Ankyrin-B metabolic syndrome combines age-dependent adiposity with pancreatic β cell insufficiency

Damaris N. Lorenzo, 1,2 Jane A. Healy, 2 Janell Hostettler, 1,2 Jonathan Davis, 1,2 Jiayu Yang, 2 Chao Wang, 3,4 Hans Ewald Hohmeier, 5,6 Mingjie Zhang, 3,4 and Vann Bennett 1,2

1 Howard Hughes Medical Institute and 2 Department of Biochemistry, Duke University, Durham, North Carolina, USA. 3 Division of Life Science, State Key Laboratory of Molecular Neuroscience, Hong Kong University of Science and Technology, Hong Kong, China. 4 Center of Systems Biology and Human Health, School of Science and Institute for Advanced Study, Hong Kong, China. 5 Department of Medicine and 6 Sarah W. Stedman Nutrition and Metabolism Center, Duke University, Durham, North Carolina, USA.

Rare functional variants of ankyrin-B have been implicated in human disease, including hereditary cardiac arrhythmia and type 2 diabetes (T2D). Here, we developed murine models to evaluate the metabolic consequences of these alterations in vivo. Specifically, we generated knockin mice that express either the human ankyrin-B variant R1788W, which is present in 0.3% of North Americans of mixed European descent and is associated with T2D, or L1622I, which is present in 7.5% of African Americans. Young Ankb R1788W/R1788W mice displayed primary pancreatic β cell insufficiency that was characterized by reduced insulin secretion in response to muscarinic agonists, combined with increased peripheral glucose uptake and concomitantly increased plasma membrane localization of glucose transporter 4 (GLUT4) in skeletal muscle and adipocytes. In contrast, older Ankb L1622I/L1622I and Ankb R1788W/R1788W mice developed increased adiposity, a phenotype that was reproduced in cultured adipocytes, and insulin resistance. GLUT4 trafficking was altered in animals expressing mutant forms of ankyrin-B, and we propose that increased cell surface expression of GLUT4 in skeletal muscle and fatty tissue of Ankb R1788W/R1788W mice leads to the observed age-dependent adiposity. Together, our data suggest that ankyrin-B deficiency results in a metabolic syndrome that combines primary pancreatic β cell insufficiency with peripheral insulin resistance and is directly relevant to the nearly one million North Americans bearing the R1788W ankyrin-B variant.

Introduction

The incidence of obesity, type 2 diabetes (T2D), and related comorbidities has markedly increased during the last 20 years (1, 2). Individual susceptibility to T2D and obesity reflects a complex mix of genetic, epigenetic, and environmental influences that are exacerbated by poor dietary habits and sedentary lifestyles (3). Initial efforts to uncover obesity and T2D susceptibility loci through GWAS have identified common genetic variants with low effect sizes (4, 5), which collectively account for only 10% of disease hereditability (6, 7). Moreover, disease variants associated with particular ethnic groups are often undetected in GWAS because these populations are either not included or are underrepresented (8). Increasing evidence suggests that gene variants with frequencies below the GWAS threshold of detection of 5% are also important contributors to complex disease susceptibility (9, 10). The growing availability of whole-exome and whole-genome sequence information for large numbers of individuals is facilitating the detection of lower-frequency variants that may be associated with common metabolic diseases. However, identifying the variants among these candidates that exhibit a physiological phenotype remains a challenge.

Ankyrin-B (AnkB) is a member of the ankyrin family of membrane adaptors that contributes to the assembly of diverse specialized plasma membrane domains (11). In addition, AnkB associates with phosphatidylinositol 3-phosphate in intracellular membranes and with the dynactin/dynein motor complex to facilitate microtubule-based organelle transport (12–14). In the heart, AnkB regulates Ca2+ dynamics through localization of the sodium pump, the sodium/calcium exchanger, and inositol-1, 4, 5-trisphosphate receptors (InsP3Rs) to transverse tubules of cardiomyocytes (15, 16). Nonsynonymous mutations in ANKB have been linked to a hereditary cardiac arrhythmia syndrome and are present at frequencies from 0.05% to 7% in human populations, depending on ethnicity (refs. 15–18 and Figure 1A).

AnkB is enriched in pancreatic β cells, where it stabilizes InsP3R and regulates intracellular Ca2+ release, functions required for enhanced insulin secretion in response to muscarinic agonists (19). AnkB-haploinsufficient (Ankb−/−) mice exhibited impaired oral glucose tolerance, with reduced cholinergic potentiation of glucose-stimulated insulin secretion. Interestingly, the cardiac arrhythmia–associated human p.R1788W AnkB variant is enriched in probands of mixed European descent and Hispanic probands.
with T2D from the American Diabetes Association GENNID cohort, relative to its rare incidence in control groups as well as in other ethnically matched populations (Figure 1B and refs. 19–21). In comparison to wild-type AnkB, expression of the R1788W AnkB variant failed to rescue insulin secretion deficits in AnkB-deficient pancreatic islets, suggesting that this variant is likely to cause defective glycemic regulation through impaired AnkB function and is a risk factor for T2D and associated disorders (19).

To evaluate the physiological consequences of human AnkB variants, we generated knockin mice bearing the p.R1788W mutation, linked to T2D, and the p.L1622I variant, the most common ANKB mutation in African Americans at over 7%, with a presence in less than 0.1% of individuals of mixed European descent (Figure 1A and refs. 17–20). The p.L1622I AnkB variant, characterized in the setting of hereditary cardiac arrhythmia, has demonstrated functional effects in cardiomyocytes (17, 18). Although this variant did not associate with T2D in the GENNID cohort, within this group it was exclusively identified in patients of African American descent, who are known to have a greater risk of diabetes than counterparts of mixed European descent. We hypothesized that p.L1622I AnkB might promote more subtle metabolic derangements that, in combination with other genetic, environmental, and lifestyle factors, could promote diabetogenesis. Here, we show that mice homozygous for either the R1788W or L1622I alterations develop a metabolic syndrome characterized by early-onset pancreatic β cell dysfunction and age- or diet-dependent increased adiposity and insulin resistance. We explore the mechanistic basis for these phenotypes, determining that these AnkB variants influence insulin secretion in pancreatic β cells acting through InsP3R8s.

Furthermore, we demonstrate for the first time to our knowledge that AnkB promotes internalization of GLUT4 from the plasma membrane and present evidence that increased cell surface GLUT4 in AnkbR1788W/R1788W mice results in age-dependent adiposity and insulin resistance.

Results

Human AnkB variants cause tissue-specific AnkB deficiency. We generated knockin mice bearing either the T2D-associated R1788W variant or the L1622I variant (Supplementary Figure 1A; supplemental material available online with this article; doi:10.1172/JCI81317DS1). R1788 and L1622 residues are located in the unstructured C-terminal regulatory domain of AnkB and are conserved from humans to chickens (Figure 1, C and D, and ref. 22). The R1788W and L1622I variants were predicted to be damaging by computational structural modeling, including the Polyphen-2 algorithm (R1788W score = 1 and L1622I score = 0.975). R1788W was also expected to be damaging by sorting intolerant from tolerant (SIFT) analysis (score = 0.001). We found that functional changes could not be attributed to effects of these human variants on AnkB secondary or tertiary structure, as recombinant AnkB polypeptides containing the R1788W and L1622I substitutions had no effect on AnkB’s secondary structure in assays of circular dichroism spectroscopy and chymotryptic digestion (Supplemental Figure 1, B and C).

L1622I and R1788W AnkB variants are predominantly found in heterozygous carriers in human populations. However, to facilitate the detection of early metabolic deficits, we initially characterized phenotypes of homozygous R1788W (AnkbR1788W/R1788W) and L1622 (AnkbL1622I/L1622I) AnkB knockin mice. Congenic AnkbR1788W/R1788W and L1622 AnkbR1622I/L1622I mice, confirmed by DNA sequence analysis, were born at the expected Mendelian ratios and exhibited no differences in Ankb mRNA levels, as assessed by quantitative PCR (qPCR) analysis (Figure 1, E and G). To investigate the functional impact of these AnkB variants in vivo, we first evaluated their effect on protein levels. Surprisingly, given the evidence for normal protein folding and unaltered mRNA levels, mice homozygous for AnkB variants R1788W and L1622I had significant reductions in AnkB protein levels in a subset of tissues, including fat, liver, skeletal muscle, pancreatic β cells, and heart. By comparison, no change in AnkB protein expression was detected in brain (Figure 1F and Supplemental Figure 1D). In addition, primary mouse embryonic fibroblast (MEF) cultures derived from AnkbR1788W/R1788W and AnkbL1622I/L1622I mice showed reduced AnkB protein expression (Figure 1F and Supplemental Figure 1D).

Analysis of protein turnover in cycloheximide-treated MEFs revealed that, in control cells, AnkB expression was reduced to 55% 4 hours after blocking protein synthesis (Supplemental Figure 2, A and B). In contrast, AnkB levels were already reduced to 36% and 47% in untreated AnkbR1788W/R1788W and AnkbL1622I/L1622I MEFs, respectively, and were unaffected by cycloheximide treatment (Supplemental Figure 2, A and B). These data suggested that these AnkB variants specifically affect the stability of a rapidly degradable AnkB subpopulation. To determine what cellular pathways accounted for the observed decrease in AnkB expression in AnkbR1622I/L1622I and AnkbR1788W/R1788W mice, we treated MEFs with inhibitors of various protein degradation systems. Our results indicate that calpain, lysosomal, and proteasomal pathways cleared the mutant AnkB polypeptides at higher rates than the control AnkB protein and were partially responsible for their lower levels (Supplemental Figure 2C).

R1788W AnkB mutation impairs insulin secretion and oral glucose tolerance. Deficits in AnkB and InsP3R levels noted in Ankb−/− mice (19) were recapitulated in islets of 3-month-old AnkbR1788W/R1788W and AnkbL1622I/L1622I mice (Figure 1F, Figure 2A, and Supplemental Figure 1D). Similar to those of Ankb−/− mice, isolated pancreatic islets of AnkbR1788W/R1788W mice exhibited over 50% reduction in glucose-stimulated insulin secretion normalized to total insulin content. By contrast, islets isolated from AnkbL1622I/L1622I mice responded normally to glucose stimulation (Figure 2B).

While fasting serum glucose levels were unaltered in both AnkbR1788W/R1788W and AnkbL1622I/L1622I mice, oral administration of glucose (2 g/kg body weight) during an oral glucose tolerance test led to both an increased level of blood glucose at 30 minutes as well as a delay in glucose clearance (Figure 2D and Supplemental Figure 3A). The areas under the curve for the blood glucose response were significantly increased by 2- and 1.7-fold in AnkbR1788W/R1788W and AnkbL1622I/L1622I knockin mice, respectively (Figure 2E), and by 1.8- and 1.4-fold in heterozygous carriers of the same genotypes, designated as AnkbR1788W/+ and AnkbL1622I/+ mice (Supplemental Figure 3B), when compared with control littermates. Consistent with impaired glucose clearance, AnkbR1788W/R1788W mice exhibited 50% reduction in serum insulin levels following glucose ingestion (Figure 2C). In contrast, both AnkbR1788W/R1788W and AnkbL1622I/L1622I mice displayed normal tolerance to glucose administered through intra-
Figure 1. Human AnkB variants cause tissue-specific AnkB reduction. (A) Variant frequency (per 100 individuals) for the most common ANKB variants in North Americans of European and African descent. Allele frequencies were compiled from the NHLBI GO Exome Sequencing Project. Asterisks indicate undetected variants. (B) Frequency of the p.R1788W ANKB variant in European Americans and Hispanics from the GENNID cohort of noninsulin-dependent diabetic patients (control and T2D cases) and from individuals of the same ethnicity sequenced as part of the entire GENNID cohort, the NHLBI GO Exome Sequencing Project, and the 1000 Genomes Project. Asterisks indicate undetected variants. (C) Model of AnkB structure, with domains indicated, showing the localization of L1622 and R1788 sites within AnkB’s unstructured C-terminal domain. (D) Evolutionary conservation of L1622 and R1788 sites. (E) Sequencing chromatograms of brain cDNA from AnkB+/+ and AnkB knockin mice. (F) Quantitative analysis of AnkB protein levels for the indicated tissues as a percentage of control (AnkB+/+). (G) Quantification of 220-kDa AnkB transcript levels by qPCR. GAPDH was used for normalization of mRNA and protein levels. Data represent mean ± SEM (n = 5 males, 10-weeks-old). Results are representative of 3 independent experiments. ***P < 0.001, 2-tailed t test.
peritoneal injection relative to that of Ankb+/+ controls (Supplemental Figure 4). The finding of normal intraperitoneal glucose tolerance but impaired oral glucose tolerance was also reported for Ankb+/+ mice and was rationalized to be due to loss of cholinergic enhancement of insulin secretion, which occurs following oral intake of glucose (19).

In pancreatic β cells, InsP3R regulates Ca2+ mobilization and cholinergic potentiation of glucose-stimulated insulin secretion. InsP3R deficiency results in hyperglycemia and glucose intolerance in young mice (23–25), without affecting insulin sensitivity (Supplemental Figure 5). Interestingly, 3-month-old InsP3R-haploinsufficient (Itp1+/−) mice exhibited abnormal oral glucose tolerance and insulin secretion deficits following oral glucose ingestion, which were similar to the phenotypes observed in young AnkbR1788W/R1788W mice (Figure 2, F–H, and ref. 25). However, AnkbL1622I/L1622I mice, which also exhibit a reduction in InsP3R, had normal glucose-dependent insulin secretion from isolated islets and normal serum insulin levels following glucose ingestion (Figure 2, B and C). Thus, InsP3R downregulation likely contributes to primary β cell insufficiency in young R1788W mice, while L1622I mice apparently employ additional compensatory mechanism(s).

Increased peripheral glucose uptake in young AnkbR1788W/R1788W mice. Despite impaired insulin secretion and abnormal oral glucose tolerance, 3-month-old AnkbR1788W/R1788W mice exhibited normal fasting glucose levels (Figure 2E, time 0, and Figure 3A, time 0). To evaluate whole-body glucose utilization, hyperinsulinemic-euglycemic clamp, combined with measurements of 2-deoxy-[3H]-glucose (2-DG) transport in vivo in skeletal muscle and white adipose tissue (WAT), was conducted at the Yale Metabolic Core Facility. Blood glucose levels were tightly maintained through a variable glucose infusion rate (GIR) in response to a constant exogenous supply of insulin. Surprisingly, clamp measurements revealed that 3-month-old AnkbR1788W/R1788W mice have increased glucose utilization compared with that of control littermates and Ankb+/− mice, as demonstrated by a significantly higher whole-body GIR (Figure 3, A and B). Moreover, 2-DG uptake by AnkbR1788W/R1788W mice was significantly elevated compared with that of controls in both skeletal muscles (30%) and WAT (76%) (Figure 3, C and D). By comparison, Ankb+/− mice exhibited normal 2-DG uptake in skeletal muscle and a more modest 43% increase in WAT (Figure 3, C and D). In contrast, both AnkbR1788W/R1788W and AnkbL1622I/L1622I 3-month-old mice showed similar increases in hepatic insulin responsiveness, as reflected by a 93% suppression of hepatic endogenous glucose production (EGP) during the clamp in AnkbR1788W/R1788W mice and a 88% suppression in AnkbR1622I/L1622I mice, compared with a 53% suppression in control mice (Supplemental Figure 6, A and B). Changes in EGP were concomitant with reductions in transcript levels for gluconeogenesis rate-limiting enzymes (Supplemental Figure 6E). Together, our results suggest that reduced gluconeogenesis rates contribute to the maintenance of normoglycemia in young AnkbR1788W/R1788W and Ankb+/− mice. It is notable that suppression of EGP was seen in both AnkbR1788W/R1788W and AnkbR1622I/L1622I animals, though higher whole-body glucose disposal was observed exclusively in AnkbR1788W/R1788W mice. This suggests that increased peripheral glucose uptake, rather than decreased gluconeogenesis, is the primary cause of the accelerated glucose disposal rate detected in the AnkbR1788W/R1788W mice.

Blood glucose uptake by skeletal muscle and adipocytes is achieved by glucose transporter 4 (GLUT4), which rapidly translocates to the cell surface from intracellular storage sites in response to insulin (26). Numerous studies suggest that activation of the serine/threonine phosphoinositide 3-kinase and phosphorylation of its downstream target PKB (also known as AKT) are key steps in the regulation of insulin-induced exocytosis of GLUT4-containing vesicles (27–29). Significantly, levels of phosphorylated S473 AKT were similar in skeletal muscle or in differentiated adipocytes across genotypes, both under basal and insulin-stimulated conditions (Figure 3, E and F). These observations suggest that the increased glucose uptake in peripheral tissues of AnkbR1788W/R1788W mice is not likely due to global alterations in AKT-dependent activation of insulin signaling. However, we cannot rule out effects of Ankb variants on specific AKT isoforms or other components of the AKT insulin-dependent signaling pathway.

Increased cell-surface GLUT4 in tissues of Ankb knockin mice. To investigate the mechanism underlying the increased glucose uptake observed in young AnkbR1788W/R1788W mice, we examined the levels of GLUT4 in plasma membrane fractions from extracts of skeletal muscle and adipose tissue either under resting conditions or following exogenous insulin stimulation. Untreated control (Ankb+/−) mice showed low levels of plasma membrane-associated GLUT4, which increased around 2-fold upon insulin stimulation (Figure 3, G and I). In contrast, under basal conditions, GLUT4 was robustly associated with plasma membranes from AnkbR1788W/R1788W mice (over 2-fold higher GLUT4 cell surface levels than in controls) and, to a lesser extent, from Ankb+/− mice. GLUT4 maintained its plasma membrane localization after insulin treatment in both homoygous knockin strains (Figure 3, G and H). Total GLUT4 levels were similar in skeletal muscle and fat across all genotypes (Figure 3I). The elevations in surface-associated GLUT4 under basal conditions were also detected in skeletal muscle and adipose tissue membrane fractions from Ankb+/− mice (Figure 3, G and H), suggesting that these deficits are, at least in part, due to reduced levels of Ankb polypeptides. The persistent plasma membrane localization of GLUT4 in skeletal muscle and adipose tissue in young AnkbR1788W/R1788W mice provides a potential compensatory mechanism to explain how these animals maintain normal fasting glucose levels in the face of impaired insulin secretion.

Ankb regulates GLUT4 internalization in differentiated adipocytes. To elucidate the role of Ankb in GLUT4 cellular localization, we first assessed glucose uptake in differentiated adipocyte cultures established from primary MEF lines derived from Ankb+/−, AnkbR1788W/R1788W, and Ankb+/− mice. Similar to the in vivo results, differentiated adipocytes from Ankb+/−, AnkbR1622I/L1622I, and AnkbR1788W/R1788W animals exhibited elevated glucose uptake under both resting conditions and following insulin stimulation (Figure 4A). Next, we assessed the cell surface expression of GLUT4 in adipocytes by transfection with a construct encoding GLUT4-GFP containing an exonaficial Myc epitope (Figure 4C). The expressed fusion protein has been shown previously to be fully active in vivo (30). At steady-state, the majority of GLUT4 molecules were intracellular in Ankb+/− adipocytes, as measured by the ratio of plasma membrane to cytosolic myc-GLUT4 signal (0.2, Figure 4, D and E, basal). However, cell surface GLUT4 localization increased 3.8-fold following insulin
in AnkbL1622I/L1622I adipocytes (Figure 4D, time 0), we hypothesized that elevation in GLUT4 on the surface of cells isolated from Ankb knockin mice resulted from a slower internalization rate. Monitoring of GLUT4 endocytosis in differentiated control adipocytes after insulin stimulation showed markedly increased plasma membrane localization (Figure 4D, time 0), which decreased to nonstimulated values (Figure 4, D and E) 30 minutes after internalization ($t_{1/2}$ of GLUT4 internalization = 7 minutes). In contrast, plasma membrane–associated Myc-GLUT4-GFP treatment (Figure 4, D and E, time 0). In contrast, cell surface GLUT4 was already elevated about 2-fold in AnkbL1622I/L1622I and 3.2-fold in AnkbR1788W/R1788W adipocytes at rest and remained high in the insulin-stimulated state (Figure 4, D and E).

In muscle cells and adipocytes under basal conditions, low levels of GLUT4 on the cell surface are maintained by a faster rate of GLUT4 endocytosis relative to re-exocytosis (31, 32). Considering that we did not observe significant increases in the rate of GLUT4 translocation to the surface in response to insulin in AnkbR1788W/R1788W or AnkbL1622I/L1622I adipocytes compared with that in AnkbL1622I/L1622I adipocytes (Figure 4D, time 0), we hypothesized that elevation in GLUT4 on the surface of cells isolated from Ankb knockin mice resulted from a slower internalization rate. Monitoring of GLUT4 endocytosis in differentiated control adipocytes after insulin stimulation showed markedly increased plasma membrane localization (Figure 4D, time 0) increased plasma membrane localization (Figure 4D, time 0), which decreased to nonstimulated values (Figure 4, D and E) 30 minutes after internalization ($t_{1/2}$ of GLUT4 internalization = 7 minutes). In contrast, plasma membrane–associated Myc-GLUT4-GFP
in Ankβ<sup>R1788W/R1788W</sup> adipocytes doubled compared with that in Ankβ<sup>+/+</sup> adipocytes at the same time interval and exhibited a substantial reduction in internalization rate (t<sub>1/2</sub> = 26 minutes) (Figure 4, D and E). Adipocytes from Ankβ<sup>L1622I/L1622I</sup> mice showed an intermediate phenotype (t<sub>1/2</sub> = 13 minutes) (Figure 4, D and E). Expression of wild-type 220-kDa AnkB-HA in Ankβ<sup>+/+</sup> adipocytes rescued the deficits in the cellular distribution and internalization of GLUT4 (t<sub>1/2</sub> = 8 minutes) (Figure 4, D and E).

AnK recruits binding partners to specialized membrane domains and associates with PtdIns(3)P-enriched membranes, such as endosomes, to promote organelle transport (11, 14). Interestingly, Ankβ directly interacts with members of the Eps15 homology domain/receptor-mediated endocytosis-1 (EHD/ RME-1) family of endosome-based scaffolding molecules and with clathrin heavy chain, both of which have well-established roles in the regulation of GLUT4 intracellular traffic (33–37). Therefore, we hypothesized that Ankβ, either directly or in combination with its binding partners, might facilitate GLUT4 retrieval from the plasma membrane. To explore this possibility, we first determined whether Ankβ and GLUT4 proteins associate in vivo. Coimmunoprecipitation experiments with skeletal muscle homogenates showed that these proteins interact (Figure 4B). Similarly, coimmunoprecipitation of lysates of HEK293 cells expressing 220-kDa Ankβ-GFP and HA-GLUT4 showed binding of Ankβ to GLUT4. Anti-GFP antibody coimmunoprecipitated wild-type HA-GLUT4 with Ankβ-GFP (Figure 4F). However, it failed to pull down HA-tagged GLUT4 harboring the A<sup>5</sup>AAA<sup>8</sup> substitutions in the N-terminal F<sup>5</sup>QQI<sup>8</sup> motif (Figures 4F and 7A). However, we found that the ANK repeat domain of Ankβ failed to associate with the N-terminal portion of GLUT4. In contrast, this domain formed a high-affinity interaction with residues 1206–1233 of the mouse L1 cell adhesion molecule (L1CAM), a known ankyrin-binding partner (Supplemental Figure 7, A and B, and ref. 41). Thus, Ankβ interaction with GLUT4 is not mediated by direct interaction of the N-terminal GLUT4 residues with the ANK repeat domain. This suggests that the observed coimmunoprecipitation of GLUT4 and Ankβ may require the intact GLUT4 polypeptide and/or involve accessory protein(s).

A single substitution of the phenylalanine residue within the F<sup>5</sup>QQI<sup>8</sup> motif (F<sup>5</sup>AA) causes loss of GLUT4 localization to cell surface clathrin lattices, reduction in GLUT4 internalization, and accumulation of GLUT4 on the plasma membrane of nonstimulated adipocytes (40). To further assess whether Ankβ facilitates GLUT4 internalization by mediating its interaction with the endocytic components through the FQQI motif, we evaluated the cellular distribution of the F<sup>5</sup>AA Myc-GLUT4-GFP mutant protein in Ankβ<sup>+/+</sup> and Ankβ<sup>R1788W/R1788W</sup> adipocytes. We found that GLUT4 was internalized at similar rates in Ankβ<sup>R1788W/R1788W</sup> adipocytes expressing wild-type Myc-GLUT4-GFP (t<sub>1/2</sub> = 25 minutes) or mutant F5A Myc-GLUT4-GFP (t<sub>1/2</sub> = 24 minutes) proteins (Figure 4, D and E) as well as in Ankβ<sup>+/+</sup> cells expressing the internalization-deficient F<sup>5</sup>AA GLUT4 mutant (t<sub>1/2</sub> = 25 minutes) (Figure 4E). These results indicate that Ankβ deficiency does not affect the internalization deficits of mutant F<sup>5</sup>AA GLUT4.

Moreover, these findings suggest that Ankβ facilitates the association of GLUT4 with endosomes through the FQQI motif, either directly through domains other than the ANK repeats or through intermediate proteins. This suggests that Ankβ might function as a heretofore undescribed adaptor for the endocytic machinery that promotes retrieval of the GLUT4 transporter from the cell surface.

**Age-dependent increase in adiposity in R1788W Ankβ mice.** Increased GLUT4 expression in adipose tissue leads to a constitutive elevation in glucose uptake with increased adiposity in mice (42, 43). Similarly, significantly elevated basal glucose uptake and GLUT4 levels occur in adipocytes from young obese Zucker rats when compared with lean littermates (44). We therefore evaluated effects of age on the extent and distribution of adipose tissue in Ankβ<sup>+/+</sup>, Ankβ<sup>L1622I/L1622I</sup>, and Ankβ<sup>R1788W/R1788W</sup> mice. Young (3-month-old) Ankβ knockin mice had a percentage of body fat indistinguishable from that of controls (Figure 5B and Figure 6D). However, older (10-month-old) Ankβ<sup>R1788W/R1788W</sup> mice exhibited increased body fat mass, as measured by nuclear magnetic resonance (Figure 5B and Figure 6D). In addition, both Ankβ<sup>R1788W/R1788W</sup> and Ankβ<sup>L1622I/L1622I</sup> mutant mice exhibited significant adipocyte hypertrophy (54% increase in adipocyte diameter in Ankβ<sup>R1788W/R1788W</sup> and 38% in Ankβ<sup>L1622I/L1622I</sup> mice; Figure 5, A and C), which was accompanied by elevated levels of circulating nonesterified fatty acids (Figure 5D). Interestingly, these increases in body fat were more prominent than the overall changes in total body weight (Figure 6, A–C). These findings, together with the similar percentages of lean mass observed among all genotypes tested (Figure 6E), suggest that expression of these Ankβ variants is associated with a redistribution of body mass, favoring the accumulation of adipose tissue. Thus, our results indicate that, while increased glucose clearance may initially compensate for hyperglycemia caused by impaired insulin production, the persistent rise in glucose uptake by insulin target organs triggers age-dependent increases in adiposity in older animals.

Adult-onset obesity is commonly caused by dysfunction of the hypothalamic circuits regulating energy homeostasis (45, 46). Surprisingly, systematic metabolic cage analysis revealed that both 3-month-old and 10-month-old Ankβ<sup>R1788W/R1788W</sup> and Ankβ<sup>L1622I/L1622I</sup> mice showed overall similar average rates of food intake, energy source utilization, and activity compared with littermate controls during both light and dark cycles (Figure 6, F–M). These results suggest that the increased body weight and adiposity observed in older Ankβ<sup>R1788W/R1788W</sup> and Ankβ<sup>L1622I/L1622I</sup> mice are not due to marked changes in appetite or activity. However, our data do not exclude small cumulative effects of decreased activity and/or increased food consumption that were below the sensitivity of these assays or other changes in water content, lean mass, and food absorption in the gut.

**Cell-autonomous increases in lipid accumulation in Ankβ mutant adipocytes.** We next asked whether increased adiposity in vivo could be recapitulated in a cell-autonomous manner in adipocytes differ-
Figure 3. Increased peripheral glucose utilization associated with persistent cell surface GLUT4 in Ankbr<sup>1788W/R1788W</sup> mice. (A) Hyperinsulinemic-euglycemic clamp analysis in 3-month-old (n = 8) mice. (B) GIR during insulin clamp. (C and D) 2-DG uptake by (C) skeletal muscle and (D) WAT. (E and F) Immunoblots of total AKT and phosphorylated AKT (p-AKT S473) levels in (E) skeletal muscle and (F) differentiated adipocytes from 3-month-old mice under basal or insulin-stimulated conditions. (G and H) Immunoblots and quantification of plasma membrane–associated GLUT4 levels after subcellular fractionation of (G) skeletal muscle and (H) WAT lysates under basal conditions or following insulin stimulation. (I) Total levels of GLUT4, as evaluated by immunoblotting. Data represent mean ± SEM (n = 8 mice). *P < 0.05, **P < 0.01, ***P < 0.001, 1-way ANOVA with post-hoc Tukey test. Results are representative of 3 independent experiments.
The differences in adipocyte differentiation in vitro correlated with an earlier and more robust transcriptional activation of genes implicated in adipogenesis (Supplemental Figure 8, B–E). However, similar changes in the expression profile of adipogenesis-related transcription factors and their targets were not observed before the onset of adiposity in adipose tissue of 3-month-old *AnkBr1788W/R1788W* and *AnkB L1622I/L1622I* mice (Supplemental Figure 8A). Importantly, these in vitro results suggest that AnkB deficiency can promote adiposity.

Figure 4. Increased GLUT4 association with plasma membranes of mutant AnkB adipocytes caused by reduced GLUT4 internalization rates. (A) 2-DC uptake in differentiated adipocytes before and after stimulation with 100 nM insulin. Data represent mean ± SEM of values from one 6-well plate per genotype for 1 of 3 repeats. (B) Coimmunoprecipitation from control mouse skeletal muscle homogenates. (C) Diagram of a GLUT4 construct with extracellular Myc and C-terminal GFP epitopes (Myc-GLUT4-GFP) used for assessing GLUT4 distribution in differentiated adipocytes. The position of the FQQI motif is indicated. (D) GLUT4 localization in differentiated adipocytes under basal conditions or following insulin stimulation and internalization at 37°C. Scale bar: 10 μm. (E) Quantification of GLUT4 association with plasma membranes of differentiated adipocytes either at rest or at indicated times following treatment with 100 nM insulin and internalization at 37°C. (F) Coimmunoprecipitation from HEK293 cells expressing either wild-type HA-GLUT4 or FQQI-A5AAA HA-GLUT4 and GFP-AnkB proteins. Data is representative of 3 independent experiments. Data represent mean ± SEM for 1 of 3 independent determinations (n = 30 cells per genotype and per condition). *P < 0.05, **P < 0.01, ***P < 0.001, 1-way ANOVA with post-hoc Tukey test.
hepatic steatosis (based on liver appearance) phenotypes normally induced by the HFD were more severe in both Ankb R1788W/R1788W and AnkbL1622I/L1622I mice than in control animals, even though the majority of these parameters were similar among young animals on a normal diet across genotypes (Figure 7, A–C, F, and G). HFD-fed homozygous knockin mice also showed higher fasting glucose levels and significant oral glucose intolerance (Figure 7, D and E). It is notable that GLUT4 overexpression in adipocytes did not protect HFD-fed AnkbL1622I/L1622I or AnkbR1788W/R1788W mice from glucose intolerance (47).

Figure 5. Age-dependent increases in adiposity in Ankb mutant mice and elevated lipid accumulation in differentiated adipocytes. (A) Epididymal WAT sections from 3-month-old and 10-month-old mice stained for the lipid-associated protein perilipin. Scale bar: 20 μm. (B) Percentage of body fat mass. (C) Epididymal white adipose size (n = 6 mice). (D) Nonesterified free fatty acid (NEFA) levels (n = 10 mice). (E) Day 8 differentiated adipocytes stained for perilipin. Scale bar: 5 μm. (F) Lipid droplet volume in differentiated adipocytes (n = 100 cells for 1 of 3 independent experiments). Data represent mean ± SEM. *P < 0.05, **P < 0.01, ***P < 0.001, 1-way ANOVA with post-hoc Tukey test.
The R1788W AnkB mutation promotes age-dependent insulin resistance. Obesity and hyperlipidemia are associated with the development of insulin resistance and with hyperglycemia in humans and other species. Therefore, using aged AnkB<sub>R1788W/R1788W</sub> or AnkB<sub>L1622I/L1622I</sub> mice maintained on a regular diet, we next evaluated whole-body glucose responses, which demonstrated the characteristic increased adiposity and elevated levels of free fatty acids (Figure 5, A-D). Not surprisingly, 10-month-old AnkB<sub>R1788W/R1788W</sub> mice exhibited elevated fasting glucose levels and impaired oral glucose tolerance (Figure 8, A and B). In addition, hyperinsulinemic-euglycemic clamp measurements revealed that AnkB<sub>R1788W/R1788W</sub> mice had become less responsive to insulin with age (Figure 8C), as evidenced by a reduction in GIR (29%), a significantly lower 2-deoxy-glucose uptake by muscle (21%) and WAT (37%) (Figure 8, D–F), and a marked loss (91%) of the ability of the liver to suppress EGP (Supplemental Figure 5, C and D). Importantly, insulin resistance was accompanied by decreased activation of insulin signaling in both WAT and skeletal muscle (Figure 8, G and H). Thus, our results indicate that age-dependent increases in adiposity combined with primary pancreatic β cell insufficiency trigger the onset of severe metabolic abnormalities in older AnkB<sub>R1788W/R1788W</sub> animals.

AnkB<sub>R1788W/+</sub> mice have shortened life spans and demonstrate obesity and hyperglycemia with age. Since the L1622I and R1788W AnkB variants are predominantly found in heterozygous carriers in human populations, we assessed the long-term effects of these variants in heterozygous mice. Overall, body weight and fasting glucose levels were increased with age in both control and AnkB<sub>R1788W/+</sub> and AnkB<sub>L1622I</sub> knockin mice, but these changes were more pronounced in AnkB<sub>R1788W/+</sub> mice (Supplemental Figure 10, A and B), which also exhibited shorter life spans. In particular, AnkB<sub>R1788W/+</sub> animals had a median survival of 794 days compared with 896 days for AnkB<sup>+/+</sup> mice and 855 days for AnkB<sub>L1622I</sub> mice (P = 0.005, Supplemental Figure 10, C and D). It is likely that metabolic deficits described here, combined with the more deleterious impact of the R1788W variant on heart function noted in humans (17), contributed to the significantly reduced life span of AnkB<sub>R1788W/+</sub> mice.

Discussion

A major challenge in understanding the recent epidemic in metabolic disorders in modern Western societies is the identification of susceptibility loci that are predominantly expressed under conditions of high-caloric diets and extended life span. We report a previously uncharacterized metabolic syndrome due to an R1788W AnkB mutation shared by nearly one million North Americans, which in mice combines pancreatic β cell dysfunction, characterized by impaired insulin secretion due to InsP3R deficiency, and age-dependent adiposity and insulin resistance, which is associated with increased cell surface GLUT4 and with sustained elevations in glucose uptake in skeletal muscle and fat. In young AnkB knockin mice bearing the T2D-associated R1788W variant, insulin secretion deficits (Figure 2, B and C) were compensated by increased peripheral glucose uptake through persistent cell surface GLUT4 (Figure 3, C, D, G, and D). AnkB<sub>R1788W/R1788W</sub> mice developed metabolic defects, including increased adiposity, elevated fasting glucose, and insulin resistance with age or following a HFD (Figures 5–8). Extrapolating to humans, the metabolic costs associated with the R1788W AnkB mutation would likely have been blunted in earlier times before modern increases in longevity and caloric intake.

An unexpected finding was that both R1788W and L1622I AnkB mutations resulted in a tissue-specific reduction in AnkB polypeptide levels (Figure 1). Interestingly, AnkB<sup>+/−</sup> haplosufficient mice also showed deficits in oral glucose tolerance and insulin secretion (19) and significant increases in cell surface GLUT4, in glucose uptake in adipocytes, and in lipohypertrophy (Figures 3–5). These findings demonstrate that the pathological effects of these AnkB variants are caused, at least in part, by AnkB haplosufficiency. However, the more pronounced metabolic phenotypes observed in AnkB<sub>R1788W/R1788W</sub> and AnkB<sub>R1788W/+</sub> mice and in cultured cells suggest additional functional effects of the R1788W variant. Interestingly, the R1788W mutation is located on the surface of a predicted amphipathic α-helix in the AnkB’s C-terminal regulatory domain and modulates the affinity of AnkB for binding partners, such as the sarcoplasmin protein obscurin (48) and heat shock protein 40 (Hsp40) (49). In contrast, the molecular consequences of the L1622I mutation are not known. Thus, while R1788W and L1622I AnkB variants are both tissue-specific hypomorphs, the R1788W mutation, in addition, results in altered intrinsic function.

AnkB<sub>L1622I/L1622I</sub> mice exhibited a milder pancreatic phenotype than R1788W mice but still shared an increased sensitivity to the metabolic consequences of a HFD (Figure 7). These results suggest the possibility that other mutations of AnkB identified as being functionally significant based on their association with cardiac arrhythmia (15–18) may also cause AnkB deficiency and increased susceptibility to pathological consequences of Western diets. These AnkB mutations occur with a cumulative prevalence of about 2% in individuals of mixed European descent and of over 7% in African Americans (ref. 16 and Figure 1A), which together represent a population large enough to contribute to public health.

An unusual feature of AnkB metabolic syndrome due to the R1788W mutation is the combination of early pancreatic β cell insufficiency with increased glucose uptake by peripheral tissues, leading to age-dependent adiposity and insulin resistance. Patients with this condition would likely present as middle-aged adults with features of both type 1 diabetes and T2D. Interestingly, a subset of diabetes referred to as “double diabetes” also presents with initial loss of pancreatic islet function in combination with insulin resistance (50, 51). However, while most of these patients have autoimmune antibodies, those with AnkB metabolic syndrome would probably experience impaired insulin secretion without involving the immune system.

Despite extensive evidence for insulin-dependent delivery of GLUT4 transporters to the plasma membrane, mechanisms for internalization of GLUT4 following cessation of insulin signaling are less well established (31, 32, 52). Multiple reports provide evidence for both clathrin-mediated endocytosis–dependent (CME-dependent) and CME-independent mechanisms of GLUT4 internalization in adipocytes and muscle cells. The degree of activation of one pathway versus the other and the type of CME-independent route used depends on the cell type...
Figure 6. Increased adiposity in AnkbR1788W/R1788W mutant mice is not caused by major changes in appetite or activity. (A and B) Representative images of congenic Ankb+/+, AnkbL1622I/L1622I, and AnkbR1788W/R1788W (A) 3-month-old and (B) 10-month-old male mice. (C) Body weight. (D) Fat mass and (E) lean mass as a percentage of body weight. (F) Average 24-hour energy expenditure. (G) Oxygen consumption. (H) CO2 production measured by CLAMS. (I) Respiratory quotient. (J and L) Activity during a 24-hour period for (J) 3-month-old and (L) 10-month-old mice. (K and M) Food consumption on a regular diet during a 24-hour period for (K) 3-month-old and (M) 10-month-old mice. Data represent mean ± SEM, (n = 8, 3-month-old mice and n = 10, 10-month-old mice) for 1 experiment. *P < 0.05, **P < 0.01, ***P < 0.001, 1-way ANOVA with post-hoc Tukey test.
and, importantly, on the insulin stimulation stage of the cell (52–59). In addition, several studies suggest that the interaction between the p2-adaptin subunit of the clathrin adaptor AP-2 and the phenylalanine (F) residue within the GLUT4 N-terminal FQQL motif is the predominant signal that sorts GLUT4 to clathrin-coated pits (60, 61). However, adaptors other than AP-2 may mediate GLUT4 association with the endocytic machinery since silencing of AP-2 in adipocytes reduced GLUT4 internalization only in insulin-stimulated cells, even though GLUT4 colocalizes with clathrin puncta and is still internalized through CME in the basal state (37, 54).

The following findings suggest that AnkB functions as a heretofore undescribed adaptor that either by direct interaction with the GLUT4 transporter or through other proteins promotes GLUT4 retrieval from the cell surface. First, AnkB coimmunoprecipitated with GLUT4 from skeletal muscle lysates and, importantly, on the insulin stimulation stage of the cell (52–59). In addition, several studies suggest that the interaction between the μ2-adaptin subunit of the clathrin adaptor AP-2 and the phenylalanine (F) residue within the GLUT4 N-terminal FQQL motif is the predominant signal that sorts GLUT4 to clathrin-coated pits (60, 61). However, adaptors other than AP-2 may mediate GLUT4 association with the endocytic machinery since silencing of AP-2 in adipocytes reduced GLUT4 internalization only in insulin-stimulated cells, even though GLUT4 colocalizes with clathrin puncta and is still internalized through CME in the basal state (37, 54).

The following findings suggest that AnkB functions as a heretofore undescribed adaptor that either by direct interaction with the GLUT4 transporter or through other proteins promotes GLUT4 retrieval from the cell surface. First, AnkB coimmunoprecipitated with GLUT4 from skeletal muscle lysates and
required the FQQI motif of GLUT4 for coimmunoprecipitation (Figure 4F). Moreover, the internalization-deficient F5A Myc-GLUT4-GFP mutant was resistant to the effects of AnkB deficiency in R1788W adipocytes (Figure 4, C–E), suggesting that AnkB regulates GLUT4 internalization through a mechanism that requires the interaction of the GLUT4 FQQI motif with the endocytic machinery.

R1788W Ankβ knockin mice developed age-dependent increases in adiposity without marked changes in food consumption or activity. Interestingly, AnkβR1788W/R1788W as well as AnkβL1622I/L1622I differentiated adipocytes showed cell-autonomous increases in adipocyte number and enlarged lipid droplets. Our observations suggest a mechanism for age-dependent adiposity that is independent of major changes in appetite or activity and is rather caused by elevation in cell surface-associated GLUT4. It is likely that persistent glucose influx, which supplies adipocytes with lipid synthesis precursors, contributes to increased adipocyte number and size.
A major challenge in the management of T2D is the phenotypic heterogeneity of the disease, which results in differences in patient responses to treatments. Identifying and characterizing distinct subcategories of T2D with unique biology, like the one described in this study, may facilitate early disease prognosis and guide the choice of treatment. In this case, patients with ANKB mutations could become prime candidates for early intervention through personalized therapies. Based on our results in knockin mice, we predict that, in the case of R1788W carriers, in addition to following a healthy lifestyle, it would be beneficial for them to manage their metabolic syndrome at early stages, using a combination strategy that promotes noncholinergic stimulation of insulin secretion by agents such as incretins (64) as well as agents such as α-glucosidase inhibitors and diets that reduce hyperglycemia.

In summary, we identify a metabolic syndrome due to the human AnkB R1788W variants present in 0.3% of individuals of mixed European descent that combines early-onset abnormal pancreatic β cell function with age- and diet-dependent increased adiposity. Intriguingly, additional AnkB variants, some of which are known to directly affect cardiac function (15–18), might also affect glucose homeostasis and related disorders with distinct presentations in different ethnic populations. In particular, the L1622I allele shown in this study to increase susceptibility to a HFD is shared by 7% of African Americans, or about the same prevalence as the trait for sickle cell anemia, and would be predicted to be present in homozygotes at a frequency of about 0.1%. Given the differences between mice and humans, it will be critical to evaluate metabolic effects of ANKB variation in human subjects, including those homozygous for the L1622I mutation.

Methods

Mouse lines and animal care. All studies were conducted in 3-month-old or 10-month-old congenic male mice. Ankb+/− mice have been previously described (18). Itpr1−/− mice were purchased from The Jackson Laboratory (stock 000019). Ankb knockin mice bearing the L1622I mutation were generated in the laboratory of Dr. Christian Franke (University of Massachusetts). All studies were conducted in 3-month-old mice and animal care was carried out according to the guide for the Care and Use of Laboratory Animals (National Institutes of Health). All studies were approved by the Institutional Animal Care and Use Committee of the Yale University School of Medicine.

Experiments were performed on 3-month-old male mice. Ankb+/− and wild-type (129/SvJ background) control mice were divided into age- and diet-matched groups and fed ad libitum either control diet or HFD for a period of 12 weeks. The body composition of 12-month-old mice and animal care was performed according to the guide for the Care and Use of Laboratory Animals (National Institutes of Health).

Tolerance tests and measurement of plasma metabolites. Mice were fasted for 14 hours, and blood glucose was measured by tail bleeding at 0, 15, 30, 60, 90, and 120 minutes after glucose administration by oral gavage. For insulin tolerance tests, mice were administered 0.75 U/kg body weight of recombinant human insulin (Humulin R, Eli Lilly) by intraperitoneal injection.

Metabolic cage analysis and hyperinsulinemic-euglycemic clamp study. Metabolic cage analysis and clamp studies were conducted at the Yale Mouse Metabolic Phenotyping Center (MMPC) following recommendations of the MMPC Consortium (65). Body composition was determined by 1H magnetic resonance spectroscopy (Bruker Minispec). The comprehensive lab animal monitoring system (CLAMS, Columbus Instruments) was used to evaluate activity, energy expenditure, feeding, drinking, and respiratory quotient over the course of 48 hours. The data presented in graphs are the 24-hour averages normalized to body weight. Clamps were performed following a previously described protocol (66).

MEF cultures and differentiation into adipocytes. Primary MEF cultures were established from postnatal day 0 mice. 48 hours after confluence, MEFs were induced to undergo adipogenic differentiation.

Gene expression analysis. Total RNA was isolated from cells or tissues using the RNeasy Kit or the RNeasy Lipid Tissue Kit (Qiagen) and DNase treatment. Quantitative PCR was performed with the Applied Biosystems 7500 Fast RT-PCR system and SYBR Green detection reagent using cDNA synthesized with the SuperScript III First-Strand Synthesis System (Invitrogen). Primer sequences are provided in the Supplemental Methods. Gene expression analysis of adipocyte genes was conducted at the Duke Microarray Shared Resource (Duke University). 500 ng of total RNA was amplified according to the MessageAmp Premier protocol (Ambion) and hybridized to the Mouse 430 2.0 GeneChip (Affymetrix) according to the manufacturer’s instructions. Partek Genomics Suite 6.6 (Partek Inc.) was used to perform data analysis.

Membrane fractionation assay. 3-month-old male mice were fasted overnight and injected subcutaneously either 0.5 U/kg insulin or saline solution. Hind limb muscles and epididymal fat pads were carefully dissected out 30 minutes after injection and snap-frozen in liquid nitrogen. Plasma membrane fractions from skeletal muscle or adipose tissue lysates were obtained by differential ultracentrifugation as described previously (67, 68). We also used a similar protocol for isolation of plasma membrane fractions from differentiated adipocytes (68).

2-Deoxy-D-glucose uptake assay. Differentiated adipocytes cultured in 12-well plates were assessed for their ability to uptake glucose 9 days after induction of differentiation following a previously published protocol (69).

Quantification of insulin signaling pathway. Changes in the activation of the insulin pathway were assessed from total protein lysates from differentiated adipocytes that were either untreated (time 0) or stimulated with 100 nM insulin. For in vivo analysis, mice were fasted for 8 hours and administered either saline solution (untreated) or 0.8 U/kg insulin through intraperitoneal injection. Cells were collected at various time points after treatment with insulin, and tissues were collected 20 minutes after treatment. Immunoblotting of total lysates was performed using anti-AKT and anti-α-AKT (S473) antibodies.

Analysis of GLUT4 translocation and endocytosis by immunofluorescence. Differentiated adipocytes expressing Myc-GLUT4-GFP alone or in combination with 3xHA-AnkB were starved in serum-free DMEM containing 0.2% BSA for 2 hours at 37°C. Cells were either treated or not (basal) with 100 nM insulin for 30 minutes at 37°C. The population of plasma membrane–associated GLUT4 was labeled with mouse anti-Myc antibody for 1 hour on ice. Cells were then either fixed with 4% PFA (basal and time 0) or incubated at 37°C for 5, 10, 15, 20, and 30 minutes to allow for GLUT4 internalization.

Image acquisition. Fluorescent antibody and dye labeling was visualized using a 780 laser scanning confocal microscope (Zeiss). Z-stacks with optical sections of 0.5-μm intervals were collected using the ×20 (0.8 NA) and ×40 oil (1.3 NA) objective lens. Single plain images of Oil Red O–stained adipocytes were taken using the ×20 objective lens of a TE200 inverted microscope (Nikon).
Statistics. GraphPad Prism (GraphPad Software) was used for statistical analysis. Two groups of measurements were compared by unpaired, 2-tailed Student’s t test. Multiple groups were compared by repeated-measures 1-way ANOVA, followed by a post-hoc Tukey test. Survival curves were compared by a Mantel-Cox test. Results are presented as mean ± SEM. P < 0.05 was considered significant.

Study approval. Experimental procedures were approved by the Institutional Animal Care and Use Committee of Duke University.

Acknowledgments
We thank J.S. Bogan for the gift of the Myc7x-GLUT4-GFP construct, Gerald I. Shulman and Michael Jurczak at the Yale MMPC for technical assistance with the studies, the Duke Transgenic Mouse Facility for help generating the knockin mice, the Duke Microarray Shared Resource facility for help with gene expression analysis, and Christopher B. Newgard, Deborah Muoio, and Larry Moss at the Duke Molecular Physiology Institute for their valuable suggestions. M. Zhang acknowledges grant support (663812 and AoE/M09/12) from the Research Grant Council of Hong Kong. Mouse metabolic cage and insulin clamp analysis were supported by Yale MMPC grant U24 DK-059635.

Address correspondence to: Vann Bennett, Duke University Medical Center, 214A Nanaline Duke, 307 Research Drive, Box 3711, Durham, North Carolina 27710, USA. Phone: 919.684.3538; E-mail: vann.bennett@dm.duke.edu.


