Cre-loxP-mediated recombination in this depot.

In spite of this lipodystrophy, body weight and length were comparable in MORE-PGKO and LC mice (Table 1). The adrenal gland, kidney, liver, lung, pancreas, thyroid, and heart weights of MORE-PGKO mice were all significantly greater than those of LC mice (Table 1). This organomegaly was similar to that seen in A-ZIP mice, another lipodystrophic mouse model (23).

Insulin resistance in MORE-PGKO mice. Previous studies using tissue-specific knockout strategies have demonstrated that adipocyte, myocyte, and hepatocyte PPARγ all play important roles in maintaining insulin sensitivity and glucose homeostasis (15, 17–20). We evaluated insulin sensitivity and glucose homeostasis in MORE-PGKO and LC mice at 6 months of age. Insulin tolerance test (ITT) showed that both male and female MORE-PGKO mice had impaired insulin sensitivity compared with LC mice (Figure 3, A and D). They had significantly elevated fasting plasma insulin levels (Figure 3, B and E), also demonstrating insulin resistance. Histologic analysis revealed enlarged pancreatic islets in MORE-PGKO mice (Figure 3, C and F). Male MORE-PGKO mice were diabetic, with impaired glucose tolerance test (GTT; Figure 4A) and elevated fed blood glucose (Figure 4B). However, female MORE-PGKO mice were not diabetic (Figure 4E) and showed enhanced glucose tolerance (Figure 4D) compared with LC mice, probably because of higher plasma insulin levels after overnight food deprivation and during GTT in female MORE-PGKO mice (Figure 4F). Conversely, male MORE-PGKO mice showed

Table 1

<table>
<thead>
<tr>
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<th>LC</th>
<th>MORE-PGKO</th>
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</thead>
<tbody>
<tr>
<td>Body weight (g)</td>
<td>34.5 ± 5.0</td>
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<td>Body length (mm)</td>
<td>91.0 ± 6.2</td>
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<td>Adrenal gland (mg/g)</td>
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<td>Kidney (mg/g)</td>
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<td>18.3 ± 0.4</td>
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<td>49.4 ± 6.6</td>
<td>69.7 ± 18</td>
<td>0.0012</td>
</tr>
<tr>
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<td>5.3 ± 0.5</td>
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<tr>
<td>Pancreas (mg/g)</td>
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<td>14.8 ± 1.8</td>
<td>0.0023</td>
</tr>
<tr>
<td>Thyroid (mg/g)</td>
<td>0.07 ± 0.01</td>
<td>0.12 ± 0.04</td>
<td>0.012</td>
</tr>
<tr>
<td>Heart (mg/g)</td>
<td>3.9 ± 0.2</td>
<td>4.3 ± 0.2</td>
<td>0.017</td>
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</table>

Individual organ weights are expressed relative to body weight. Results were similar for female mice. n = 8–11 per group.

Figure 1

MORE-PGKO mice show extensive recombination. Representative Southern blot results of tissues from female LC and MORE-PGKO (KO) mice. The 14-kb endogenous WT allele appeared only in LC mice. The 8-kb Cre-recombined null allele appeared only in MORE-PGKO mice. The 10-kb floxed allele was rare in MORE-PGKO mice. A, aorta; B, brain; F, fat; H, heart; I, intestine; Li, liver; Lu, lung; M, skeletal muscle; P, pancreas; K, kidney; Sp, spleen; St, stomach.
significantly lower plasma insulin levels than did LC mice during GTT (Figure 4C).

Both male and female MORE-PGKO mice had significantly decreased plasma leptin and adiponectin levels (Figure 5, A and B) as well as elevated plasma nonesterified fatty acids (NEFAs) and triglycerides (Figure 5, C and D) compared with LC mice, which can contribute to insulin resistance. Although adipose-specific and skeletal muscle–specific PPARγ knockout mice both displayed fatty liver as well as insulin resistance (15, 17), the livers of MORE-PGKO mice were histologically similar to those of LC mice, without apparent lipid accumulation, hepatocellular damage, or inflammation (Figure 6). This is consistent with previ-
ous data showing that liver-specific PPARγ deficiency protects against hepatic steatosis in leptin-deficient mice and lipodystrophic A-ZIP mice (19, 20).

Hypotension in MORE-PGKO mice. Dominant-negative mutations of PPARγ cause hypertension, a cardinal characteristic of the MetS (1, 2), in both humans and mice (10, 11). TZDs also decrease blood pressure in insulin-resistant patients and animal models (25–27). We therefore sought to determine the consequences of PPARγ deficiency on blood pressure regulation. Blood pressure and heart rate were monitored by radiotelemetry. Surprisingly, despite severe insulin resistance, both systolic and diastolic blood pressure of MORE-PGKO mice were significantly lower than those of LC mice (Figure 7, A and B). In addition, the heart rates of MORE-PGKO mice were significantly higher than those of LC mice (Figure 7C).

Dietary salt loading caused small, nearly identical blood pressure increases in both MORE-PGKO and LC mice (Figure 8) that were consistent with previously reported changes in WT mice (28). Compared with LC mice, MORE-PGKO mice remained hypotensive with a high-salt diet (Figure 8). Salt loading did not significantly change heart rate in either genotype (data not shown). Food consumption (both normal and high-salt diets) was not significantly different between MORE-PGKO and LC mice.

In order to determine whether hypotension was caused by impaired vasoactivity, we tested the responses of aortic rings to graded levels of phenylephrine and acetylcholine. Aortic rings from MORE-PGKO mice showed significantly lower maximum contraction in response to phenylephrine than did rings from LC mice (Figure 9A), although –log[EC50] values were not significantly different between groups (LC, 7.46 ± 0.26; MORE-PGKO,
7.25 ± 0.93; \( P = 0.29 \)). This suggests decreased vascular smooth muscle contractility in MORE-PGKO mice. In addition, the dose-response curve for MORE-PGKO mouse aortic rings responding to acetylcholine significantly shifted to the left compared with the curve for LC mice (–log[EC50], LC, 7.97 ± 0.16; MORE-PGKO, 8.32 ± 0.42; \( P = 0.024 \); Figure 9B), indicating that MORE-PGKO mouse aortas are more sensitive to relaxation by acetylcholine than are those of LC mice.

Overexpression of eNOS results in hypotension (29), and the PPAR\( \gamma \) agonist rosiglitazone increases NO production in cultured endothelial cells (potentially by increasing eNOS activity), reflected by increased phosphorylation of eNOS at Ser1177 (30). In order to determine whether eNOS contributes to the hypotension phenotype, we examined its expression and phosphorylation (Ser1177) in aortas from MORE-PGKO and LC mice. The mRNA and protein phosphorylation levels of eNOS were not significantly different between MORE-PGKO and LC mice (Figure 10). MORE-PGKO mouse aortas had normal histology and expression of macrophage marker F4/80 compared with that of LC mice (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI28859DS1).

Renin-angiotensin-aldosterone system (RAS) is important in blood pressure regulation and vascular volume control (31). Renal collecting duct–specific PPAR\( \gamma \) knockout mice have either elevated (32) or normal plasma aldosterone levels (33). Both plasma aldosterone levels and plasma renin activity (PRA) were increased in MORE-PGKO mice compared with LC mice (Figure 11, A and B). However, the RAS gene expression, including that of angiotensinogen (Agt) and renin, was not changed in kidney, liver, or fat (Figure 11, C–E).

Although lipodystrophy is associated with hypertension in humans (34, 35), to our knowledge, this association has not previously been tested in mice. We therefore measured blood pressure
in A-ZIP mice (23), another lipodystrophic model. A-ZIP mice had higher systolic and diastolic blood pressure and lower heart rate than did WT mice (Figure 12), indicating that hypotension in MORE-PGKO mice is not attributable to lipodystrophy.

Discussion

Recent in vivo studies using conditional knockout mouse models have established the importance of PPARγ in different aspects of the MetS (14–21). However, when investigating the mechanisms of diseases involving multiple organs, a cell type–specific approach may not be sufficient. Here we developed and studied a global knockout mouse model of PPARγ (both PPARγ1 and PPARγ2), MORE-PGKO mice, in order to understand the integrative roles of PPARγ in the complex of MetS.

MORE-PGKO mice had severe lipodystrophy, which further supports the regulatory role of PPARγ in adiposity in vivo. Female MORE-PGKO mice had a small amount of peritoneal fat, but male MORE-PGKO mice had no detectable fat depots. This probably reflects escape from Cre-loxP–mediated recombination in the peritoneal fat of female MORE-PGKO mice, although recombination was similar in other tissues from males and females. Another possibility is that this is related to effects of estrogen on growth of specific adipose depots (36). Organomegaly in MORE-PGKO mice was similar to that previously reported in lipodystrophic A-ZIP mice (23), suggesting a link between organomegaly and lipodystrophy. However, we have previously shown that a lack of PPARγ specifically in cardiomyocytes results in cardiac hypertrophy without lipodystrophy (37). In addition, PPARγ deficiency specifically in pancreatic islets causes islet hyperplasia (38). Furthermore, PPARγ agonists can inhibit proliferation of vascular and airway smooth muscle cells (39, 40), bile ducts (41), and kidney fibroblasts (42) as well as a variety of cancer cells in vitro and/or in vivo (43, 44). These findings suggest that PPARγ can act as an intrinsic growth inhibitor and that generalized PPARγ deficiency might cause generalized organomegaly independent of lipodystrophy. However, whether PPARγ plays a similar growth-inhibitory role in organs other than the heart and pancreas remains to be determined.

The severe insulin resistance we observed in MORE-PGKO mice demonstrated that PPARγ is a key regulator of insulin sensitivity. However, insulin resistance and glucose intolerance are not always associated (13). Male, but not female, MORE-PGKO mice were diabetic. In fact, female MORE-PGKO mice showed greater glucose tolerance than did female LC mice, likely because of much higher plasma insulin levels and changes with fasting during GTT. In addition, MORE-PGKO mice lacked hepatic steatosis, another feature observed in mice with insulin resistance, including mice with adipose-specific and muscle-specific PPARγ deficiency (15, 17). This probably reflects the protective effect of hepatic PPARγ deficiency that has been previously shown to improve hepatic steatosis in leptin-deficient (ob/ob) mice (19) and lipodystrophic A-ZIP mice (20), both of which are insulin resistant.

Lipodystrophy and insulin resistance are both associated with hypertension, another component of the MetS, in humans (34, 35). Our results demonstrated the association of hypertension, lipodystrophy, and insulin resistance in A-ZIP mice. Intriguingly, the MORE-PGKO mice were lipodystrophic and insulin resistant but hypotensive, suggesting that hypertension is
separable from lipodystrophy or insulin resistance. Paradoxically, the PPARγ-agonist TZDs have previously been shown to decrease blood pressure (25–27). Knockout phenotypes that mimic agonist phenotypes suggest that blood pressure control by PPARγ is likely mediated by gene suppression that both agonist and genetic deficiency are capable of relieving. The phenomenon that deletion of PPARγ as well as its agonists have similar effects resembles the previous finding that pioglitazone, a TZD, and PPARγ knockout both increase sodium absorption in cultured inner medullary collecting ducts (32). Another example is the effects of PPARδ, another member of the PPAR family, in atherogenic inflammation: inactivation of PPARδ as well as its agonist has an antiinflammatory effect by dissociating PPARδ from transcriptional repressor BCL-6 (45).

It is unlikely that the hypotension in MORE-PGKO mice is because of lower sympathetic drive caused by lower plasma leptin levels (46) because their heart rates were significantly higher than those of LC mice. Both aldosterone levels and PRA were mildly elevated in the plasma of MORE-PGKO mice, indicating activation of the RAS. The failure of salt loading to correct the hypotension in MORE-PGKO mice suggests that a simple lack of salt retention is not responsible for the phenotype, consistent with the normal blood pressure observed in collecting duct–specific PPARγ knockout mice (33). Therefore, increased vasorelaxation and decreased vasoconstriction are likely significant contributors to the hypotension, although a combination with mild renal effects cannot be ruled out.

Baseline blood pressure is normal in mice with endothelial cell–restricted PPARγ deficiency (21), although a different method was used to measure blood pressure than the one used in the current study. It is likely that PPARγ in other cell types, such as vascular smooth muscle cells, are more important in mediating the effect of PPARγ on blood pressure regulation. Further investigation of mice with smooth muscle cell–specific PPARγ deficiency would help clarify this situation.

eNOS is an important regulator of blood pressure and vascular reactivity (29), but changes in expression or phosphorylation were not found in MORE-PGKO aortas. The involvement of macrophages in the vasculature of MORE-PGKO mice is also unlikely to contribute to the hypotension phenotype, because no differences were found.

In summary, we demonstrated that allowing PPARγ expression in trophoblasts was sufficient to rescue embryonic lethality and confirmed that PPARγ was necessary for adipogenesis and normal insulin sensitivity in mice. Surprisingly, PPARγ deficiency caused increased vascular relaxation and decreased contraction, contributing to hypotension rather than the expected hypertension associated with lipodystrophy. The MORE-PGKO mouse model will provide a useful tool for further investigating the extent to which PPARγ mediates the effects of TZDs in the MetS.

**Figure 7**
Hypotension in MORE-PGKO mice. Systolic (A) and diastolic (B) blood pressure and heart rate (C) of MORE-PGKO and LC mice. MORE-PGKO mice showed significantly lower systolic and diastolic pressure and higher heart rates than did LC mice. Results were similar in both genders and were combined (n = 8, 4 males and 4 females, for each genotype). Data collected over 2 weeks were averaged and presented as light/dark cycle over a 24-hour period. *P < 0.05, **P < 0.01 versus LC, Bonferroni post-test.
Our results also establish A-ZIP mice as a model of hypertensive lipodystrophy similar to that of humans.

Methods

Materials. Reagents were purchased from Sigma-Aldrich unless otherwise specified.

Animals. In MORE mice (24), Cre recombinase is expressed and enzymatically active in the germline. In order to obtain appropriate WT LC mice, MORE mice were first bred to floxed PPARγ mice (47) to obtain heterozygous PPARγ mice (floxed PPARγ/WT, MORE-positive/WT; designated PPARγ<sup>F</sup>/WT,MORE<sup>+/WT</sup> and are functionally PPARγ<sup>F</sup>/WT,MORE<sup>+/WT</sup> in the germline), which were further bred to homozygous floxed PPARγ mice to generate both MORE-PGKO mice (floxed PPARγ/null, MORE-positive/WT; designated PPARγ<sup>F</sup>/–,MORE<sup>+/WT</sup> and LC mice (PPARγ<sup>F</sup>/WT,MORE<sup>+/WT</sup>/WT). These mice were on a mixed C57BL/6, NIH Swiss, and 129SvEv background. A-ZIP lipodystrophic mice on FVB/N background were produced as described previously (23). All animal protocols were approved by the University Committee on Use and Care of Animals of the University of Michigan and by the Standing Committee On Animals of Harvard Medical School.

Southern blot. Genomic DNA was isolated from various tissues of MORE-PGKO and LC mice, digested with BamHI, separated by electrophoresis, transferred to nylon membrane, and hybridized with a <sup>32</sup>P-labeled DNA probe derived from the murine PPARγ gene (3′-probe) as described previously (37, 47).

Fat mass estimation. DEXA (Norland Medical Systems Inc.) scanning was used to evaluate the ratio of fat mass to total mass in 6-month-old MORE-PGKO and LC mice (n = 8–11 for separate male and female groups of each genotype). Macroscopic and histologic analyses of adipose tissue were performed to visualize fat mass. Epididymal fat pads, perirenal fat pads, and interscapular brown adipose tissue were weighed. The ratios of these fat pad weights to body weight were calculated.

Histologic analysis. Tissues were fixed in 10% neutral buffered formalin overnight and embedded in paraffin. Sections (5 μm) were stained with H&E. Digital images were obtained using an Olympus BX51 microscope (Olympus).

Gene expression analysis. The following genes were assayed using quantitative RT-PCR as described previously (48): PPARγ, eNOS, renn, Agt, and β-actin. All primer-probe sets were purchased from Applied Biosystems.

Figure 8
High-salt diet increases blood pressure in both MORE-PGKO and LC mice. n = 5 per group. Mean systolic (A) and diastolic (B) blood pressure of male LC and MORE-PGKO mice on chow with normal (N) or high (H) dietary salt during light and dark cycles.

Figure 9
MORE-PGKO mice show impaired vasoactivity. Phenylephrine (A) and acetylcholine (B) dose-response curves of aortic rings from MORE-PGKO and LC mice. Results were similar in both genders and were combined (n = 8, 4 males and 4 females, for each genotype). *P < 0.05, **P < 0.01 versus LC, Bonferroni post-test.
GTT and ITT. GTT and ITT were performed in 6-month-old MORE-PGKO and LC mice (n = 8–11 for separate male and female groups of each genotype) as previously described (18, 49–51), with the following modifications. For GTT, mice were fasted overnight for 14 hours, and glucose (2 g/kg body weight) was injected intraperitoneally. Blood glucose was measured by tail bleeding using Elite glucometer (Bayer) at 0, 15, 30, 60, and 120 minutes after glucose injection. Tail blood (50 μl) was collected at 0, 30, and 120 minutes after glucose injection for insulin measurements. For ITT, randomly fed mice at 7–9 hours into the light cycle were injected with insulin (1 IU/kg body weight; Eli Lilly and Co.) intraperitoneally, and blood glucose was measured as for GTT. Tail blood (100 μl) was collected at the time of injection for blood profile analyses.

Blood profile. Plasma insulin and leptin were measured with ELISA kits from Crystal Chem. Plasma adiponectin was measured with ELISA kit from Linco. Plasma NEFAs and triglycerides were assayed using commercial kits (Wako). Plasma aldosterone was measured by a RIA kit from Diagnostic Products Corporation as described previously (32, 33). PRA was measured by indirect RIA kit (DiaSorin) as the generation of Ang I per μl of plasma per hour after addition of excess Agt (ng Ang I/μl/h) (52, 53).

Direct measurement of blood pressure by radiotelemetry. Blood pressure and heart rate of MORE-PGKO and LC mice as well as A-ZIP mice and their WT controls (n = 8, 4 male and 4 female, per group; all 3–4 months old) were measured by radiotelemetry (PA-C20; Data Sciences International) as described previously (54). Briefly, the left common carotid artery was cannulated with the implant catheter, and the implant body was secured in the abdominal cavity. Data were collected for 4 weeks beginning immediately after implantation. Mean systolic blood pressure, diastolic blood pressure, and heart rate were obtained by averaging the data of the final 2 weeks of the salt loading experiment, 3-month-old male MORE-PGKO and LC mice. Actin was used as an endogenous control. Results were similar for female MORE-PGKO and LC mice.

Vasoactivity study. Rings (2 mm long) of thoracic aorta from 3- to 4-month-old MORE-PGKO and LC mice (n = 4 males and 4 females of each genotype) were mounted in a myograph system (Danish MyoTechnology A/S) and contracted with KCl (55). Phenylephrine dose-response curves were then performed, and the results were expressed as contraction percentage of maximum response to KCl. Acetylcholine dose-response curves were performed after the aortic rings were first contracted with phenylephrine, and the results were expressed as percentage of the maximum response to acetylcholine.

Western blot analysis. Total protein was isolated from aortas, subjected to electrophoresis, and transferred to PVDF membranes as described previously (37, 56). Membranes were incubated with primary antibodies recognizing phosphorylated eNOS (Ser1177; Cell Signaling Technology), total eNOS (ABR–Affinity BioReagents), and actin (Santa Cruz Biotechnology Inc.). The membranes were then incubated in secondary antiseraums conjugated with horseradish peroxidase and detected as previously described (56).

Statistics. Mean ± SEM values were analyzed using Prism (version 3; GraphPad Software). Statistical comparisons between groups were performed by Student’s t test unless all values in a group equaled 0, in which case Fisher’s exact test was used. Curves were compared by 2-way ANOVA with appropriate post hoc comparisons. Cumulative phase angles were compared by t test. Curves were compared by 2-way ANOVA with appropriate post hoc comparisons. Curves were compared by 2-way ANOVA with appropriate post hoc comparisons.
ANOVA and Bonferroni post-tests. Groups were considered significantly different if \( P \) values were ≤ 0.05.

Acknowledgments

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Figure 12

Hypertension in A-ZIP lipodystrophic mice. Systolic (A) and diastolic (B) blood pressure and heart rate (C) of WT and A-ZIP mice. A-ZIP mice showed significantly higher systolic and diastolic blood pressure and lower heart rates than did WT mice. Results were similar in both genders and were combined (\( n = 8 \), 4 males and 4 females, for each genotype). Data collected over 2 weeks were averaged and presented as light/dark cycle over a 24-hour period. *\( P < 0.05 \), **\( P < 0.01 \), ***\( P < 0.001 \) versus WT, Bonferroni post-test.