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Introduction
Insulin regulates peripheral energy homeostasis by acting on multiple tissues to control carbohydrate, lipid, and protein metabolism (1). Gene targeting in mice has shown that β cell deletion of the insulin receptor causes reduced first-phase insulin release, reduced β cell insulin content, and progressive deterioration in glucose tolerance (2). Early studies of the effects of insulin in the CNS demonstrated a role for intracerebroventricularly administered insulin in the control of food intake and body weight (3). Mouse brain insulin receptor deletion causes mild hyperglycaemia and adiposity in female mice, diet-sensitive obesity, and defects in reproductive function (4). Results from studies in which insulin mimetics and insulin receptor antisense were centrally administered also support a role for CNS insulin signaling in energy homeostasis regulation (5, 6). Insulin signaling mechanisms therefore regulate β cell and CNS function, but it is unclear which postreceptor components mediate which physiological effects and, in the case of the CNS, which neuronal populations are involved. It is also unclear how insulin signaling components interact with other molecules involved in energy homeostasis, such as leptin.

Insulin receptor substrate (IRS) proteins lie downstream of the activated insulin and type 1 insulin-like growth factor receptor (7). Gene targeting studies have revealed distinct physiological roles for the 4 major IRS proteins (7). Mice lacking IRS1 display reduced islet mass, impaired glucose tolerance, and β cell dysfunction (10). These studies have suggested that IRS2 is the major mediator of the metabolic effects of insulin and have identified a novel role for IRS2 in insulin resistance (12). IRS2 signaling also plays complex roles in neuroendocrine function. Female IRS2-null mice are infertile, hyperphagic, and develop obesity (11). IRS2-null mice are reported to be leptin resistant (12), which suggests that IRS2 acts as a point of convergence for leptin and insulin signaling.

It has recently been reported that mice lacking IRS2 in both pancreatic β cells and a poorly defined hypothalamic neuronal population (generated using a rat insulin 2 promoter Cre [RIPCre] recombinase transgene) display reduced islet mass, impaired glucose tolerance, and hypothalamic dysfunction (13, 14). Unlike mice with global deletion of IRS2, these mice did not develop progressive diabetes. One study (13) reported that they displayed a hypothalamic phenotype suggesting melanocortin dysfunction, proposed that RIPCre neurons...
are proopiomelanocortin (POMC) neurons, and noted β cell recovery with time through an undetermined mechanism. Another study (14) demonstrated reduced islet mass, obesity, and leptin resistance but did not report other features of hypothalamic dysfunction or long-term analysis of islet function. Consequently, these studies have raised a number of critical but unanswered questions (15). In particular it is unclear (a) why there is a relative recovery of the β cell phenotype; (b) what role the CNS plays in the β cell phenotype and the impairment of glucose homeostasis; (c) what is the role of Irs2 in leptin action; and (d) what are the identity and characteristics of the hypothalamic neuronal population involved (15).

To address these questions, we generated mice lacking Irs2 in specific cell types: RIPCreIrs2KO mice, lacking Irs2 in β cells and a poorly characterized population of hypothalamic neurons; NesCreIrs2KO mice, lacking Irs2 in all neurons but not the endocrine pancreas; and POMCCreIrs2KO mice, lacking Irs2 in POMC-expressing cells.

Results

Generation of cells with a floxed allele of Irs2. We generated mice with a floxed allele of Irs2 (Irs2lox mice) to permit its deletion in different cell types and tissues (Figure 1, A–E). Irs2lox mice, when bred to homozygosity, were phenotypically indistinguishable from wild-type animals and displayed normal Irs2 expression (data not shown).

Using 3 Cre recombinase transgenic lines, we generated RIPCreIrs2KO, NesCreIrs2KO, and POMCCreIrs2KO mice. We also intercrossed RIPCre and POMCCre mice with ZEG (16) and Rosa26lacZ (17) indicator mice on both wild-type and Irs2lox/lox backgrounds. This allowed identification of cell types in which recombination occurred and permitted Irs2 deletion and GFP or lacZ expression in the same cell types.

Figure 1

Generation of Irs2floxed mice and characteristics of RIPCre and POMCCre mice. (A) Schema of targeting construct design, simplified restriction map of the Irs2 locus, the locus after homologous recombination and the deletion of neomycin cassette (Neo), and Southern blotting and PCR genotyping strategies used to identify these events. External probe A was used to identify homologous recombination (HR), probe B to detect the selection cassette, and probe C to detect the coding region of Irs2. HSV-tk, herpes simplex virus thymidine kinase. (B) Southern blot analysis with probe A demonstrating homologous recombination after targeting. (C and D) Southern blots using probe B after Cre-mediated recombination demonstrating deletion (Del) of the neomycin cassette and using probe C to demonstrate retention of Irs2 coding region confirming type 2 recombination. (E) PCR analysis with primers P1 and P2 of HR clone that has lost the neomycin cassette but retained the loxP site downstream of the Irs2 coding region. (F–H) We examined Cre expression in RIPCre mice by analyzing GFP expression in pancreatic sections costained with insulin and hypothalamic sections from RIPCreZEG mice. (I) We examined Cre expression in POMC Cre mice by analyzing hypothalamic sections from POMCCreZEG mice. (J) Combined ISH for POMC and ICC for GFP was performed in POMCCreZEG mice to confirm Cre expression in POMC neurons. (K) Immunohistochemistry for Irs2 (red) was performed in hypothalamic sections from POMCCreZEG mice. Scale bars: 50 μm (H, I, and K) and 10 μm (J), 3V, third ventricle; ME, median eminence.
Irs2 expression was found in pancreatic islets and in the arcuate, the VMH, and the paraventricular nucleus (Figure 1K and Figure 2B and data not shown).

To confirm deletion of Irs2 in the RIPCreIrs2KO, NesCreIrs2KO, and POMCCreIrs2KO mice, we analyzed recombination of the Irs2flax allele by PCR in islet and hypothalamic DNA. In 4-week-old mice, the recombined allele was detected in both tissues in RIPCreIrs2KO mice (Figure 2A) but only in the hypothalami of NesCreIrs2KO and POMCCreIrs2KO mice (Figure 2A). Immunoprecipitation and Western blotting confirmed the lack of Irs2 expression in isolated islets from RIPCreIrs2KO and in either whole brain or hypothalamic lysates from NesCreIrs2KO mice (Figure 2B). ICC analysis of RIPCreZEG mice demonstrated Irs2 colocalization with GFP in some but not all hypothalamic arcuate neurons (Figure 2D, left panel). In RIPCreIrs2KOZEG mice, there was no colocalization of Irs2 and GFP, which confirmed Irs2 deletion in RIPCre neurons (Figure 2D, right). Irs2 expression remained in other hypothalamic neurons. Analysis of Irs2 expression in POMCCre mice demonstrated colocalization of Irs2 with GFP in some POMC neurons (Figure 2E). Irs2 deletion was confirmed in POMCCreIrs2KOZOE mice (we were unable to breed POMCCreIrs2KOZOE mice), but Irs2 expression remained in other hypothalamic arcuate neurons (Figure 2F). Normal Irs2 expression was detected in muscle, liver, adipose, and other tissues in mice of all 3 lines (data not shown).

Glucose homeostasis in RIPCreIrs2KO, NesCreIrs2KO, and POMCCreIrs2KO mice. In 4-week-old male RIPCreIrs2KO, NesCreIrs2KO, and POMCCreIrs2KO mice, fasting blood glucose levels were indistinguishable from those of control animals (data not shown). At 12 weeks of age, when male Irs2-null mice are diabetic, RIPCreIrs2KO mice had higher fasting blood glucose levels than control mice, and glucose tolerance tests revealed defective glucose disposal (Figure 3, A and B). However, the elevation in fasting blood glucose levels was modest, and thus RIPCreIrs2KO mice did not display the overt diabetes and catabolic state that Irs2-null mice develop. Fasting glucose levels were also elevated in NesCreIrs2KO mice, and glucose tolerance was mildly impaired (Figure 3, A and C). RIPCreIrs2KO and NesCreIrs2KO mice at 6 months of age and older did not develop markedly elevated fasting blood glucose levels but remained glucose intolerant (Figure 3D and data not shown). In contrast, fasting glucose levels and glucose tolerance were equivalent to those of controls in POMCCreIrs2KO mice at 12 weeks and older (Figure 3A and data not shown).

At 12 weeks, RIPCreIrs2KO and NesCreIrs2KO mice, but not POMCCreIrs2KO mice, displayed fasting hyperinsulinemia compared with control animals, which suggests development of insulin resistance (Figure 3E). At 6 months, both RIPCreIrs2KO and NesCreIrs2KO mice remained hyperinsulinemic (data not shown). At 4 weeks, when there was a 50% reduction in β cell mass in Irs2-null mice (10), no decrease in β cell mass was seen in RIPCreIrs2KO compared with control animals (data not shown). However, by 12 weeks, RIPCreIrs2KO mice had a 40% reduction in β cell mass compared with control mice (Figure 3F, left), consistent with the deterioration in glucose homeostasis at this age. In contrast, 12-week-old NesCreIrs2KO mice exhibited an increased β cell mass, evidence of a compensatory response to insulin resistance (Figure 3F, left). POMCCreIrs2KO did not display altered β cell mass compared with age-matched controls (data not shown).

To investigate the preservation of glucose homeostasis compared with Irs2-null mice, we examined β cell mass in 9-month-old RIPCreIrs2KO mice. RIPCreIrs2KO mice had a reduced islet area compared with control animals (Figure 3F, right), but the area had increased from that seen in 12-week-old RIPCreIrs2KO mice, which suggests partial recovery in β cell mass (Figure 3F, right).

Hypothalamic function in RIPCreIrs2KO, NesCreIrs2KO, and POMCCreIrs2KO mice. RIPCreIrs2KO and NesCreIrs2KO mice displayed increased body weight compared with control mice starting around 4–5 weeks of age. By 12 weeks of age, they were 20% heavier than control animals (Figure 4A). RIPCreIrs2KO and NesCreIrs2KO mice were significantly longer than control mice (Supplemental Figure 1; supplemental material available online with this article; doi:10.1172/JCI200524445DS1). MRI scanning performed at 16...
weeks showed increased fat mass that was more significant in NesCreIrs2KO mice (Figure 4B). At 12 weeks, 24-hour food consumption was greater in both RIPCreIrs2KO and NesCreIrs2KO mice than in control animals (Figure 4C). In contrast, analysis of resting metabolic rate, core temperature, and activity revealed no differences between RIPCreIrs2KO or NesCreIrs2KO mice and control animals (Supplemental Table 1). Furthermore POMCCreIrs2KO mice did not display increased body weight (Figure 4A), were of the same length as controls (Supplemental Figure 1), and did not eat more than control mice (Figure 4C).

At 12 weeks of age when RIPCreIrs2KO and NesCreIrs2KO mice displayed marked obesity, leptin levels were significantly elevated in these mice but not in control mice (Figure 4A). In NesCreIrs2KO mice, leptin levels were significantly elevated after a 16-hour overnight fast (Figure 4B). To further investigate the temporal development of the feeding phenotype, we measured cumulative food intake with a 20% decrease in cumulative food intake (Figure 4C). In contrast, analysis of resting metabolic rate, core temperature, and activity revealed no differences between RIPCreIrs2KO or NesCreIrs2KO mice and control animals (Supplemental Figure 1). Furthermore POMCCreIrs2KO mice did not display increased body weight (Figure 4A), were of the same length as controls (Supplemental Figure 1), and did not eat more than control mice (Figure 4C).

Characterization of β cell recovery in RIPCreIrs2KO mice. Our studies and others (13, 15, 19) raise the question of the mechanism for the relative recovery of the β cell mass in RIPCreIrs2KO mice with age. Analysis of 4-week-old RIPCreIrs2KOZEG mice demonstrated that a vast majority of insulin-positive cells coexpressed GFP, which suggests that recombination was occurring in most β cells (Figure 5, A and B). However, we detected small numbers of insulin-expressing cells that were GFP negative (Figure 5B), which indicates lack of recombination and that the RIPCre promoter was not active in all β cells. It would be predicted that Ins2 would not be deleted in these cells and therefore they are involved in the preservation or recovery of β cell mass. Indeed, analysis of 9-month-old RIPCreIrs2KOZEG mice demonstrated that in many islets, a significant proportion of the insulin-positive β cells no longer expressed GFP (Figure 5, C and D). Comparison of the numbers of GFP-positive β cells in 9-month-old mice demonstrated that GFP expression loss occurred significantly more often in RIPCreIrs2KOZEG than control mice (percentage of GFP-negative β cells: RIPCreIrs2KOZEG, 50.9% ± 1.2% vs. RIPCreZEG, 10.1% ± 1.1%; P < 0.001; 2,062 β cells sampled in 54 islets from 4 RIPCreIrs2KOZEG mice and 2,987 β cells sampled in 45 islets from 3 RIPCreZEG mice).

To further characterize the mechanism of β cell recovery, we used a duct-specific lectin (Dolichos biflorus agglutinin [DBA]) to identify duct epithelial cells (20). GFP or insulin expression was not detected in DBA-positive duct structures in RIPCreIrs2KOZEG mice at 9 months of age (Figure 5, E–G). Likewise, in 9-month-old RIPCreIrs2KOZEG mice, in 296 islets sampled from 4 mice, no
DBA-positive cells were found within islets (Figure 5, E–G). These findings suggest that duct cells that have differentiated to insulin-producing cells within ducts or duct cells that migrate to islet structures before converting to β cells do not make a significant contribution to the β cell repopulation. These findings do not exclude the possibility that ductal precursor cells that express neither duct markers nor insulin are involved in this process.

Characterization of RIPCre neurons and comparison with POMC neurons. Our studies demonstrate that RIPCre neurons play a role in energy homeostasis and growth and that deletion of Ins2 in these cells generates a phenotype with similarities to that of melanocortin pathway mutants. However, our genetic evidence suggests that deletion of Ins2 in POMC neurons does not account for the hypothalamic phenotype seen in RIPCreIns2KO or NesCreIns2KO mice. RT-PCR analysis of expression of hypothalamic feeding peptides in RIPCreIns2KO and NesCreIns2KO mice demonstrated reduced POMC expression (POMC mRNA: 42% reduction in NesCreIns2KO mice normalized to control mice, \( P < 0.05, n = 5 \); 48% reduction in RIPCreIns2KO mice normalized to control mice, \( P < 0.05, n = 8 \)) but normal neuropeptide Y (NPY) and agouti-related protein (AgRP) expression (data not shown), which suggests an effect on melanocortin circuits. No alteration in hypothalamic POMC, AgRP, and NPY expression was seen in POMCCreIns2KO mice. Therefore, to characterize the relationship between RIPCre neurons and melanocortin pathways, we performed ISH/ICC analysis and electrophysiological studies of RIPCre and POMCCre neurons. POMC- and GFP-expressing neurons in the hypothalamus were clearly identified by ISH and ICC, respectively, in the same slices (Figure 6, A and B), and no GFP-positive neurons colocalized with POMC mRNA. POMC and RIPCre neurons, however, are closely associated with each other. Similarly, no coexpression of GFP was demonstrated in NPY neurons, but close apposition of these neuronal populations was observed (Figure 6, C and D). Combined ICC for α melanocyte-stimulating hormone (αMSH) and GFP demonstrated no coexpression, but many GFP neurons were in close proximity to αMSH-positive fibers, which suggests that they may be targets for POMC neurons (Figure 6E).

Whole-cell current-clamp recordings were made from GFP-positive hypothalamic arcuate neurons from RIPCreZEG mice. Under control recording conditions, RIPCre neurons spontaneously fired action potentials and had a mean resting potential of –50 ± 1 mV.
and input resistance of $1,862 \pm 197 \text{ M}\Omega$ ($n = 62$). Bath application of the αMSH analog melanotin II (MTII) (a nonselective melanocortin 3/4 receptor agonist) at 100 nM caused long-lasting depolarization in the majority of cells tested (from $-53 \pm 3 \text{ mV}$ to $-48 \pm 2 \text{ mV}; P < 0.01; n = 9/13$), associated with increased spike firing (Figure 6F, G and H).

To demonstrate the MTII excitatory effect more clearly, we applied a small hyperpolarizing current to individual neurons, such that action potential frequency was markedly reduced. Under these conditions, MTII (100 nM) caused depolarization and increased firing frequency (Figure 6H). Furthermore, the application of MTII (100 nM) to RIPCre neurons, following incubation of the slice with tetrodotoxin (TTX; 500 nM) to block spike firing and prevent synaptic transmission, caused significant depolarization ($n = 4$), which indicates a direct action (Figure 6I). To further characterize RIPCre neurons and differentiate them from POMC (and NPY) neurons, we determined the actions of insulin and leptin on the electrical properties of RIPCre and POMC neurons. Leptin has been demonstrated to depolarize POMC neurons (21), and our current results support this finding. Pressure ejection of leptin (50 nM) for 60 seconds directly above the recorded neuron caused an increase in spike frequency and a small depolarization (from $-47 \pm 2 \text{ mV}$ to $-44 \pm 3 \text{ mV}; P < 0.04$) of a minority population of POMC neurons ($n = 5/13$) (Figure 7, A–C). In contrast, leptin (50 nM) did not affect action potential frequency or resting membrane potential ($-46 \pm 3 \text{ mV}$ to $-47 \pm 3 \text{ mV}; n = 10$; Figure 7, D and E) in RIPCre neurons. Pressure ejection of insulin (10 nM) for 60 seconds produced a significant increase in action potential frequency, concomitant with cell depolarization (from $-49 \pm 4$ to $-40 \pm 4 \text{ mV}; P < 0.01$) in the majority of recordings ($n = 6/9$) from RIPCre neurons (Figure 7, F–H). In contrast, pressure injection of insulin (10 nM) onto POMC neurons resulted in hyperpolarization (from $-48 \pm 1 \text{ mV}$ to $-55 \pm 2 \text{ mV}; P < 0.001$) and reduced action potential frequency in most CNS. To elucidate the contribution of Irs2 to this complex phenotype in specific tissues, we generated mice with a conditional allele of Irs2 to permit deletion in a number of cell types. We initially used RIPCre transgenic mice to generate RIPCreIrs2KO mice lacking Irs2 in β cells and a population of hypothalamic neurons, and indeed it has recently been reported that RIPCreIrs2KO mice display phenotypes due to this mixed pattern of deletion (13, 14). These mice developed reduced β cell mass, obesity, somatic overgrowth, hyperinsulinemia, and hyperleptinemia but did not progress to an overt diabetic phenotype because of an unexplained recovery in β cell mass. However, because of the inability to dissociate β cell and hypothalamic phenotypes, a number of questions regarding the role of Irs2 remain unanswered (15).

To directly address such issues that we encountered with our own RIPCreIrs2KO mice, we also generated NesCreIrs2KO mice, lacking Irs2 in all neurons but not the endocrine pancreas, and PomccIrs2KO mice, lacking Irs2 in hypothalamic POMC neurons, which have been implicated in mediating leptin’s effects on energy homeostasis (23).

Consistent with recent reports (13, 14), 12-week-old RIPCreIrs2KO mice had reduced β cell mass and displayed mildly elevated fasting blood glucose levels, impaired glucose tolerance, and hyperinsulinemia. They developed a marked hypothalamic phenotype resembling that of mice with defects in melanocortin action. In contrast, NesCreIrs2KO mice, which have normal β cell Irs2 expression, had increased β cell mass in response to insulin resistance and displayed the obesity and overgrowth syndrome. These findings demonstrate that β cell mass reduction is due to Irs2 deletion solely in β cells and not in the CNS. However, NesCreIrs2KO did develop elevated fasting blood glucose levels and hyperinsulinemia, which suggests that neuronal Irs2 pathways are required for normal glucose homeostasis but that the increase in β cell mass seen in NesCreIrs2KO mice could compensate for insulin resistance.

**Figure 5**

Analysis of recombination and duct-marker expression in islets from RIPCreIrs2KO mice. (A–D) Immunofluorescence staining for insulin (red) in islets from 4-week-old (A and B) and 9-month-old (C and D) RIPCreIrs2KO mice stained for DNA (DAPI, blue). White arrows indicate β cells that do not express GFP and have not undergone recombination. Many cells in the 9-month-old islet do not express GFP. (E–G) Labeling of ducts with DBA in islets from 9-month-old RIPCreIrs2KO mice stained blue for either DNA (E and F) or insulin (G). No duct marker is expressed in GFP-negative β cells. Scale bars: 100 μm.

The effect of insulin, MTII, and leptin on neuronal activity was extremely slow to reverse and was, in many cases, irreversible over the time course of the experiments. These experiments suggest that RIPCre neurons are a population distinct from either POMC or NPY neurons but lie within a melanocortin- and insulin-regulated neuronal mechanism.

**Discussion**

Through global gene deletion, we and others have previously implicated Irs2 in the regulation of β cell function and peripheral glucose metabolism and the central regulation of energy homeostasis (10, 11, 19, 22). These studies have suggested but not proved that Irs2 mediates both β cell growth and leptin and insulin action in the...
In contrast to Lin et al. (13), we found that in 9-month-old RIPCreIr2KO mice, there was a persistent reduction in β cell mass compared with that of control mice, although the mass was increased compared with that of 3-month-old RIPCreIr2KO mice. At later ages, therefore, RIPCreIr2KO mice had glucose intolerance but did not display marked fasting hyperglycemia and never developed the marked progressive diabetic phenotype of the global Irs2-null mouse.

Anatomical studies of recombination using a GFP indicator mouse in either a wild-type or Irs2flox background revealed that as early as 4 weeks, small numbers of β cells did not harbor the recombination event. By 9 months of age, significant numbers of β cells in RIPCreIr2KO mice not expressing GFP had repopulated the islets, but many GFP-positive cells undergoing recombination were still present, which explains the persistent reduction in β cell mass. Our studies demonstrate that the reduction in recombination occurs predominantly in RIPCreIr2KOZEG as opposed to RIPCreZEG islets, which suggests that Irs2 signaling itself is required for expression of the RIPCre transgene. Indeed, insulin gene expression appears to require an intact insulin signaling pathway in the β cell (2, 24, 25). With Irs2 largely absent within β cells, under conditions of insulin resistance there may be potential selection for pre-existing β cells not expressing the RIPCre transgene and therefore not undergoing deletion of Irs2 to expand and repopulate the islet. In 9-month-old mice, we did not detect GFP expression and Cre-mediated recombination in pancreatic ducts, and so potentially the GFP-negative β cells could have arisen from duct cells. However, we could not detect expression of a duct cell marker within islets of 9-month-old RIPCreIr2KO mice nor could we find insulin expression in ducts. The absence of deletion in ducts also suggests that the β cell phenotype in RIPCreIr2KO mice results from loss of

Figure 6
Characterization of RIPCre neurons by ICC, ISH, and electrophysiological analysis. (A and B) Dual ISH for POMC and ICC for GFP was performed on hypothalamic sections from RIPCreZEG mice. Representative low- (A) and high-power (B) views are shown. (C and D) Dual ISH for NPY and ICC for GFP was performed on hypothalamic sections from RIPCreZEG mice. Representative low- (C) and high-power (D) views are shown. ARC, arcuate nucleus. (E) Fluorescence ICC for POMC was performed on hypothalamic sections from RIPCreZEG mice. Neurons expressing POMC are red and those expressing GFP are green. Arrows indicate noncolocalized neurons and GFP neurons that are abutted by POMC fibers. A representative confocal image is shown. Scale bars: 50 μm (A and C) and 10 μm (B, D, and E). (F–I) Current-clamp recordings were made from RIPCreZEG neurons in the absence and presence of the mixed melanocortin 3/4 receptor agonist MTII, as indicated above the traces. (F) Representative recording demonstrating MTII depolarization, resulting in a small increase in spike frequency (G). This depolarization is more clearly represented in neurons that have sub-spike thresholds, achieved by application of a constant hyperpolarizing current (H). MTII also depolarizes neurons in the presence of TTX (I), indicative of a direct action. The action potentials in panel H have been truncated to show clearly the depolarizing effect of MTII. Note that the MTII depolarization was irreversible over the time course of the recordings in cells represented in H and I.
Irs2 in mature insulin-producing cells rather than any precursor or stem cell population. Taken together, these findings provide anatomical evidence that a small number of preexisting β cells that escape Irs2 deletion at an early stage in RIPCreIrs2KO mice repopulate the islets. Our studies cannot categorically exclude the possibility that duct-derived precursors expressing neither insulin nor duct markers contribute to the repopulation process. However, recent lineage tracing studies (20) suggest that proliferation of mature β cells is the major mechanism for β cell expansion in adult mice. Our findings suggest that persistent Irs2 expression in mature β cells is required for the compensatory response to insulin resistance. The preservation of β cell mass in NesCreIrs2KO mice dissociates hypothalamic dysfunction from reduced β cell mass. It has been suggested that nestin expression is a characteristic of pancreatic islet stem cells (26, 27), but recently lineage tracing studies utilizing the same NesCre mouse used in our studies, demonstrated that pancreatic exocrine, ductal, and endothelial cells but not islet endocrine cells express nestin (18, 28). The finding of increased islet mass in NesCreIrs2KO mice further supports the concept that nestin-positive cells do not contribute to the endocrine pancreas, as deletion of Irs2 in these cells would have been anticipated to reduce β cell mass.

The hypothalamic phenotypes seen in RIPCreIrs2KO and NesCreIrs2KO mice suggested defects in melanocortin function, and in RIPCreIrs2KO and NesCreIrs2KO mice, we detected reduced hypothalamic POMC expression. Deletion of the leptin receptor in POMC neurons generates mice that have some phenotypic features in common with RIPCreIrs2KO mice, including obesity (23). It has been suggested that RIPCre neurons are POMC neurons and that Irs2 signaling events are required for central leptin action (14, 29). Our results, however, do not confirm these findings and provide evidence to define the role and sites of Irs2 pathways in the CNS. We demonstrate that deletion of Irs2 in POMC neurons did not recapitulate the hypothalamic phenotype and that RIPCre neurons do not express POMC or NPY. Furthermore, RIPCreIrs2KO and NesCreIRSKO mice were sensitive to peripherally administered leptin at 6 weeks of age. Five-week-old RIPCreIrs2KO and NesCreIRSKO mice are hyperphagic but not hyperleptinemic, which suggests that elevated leptin levels seen at later ages result from greater adiposity due to increased energy intake rather than primary central leptin resistance. This is consistent with the observation that central leptin administration to Irs2-null mice causes reduction in fat mass (12). Young nonobese hyperphagic melanocortin 4 receptor–null mice, which resemble RIPCreIrs2KO and NesCreIRSKO mice, are sensitive
to the anorexigenic effects of leptin but develop leptin resistance as they become obese (30). In preliminary observations, mice lacking Irs2 in AgRP neurons, generated using an AgRPCre transgenic mouse (31), also do not develop a marked hypothalamic phenotype (H. Al-Qassab et al., unpublished observations). Therefore, these studies implicate Irs2 signaling in neurons distinct from POMC and AgRP/NPY populations in the regulation of energy homeostasis and growth and suggest that Irs2 is not an obligatory component of CNS leptin signaling pathways.

Electrophysiological analysis showed that RIPCre neurons depolarized in response to insulin and MTII, which suggests that they could be mediating the effects of insulin in the CNS in a melanocortin-sensitive neuronal pathway. Furthermore, we found no electrophysiological effect of leptin upon arcuate RIPCre neurons, which suggests that this is not a leptin-sensitive neuronal population and therefore may explain why RIPCreIrs2KO mice are not leptin resistant. In contrast, leptin depolarized and insulin hyperpolarized POMC neurons. These opposite effects upon neuronal activity perhaps explain the lack of phenotype in POMCCreIrs2KO mice and distinguish RIPCre neurons electrophysiologically from both POMC and NPY neurons (21, 32). It remains possible that the RIPCre neurons in other hypothalamic regions such as the VMH may be mediating the effects of leptin, and we have started to examine this possibility using additional mouse mutants. However, our observation of leptin sensitivity in the NesCreIrs2KO mice suggests that CNS Irs2 is not required for leptin action.

Our studies have demonstrated that CNS Irs2 pathways acting in a neuronal population distinct from POMC and NPY neurons regulate energy homeostasis and growth. Arcuate RIPCre neurons are targets for insulin and melanocortin but not leptin action, but it remains to be determined what neuropeptides and transmitters are expressed in this population. Irs2 in β cells is required for the maintenance of β cell mass, but β cells that escape Cre-mediated recombination are able to repopulate islets with time. Our observations clarify the role of Irs2 in β cell function and energy homeostasis and suggest that modulation of Irs2 function may be of use in the treatment of diabetes and obesity.

Methods

Generation of floxed allele of Irs2 and genotyping strategies for Irs2lox and Cre and ZEG transgenic mice. The murine Irs2 gene was cloned from a 129Sv mouse genomic PAC library. We cloned 2.8 kb of 5′ homology from the Irs2 locus upstream of a floxed neomycin cassette using appropriate linkers. The Irs2 coding region was subcloned with linkers, introducing a loxP site 3′ to the Irs2 coding region. A 3-kb fragment of 3′ region of the Irs2 locus was cloned downstream of this loxP site using linkers to generate the final construct (Figure 1A). 129Sv/EvTac1 ES cells were transfected with Pac1-linearized targeting construct by electroporation. Cells were subjected to selection with G418 and ganciclovir; surviving clones were expanded and genomic DNA extracted and subjected to analysis by PCR and Southern blotting to detect homologous recombination (Figure 1, B–D). Targeted clones were subsequently transfected with Cre recombinase plasmid to remove the neomycin cassette and screened by Southern blotting and PCR (Figure 1, C and D). ES cells were injected into blastocysts derived from C57BL/6J mice, which were implanted into pseudopregnant CD-1 foster mothers. Chimeric male mice were bred with C57BL/6J mice and germ-line transmission confirmed by PCR and Southern blot analysis. Irs2lox mice were intercrossed with C57BL/6J RIPCre and C57BL/6J NesCre mice obtained from the Jackson Laboratory (33, 34) to generate compound heterozygous mice. Double-heterozygous mice were crossed with Irs2lox mice to obtain WT, Irs2lox/+, Cre+, and CreIrs2lox/lox mice. Mice lacking Irs2 in RIPCre-expressing cells were designated RIPCreIrs2KO and in nestin Cre–expressing cells NesCreIrs2KO. POMCCre mice were generated using a BAC transgenic approach (35). POMCCre mice were intercrossed with Irs2lox mice and bred as described above to obtain POMCCreIrs2KO mice. ZEG mice, which carry a Cre-activated GFP transgene (16), or Rosa26lacZ indicator mice (17) were intercrossed with the RIPCre and POMCCre and Irs2lox mice to obtain mice that express GFP or lacZ in RIPCre- or POMCCre-expressing cells as well as mice that also lack Irs2 and express the indicator gene. Mice were maintained on a 12-hour light/dark cycle with free access to water and standard mouse chow (4% fat, RM1; Special Diet Services) and housed in specific pathogen–free barrier facilities. Mice were handled and all in vivo studies performed with approval by the Home Office (London, United Kingdom). All knockout and transgenic mice were studied with appropriate littermate controls. Wild-type, Cre transgenic and Irs2lox/lox mice were phenotypically indistinguishable, and equal numbers of the mice of these genotypes were used as controls. Genotyping of the mice was performed by PCR amplification of tail DNA. For Irs2lox genotyping, we used primers that flank the 3′ loxP site (Figure 1E): loxP forward, 5′-ACCTGAAGGAAGCCACAGTGCC-3′ and loxP reverse, 5′-AGTCCTACTTTCTGACAGAC-3′. Cre recombinase, Rosa26lacZ, and ZEG genotyping was performed as previously described (16, 17). For detection of Cre-mediated excision of Irs2, genomic DNA was isolated from pancreatic islets, hypothalamus, and other tissues by heating in 50 mM NaOH at 100 °C for 10 minutes. Primers located 1.1 kb upstream of the 5′ loxP site (5′-GGGAACTCAGACAGAATG-3′) and 0.2 kb downstream of the 3′ loxP site (loxP 5′-AGTCCTACTTTCTGACAGAC-3′) amplified a 1.3-kb product if recombination had occurred.

Immunoprecipitation and Western blotting. Tissues were removed and homogenized in lysis buffer, solubilized for 30 minutes on ice, and clarified by centrifugation. Supernatants were immunoprecipitated for 2 hours with an anti-Irs2 antibody as described (10). Immune complexes were collected with 100 μl of 50% slurry of protein A–sepharose, washed with buffer, resolved on 7.5% SDS-PAGE, and transferred to nitrocellulose. The blots were probed with polyclonal antibody against Irs2 and enhanced chemiluminescence (Amersham Biosciences) detection.

Metabolic studies. Body weights were determined using a Sartorius BP610 balance. Nose to anus length was measured either postmortem or in anesthetized mice, with the observer blinded to genotype. Blood samples were collected from mice via tail vein bleeds or from cardiac puncture performed on terminally anesthetized mice. Blood glucose was measured using a Glucometer Elite (Bayer Corp.). We determined plasma insulin levels using an ultrasensitive rat insulin ELISA (Crystalchem Inc.) using a mouse standard and leptin levels using a mouse leptin ELISA (Crystalchem Inc.). Glucose tolerance tests were performed on mice after a 16-hour overnight fast. Animals were injected intraperitoneally with 0.5-g glucose (1.5 g/kg) and blood glucose levels determined at the time points indicated in Figure 3, B and C.

Food intake and leptin and treatment studies. Mice were singly housed and allowed to acclimatize for a week prior to study. Food intake and body weight were measured for 3 consecutive days at the ages indicated in the Figure 4 legend. As necessary, we acclimatized mice by subjecting them to overnight fasts and sham injections prior to the study protocol. For peripheral leptin treatment, mice were treated with either 5 mg/kg recombinant mouse leptin (R&D Systems) or vehicle injected i.p. 1 hour before lights out for 3 consecutive days. Food and body weights were recorded before injection, during the treatment period, and for a 3-day wash-out period following the study.

ICC and ISH. Brains were harvested following transcardiac perfusion with RNase-free PBS and 4% paraformaldehyde, postfixed in 4% paraformaldehyde, and transferred to RNase-free 30% sucrose prior to freezing. Thirty-micrometer floating sections were cut using a sliding microtome. Twenty-micrometer cryostat sections were thaw-mounted on polylysine-coated microscope slides. Floating sections were washed
in potassium PBS (KPBS), pH 7.4, blocked with 2% serum in KPBS/0.4% Triton X-100 (KPBS-T) for 1 hour at room temperature, and then incubated with primary antibodies in KPBS-T for 48 hours at 4°C. Slices were incubated with appropriate secondary antibodies for 1 hour in KPBS-T before being washed in KPBS. For chromogenic detection, slices were incubated for 60 minutes in avidin-DH/biotinylated HRP (ABC Elite; Vector Laboratories) in KPBS-T and washed in KPBS, then in 150 mM sodium acetate, and nickel-enhanced detection was performed with a 3,3′-diaminobenzidine substrate kit for peroxidase (Vector Laboratories). For some anti-αMSH staining experiments, mice were treated with intracerebroventricular colchicine (2 μl of 20 mg/ml solution in 0.9% saline infused over 10 minutes into the lateral ventricle) 24 hours prior to perfusion fixation and biotinylated tyramide amplification and fluorescent detection with streptavidin-conjugated Alexa Fluor 594 (Invitrogen Corp.) were used. Slices were mounted on polylysine-coated microscope slides and coverslipped with buffered glycerol. Primary antibodies used were rabbit anti-Irs2 antibody (United Biomedical Inc.), sheep anti-NPY (Chemicon International), sheep anti-αMSH (Chemicon International), mouse monoclonal anti-GFP (Chemicon International), and mouse monoclonal anti-lacZ (Promega). For combined ISH and ICC, ISH was performed first under RNase-free conditions. In brief, sections on slides were fixed in formaldehyde solution for 20 minutes on ice, washed twice in PBS for 5 minutes, and acetylated at room temperature for 10 minutes in 0.1 M triethanolamine (pH 8.0) with acetic anhydride. After washing, sections were dehydrated and air dried for 30 minutes. Hybridization buffer containing 2 ng/μl digoxigenin-labeled (DIG-labeled) probe was applied and sections covered with glass coverslips and hybridized overnight at 60°C. Sections were placed in 4× SSC to remove coverslips, RNase A treated for 30 minutes at 37°C, and washed in 2× SSC at 6°C for 10 minutes, 1× SSC at 6°C for 10 minutes, 0.5× SSC at 60°C twice for 10 minutes, and 0.1x SSC at 60°C for 30 minutes. Sections were cooled to room temperature in 0.1x SSC, and we used alkaline phosphatase-conjugated anti-DIG antibody to detect bound DIG-labeled probe (Roche Diagnostics Ltd.) as described above. We generated ISH riboprobes using mouse sequences (POMC, GenBank accession number NM_008895; NPY, GenBank accession number NM_023456). Imaging was performed with an Olympus BX51 microscope with either a Hamamatsu 95 black-and-white camera or a Jenoptik ProRes C14 color camera combined with SimplePCI capture and deconvolution software (version 5.2.1, Comipix Inc.). Confocal microscopy was performed on a Bio-Rad MRC1000 microscope. Colocalization of either POMC or NPY with GFP was scored in 4 arcuate sections from each of 4 RIPCrewZEG mice by 2 independent observers.

Pancreatic ICC and measurement of islet mass and number. Pancreata were removed, cleared of fat and lymph nodes, fixed in Bouins solution, embedded in paraffin, and cut into 5-μm sections. For GFP detection, pancreata were harvested from paraformaldehyde-perfused mice as described above. For insulin staining, sections were incubated with blocking solution containing PBS buffer with 5% normal chicken serum and 2% BSA for 30 minutes, and mouse anti-insulin antibody (clone K36aC10; Sigma-Aldrich) was then applied for 2 hours at room temperature or overnight at 4°C. The sections were then incubated with chicken anti-mouse IgG–Alexa Fluor 488 conjugate (Invitrogen Corp.) for 2 hours at room temperature. Transmitted light and fluorescent images were captured as described above. For quantification of β cell area, we analyzed 5 pancreata per genotype at each time point using 4 sections at least 150 μm apart from each other. For each section, the total area occupied by insulin-positive cells was scored using SimplePCI software. Results are expressed as the percentage of the total pancreatic area. Pancreatic ducts were labeled with biotinylated DBA lectin (Vector Laboratories), detection performed with fluorescently labeled streptavidin 610 (Invitrogen Corp.), and DAPI used to identify nuclei.

**Islet isolation.** Mice were sacrificed, the common bile duct cannulated, and its duodenal end occluded by clamping. Two milliliters of collagenase V solution (1 mg/ml in HBSS) was injected into the duct to distend the pancreas. The pancreas was excised, incubated at 37°C for 15 minutes, and mechanically disrupted in 10 ml of HBSS. Cellular components were collected by centrifugation (220 g for 1 minute) and resuspended in 10 ml of HBSS. Islets were handpicked under a microscope and washed once in HBSS. Islets were centrifuged at 15,600 g for 5 minutes and stored at −20°C until DNA or protein extraction.

**mRNA quantification by RT-PCR analysis.** Hypothalami were removed from fasted mice, homogenized in Trizol reagent (Invitrogen Corp.), and 1 μg RNA was reverse transcribed using Taqman reagents (Applied Labs.). Quantitative PCR was carried out with first-strand cDNA with primers for hypothalamic feeding peptides (23), using hypoxanthine ribosyl transferase as an internal control on an ABI Prism 7900HT sequence detection system, with SYBR green reagents (Applied Biosystems Inc.), and data analyzed according to the 2−ΔΔt method (36).

**Electrophysiology.** Male RIPCrewZEG and POMCCrewZEG mice (8–16 weeks old) were killed by cervical dislocation. Brains were rapidly transferred to a ice-cold slicing solution containing 2.5 mM KCl, 1.25 mM NaH2PO4, 28 mM NaHCO3, 0.5 mM CaCl2, 7 mM MgCl2, 7 mM t-glucose, 1 mM ascorbate, 3 mM pyruvate, and 235 mM sucrose and equilibrated with 95% O2/5% CO2 to give a pH of 7.4, and 350-μm coronal brain slices were prepared using a Vibratome Series 1000. Arcuate nucleus slices were kept at room temperature in a normal external solution containing 125 mM NaCl, 2.5 mM KCl, 1.25 mM NaH2PO4, 25 mM NaHCO3, 2 mM CaCl2, 1 mM MgCl2, 10 mM t-glucose, 15 mM t-mannitol, 1 mM ascorbate, and 3 mM pyruvate equilibrated with 95% O2/5% CO2, pH 7.4. For whole-cell recordings, slices were continuously perfused with normal external solution (in the absence of ascorbate and pyruvate and with CaCl2 and MgCl2 concentrations adjusted to 0.5 and 2.5 mM, respectively) at a rate of 5–10 ml/min. Neurons were identified using epifluorescence and differential interference contrast optics using an upright Zeiss AxioSkop 2 FS plus microscope. Borosilicate patch pipettes (5–8 MΩ) were filled with an internal solution containing 130 mM potassium gluconate, 10 mM KCl, 0.5 mM EGTA, 10 mM HEPES, 1 mM NaCl, 0.28 mM CaCl2, 3 mM MgCl2, 3 mM NaATP, 0.3 mM tris-GTP, 14 mM phosphocreatine (pH 7.2). Data were recorded and stored on digital audio tape, which enabled off-line analysis. We examined changes in input resistance by monitoring membrane potential responses to negative, rectangular current pulses (5–20 pA, 200 ms, 0.05 Hz) injected via the recording electrode. Application of drugs was made via the bath perfusion system or locally by pressure ejection from a broken pipette (approximately 5-μm tip) positioned approximately 50 μm above the recorded neuron at the concentrations indicated.

**MRI and magnetic resonance spectroscopy.** Mice were scanned using a 4.7T Varian system under isoflurane anesthesia. Whole-body images (between 50 and 60 slices; 2 mm thick) were obtained for each mouse using a spin-echo sequence (TR4500/TE20). Semiautomated image segmentation software (sliceOmatic version 4.2; Tomovision) was used to separate and quantify tissue volumes (37). For adipose tissue, the volume was multiplied by a correction factor of 0.9 in order to account for hydration (38).

**Statistical analysis.** All statistics were performed using GraphPad Prism 4 software and paired and unpaired Student’s t tests and 2-way ANOVA, with Bonferroni post-hoc tests performed as appropriate.

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