Metabolic effects of leptin receptor knockdown or reconstitution in adipose tissues

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The relative contribution of peripheral and central leptin signalling to the regulation of metabolism and the mechanisms through which leptin affects glucose homeostasis have not been fully elucidated. We generated complementary lines of mice with either leptin receptor (Lepr) knockdown or reconstitution in adipose tissues using Cre-lox methodology. Lepr knockdown mice were modestly lighter and had lower plasma insulin concentrations following an oral glucose challenge compared to controls, despite similar insulin sensitivity. We rendered male mice diabetic using streptozotocin (STZ) and found that upon prolonged leptin therapy, Lepr knockdown mice had an accelerated decrease in blood glucose compared to controls that was associated with higher plasma concentrations of leptin and leptin receptor. Mice with transcriptional blockade of Lepr (LeprloxTB/loxTB) were obese and hyperglycemic and reconstitution of Lepr in adipose tissues of LeprloxTB/loxTB mice resulted in males reaching a higher maximal body weight. Although mice with adipose tissue Lepr reconstitution had lower blood glucose levels at several ages, their plasma insulin concentrations during an oral glucose test were elevated. Thus, attenuation or restoration of Lepr in adipocytes alters the plasma insulin profile following glucose ingestion, modifies the glucose-lowering effect of prolonged leptin therapy in insulin-deficient diabetes, and may modulate weight gain.

Multiple studies have reported that leptin action in the central nervous system (CNS) is critical for its regulation of food intake, thermogenesis, glucose homeostasis, and insulin sensitivity1–8. The contribution of peripheral leptin signalling to the regulation of body weight and glucose metabolism has not been fully elucidated. From the single leptin receptor (Lepr) gene, multiple isoforms of Lepr exist, and it is believed that the long isoform (Leprb) is responsible for the majority of leptin's actions9,10. Using a tamoxifen-inducible model, where Leprb expression was reduced to different extents in peripheral tissues, but was not affected in the CNS, Guo et al.11 observed no alterations in metabolism, including insulin sensitivity, except for hyperleptinemia, which was higher in females. The hyperleptinemia resulted from increased leptin secretion from adipose tissue, possibly due to interference in a negative feedback loop, but leptin action was apparently not affected because plasma levels of free leptin were not changed. Mice with liver-specific Lepr knockdown on standard chow have improved insulin sensitivity and glucose tolerance, but when on high fat diet, only the latter is ameliorated12. Using antisense RNA expressed under the control of the phosphoenolpyruvate carboxykinase (PEPCK) promoter, Huan et al.13 downregulated expression of leptin receptor isoforms in mouse white adipose tissue (WAT). This resulted in obesity, insulin resistance, glucose intolerance, and ectopic fat accumulation. However, it has been suggested that the antisense mRNA approach to reduce gene expression, as used by Huan et al.13, may itself lead to obesity in rodents14. Wang et al.15 reported that overexpression of Leprb in adipose tissue using an aP2 promoter inhibited the elevation in body weight caused by high fat diet, but these alterations may not have been adipose tissue-specific because subsequently, studies with aP2-Cre mice also found Cre-mediated recombination in non-adipose tissues16. In the current study, our objective was to clarify the role of adipose tissue leptin signalling in metabolism using two complementary approaches based on Cre-lox methodology: generating mice lacking Leprb in adipose tissues and mice that express Leprb only in adipose tissues.

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Methods

Study design. Mice expressing Cre under the control of the Adiponectin promoter (AdipoqCre) were obtained from The Jackson Laboratory (Bar Harbor, ME, USA, stock# 010803). AdipoqCre mice produce recombination selectively in white and brown adipose tissues.17, AdipoqCre mice were mated with: 1) mice with floxed exon 17 of the leptin receptor gene (LeprΔ17) to generate mice that have diminished expression of Leprb due to excision of exon 17 in adipose tissues (ATLeprEXP colony) and 2) mice containing a transcripational blockade of the leptin receptor gene (LeprΔ17TB/loxTB), The Jackson Laboratory, stock#018989) to generate mice that express leptin receptors only in adipose tissues (ATLeprEXP colony). All AdipoqCre+ mice studied were heterozygous for Cre. Littermate controls were used for all studies. AdipoqCre, LeprΔ17TB/loxTB, and LeprΔ17TB/loxTB mice were on a mixed C57BL/6 and FVB background; contribution of 129 was minor. Ear notch samples were obtained to determine genotypes and genotyping is described in Supplemental Information. To assess Cre activity by immunofluorescence, AdipoqCre;LeprΔ17TB/loxTB mice were mated with mTmG mice19,20, which were on a mixed C57BL/6 and 129 background, and AdipoqCre;LeprΔ17TB/loxTB;ROSA26mTmG mice as well as AdipoqCre;LeprΔ17TB/loxTB;ROSA26mTmG;Cre+ mouse were studied. Breeders were fed Harlan diet 2919 and maintenance diet was standard chow (Harlan #2918). For high fat diet (HFD) and low fat diet (LFD) studies, mice were placed on HFD (60 kcal%, Cat# D12492i, Research Diets) or LFD (10 kcal%, Cat# D12450Bi, Research Diets) at 9 weeks of age. Mice were housed in a 12 h:12 h light-dark cycle, food and water were available ad libitum, and mice were euthanized following a 4 h fast unless stated otherwise. All procedures were approved by the University of British Columbia Animal Care Committee and followed the guidelines of the Canadian Council on Animal Care.

Streptozotocin (STZ) and leptin therapy studies. Two studies were performed to investigate the effects of leptin therapy in STZ-induced diabetes: (1) In the prolonged leptin therapy study, at 10–12 weeks of age, male AdipoqCre;LeprΔ17TB/loxTB mice and their AdipoqCre;LeprΔ17TB/loxTB male littermates were given 180 mg per kg body weight of STZ intraperitoneally (i.p.) (Sigma-Aldrich) on Day −8, while controls were not injected. Diabetes in STZ-injected mice was defined as blood glucose concentrations ≥18.0 mM following a 4 h morning fast on Days −3 and −1. On Day 0, diabetic STZ-mice were implanted subcutaneously with mini-osmotic pumps (DURECT Corporation, Cupertino, CA, USA) containing either recombinant murine leptin (Peprotech, Rocky Hill, NJ, USA) released at a dose of 20 μg/day or vehicle, as previously described21, for 8 days. Mice that did not receive STZ underwent sham surgery on Day 0, whereby a skin incision was made, subcutaneous tunneling was done, and finally, the opening was sutured. (2) In the acute leptin therapy study, STZ (180 mg per kg body weight, i.p.) was administered to aged-matched (13–36 weeks old) male LeprΔ17TB/loxTB mice19,20, which were on a mixed C57BL/6J and 129 background, and LeprΔ17TB/loxTB;ROSA26mTmG mice on Day −8. Criteria for diabetes was the same as for the prolonged leptin therapy study. On Day 0, following a 4 h morning fast, mice were injected i.p. with recombinant murine leptin (Peprotech) at a dose of 3 mg per kg body weight, as utilized by Burnett et al.22. Plasma was obtained from saphenous vein blood collected immediately before injecting leptin (0 h), as well as at 0.25 h, 0.5 h, 1 h, 2 h, 4 h, and 6 h after injecting leptin. Half-life of leptin was calculated using a one phase decay curve22 (GraphPad Prism 7), with plasma leptin concentrations from 0.5 h to 6 h after injecting leptin.

Immunofluorescence analysis and hematoxylin and eosin (H&E) staining. Methodology for these studies are found in Supplemental Material.

Plasma assays. Body weight and blood glucose measurements were obtained following a 4 h morning fast. Blood glucose was measured in samples obtained from saphenous vein blood using a One Touch Ultra Glucometer (Life Scan, Burnaby, Canada). When blood glucose concentration exceeded the limit of detection of the glucometer (>33.3 mM), it was assigned to equal 33.3 mM. Plasma was obtained from saphenous vein blood or from cardiac blood collected following a 4 h morning fast unless stated otherwise. Plasma insulin was measured using the mouse ultrasensitive insulin ELISA from ALPCO Diagnostics (Salem, NH, USA) and plasma leptin was measured using the mouse leptin ELISA from Crystal Chem (Downers Grove, IL, USA). Plasma insulin concentrations of mice from the ATLeprEXP colony at 10 weeks of age and during glucose tolerance tests were measured using the Stellux Chemi Rodent Insulin ELISA (Alpco). The assay for plasma free fatty acids (FFAs) (Wako Chemicals, Richmond, VA, USA) and the assay for glycerol and triglycerides (TGs) (Sigma-Aldrich) were performed as previously stated23. For analyses of plasma obtained from cardiac blood in the ATLeprEXP colony, a custom-made multiplex assay for mouse leptin, insulin, and resistin (Milliplex, EMD Millipore, Billerica, MA, USA) was used according to the manufacturer’s instructions; undiluted samples from all mice were assessed for insulin and resistin, while samples were diluted 1:8 for plasma assay. The concentration of leptin receptor in plasma was determined with a mouse leptin receptor ELISA from R&D Systems (Minneapolis, MN, USA; Catalogue #DY008 and #DY497) and by following the manufacturer’s instructions.

In vivo metabolic tests. Mice were fasted for 4 h, starting in the morning, before performing i.p. or oral glucose tolerance tests (IPGTTs and OGTTs, respectively) or i.p. insulin tolerance tests (ITTs, 0.65 or 0.75U human synthetic insulin (Novolin ge Toronto, Novo Nordisk, Mississauga, Canada) per kg body weight)12,24. The higher insulin dose was used in older mice due to age-associated insulin resistance. For mice in the AdipoqKO colony, which contains mice with Lepr knockdown, the dose for IPGTTs and OGTTs was 1.5 g glucose per kg body weight. For mice in the ATLeprEXP colony, which contains mice with Lepr reconstitution, the dose for IPGTTs and OGTTs was 1 g glucose per kg body weight. Doses for IPGTTs, OGTTs, and ITTs were normalized for body weight and consequently, in studies where body weight differed between groups, the absolute doses differed between groups. Area under the curve (AUC) was calculated using GraphPad Prism 7; baseline was 0 mM for glucose and 0 ng/ml for insulin. For fasting-refeeding experiments, mice were fasted starting at 5:30–6:30 pm and blood was obtained at 0 h (immediately before start of fast), 4, 12, and 16 h of fasting. Food was then given to
the mice and blood samples were obtained 1 h and 2 h following the refeeding. Blood obtained from the saphenous vein was used for all in vivo metabolic tests.

**Metabolic cages and DEXA analysis.** In order to assess food intake, activity, energy expenditure, and substrate utilization, mice were placed in Phenomaster metabolic cages (TSE Systems) at 23 °C, as previously described. Dual energy x-ray absorbance (DEXA) measurements (Lunar PIXImus 2.0 Densitometer, Inside Outside Sales, Madison, WI, USA) were obtained in the non-fasting state. For mice from the AdipoqKO colony, DEXA analysis was carried out following metabolic cage studies, on the same day.

**Western blot analysis and hepatic glycogen content.** Female AdipoqCre<sup>Lepr<sup>lox/lox</sup></sup> and AdipoqCre<sup>Lepr<sup>fl/fl</sup></sup> mice were fasted overnight for 16 h, anesthetized with isoflurane, and injected with either 200 μl of 5% body weight insulin (Novolin Ge Toronto) or vehicle via the portal vein. Vehicle consisted of 0.1% fatty acid-free bovine serum albumin (BSA; Equitech-Bio Inc, Kerrville, TX, USA), prepared by dissolving BSA in 0.9% NaCl. One minute following injection, livers were freeze-clamped and stored at −80 °C. Approximately 50 mg of each liver sample was homogenized with the following buffer: 1X RIPA lysis buffer (EMD Millipore, Burlington, MA, USA). Complete mini EDTA-free protease inhibitor cocktail (Roche, Basel, Switzerland), 1 μg/ml pepstatin A (Sigma), 20 μM sodium fluoride (Sigma), 2 mM sodium pyrophosphate (Sigma), and 25 mM β-glycerophosphate (Sigma). Samples were rocked at 4 °C for 40 min, centrifuged at 12,000 g for 4 °C for 10 min, and lastly, the supernatant of each sample was collected and frozen at −80 °C. The Bradford method was used to measure protein concentration (Bio-Rad, Hercules, CA, USA). Samples were mixed in a 1:1 ratio with 2X Laemmli buffer containing 10% β-mercaptoethanol and boiled for 5 min before electrophoretic separation. After protein transfer to polyvinylidene difluoride membranes, membranes were blocked ( Odyssey Classic Infrared Imaging System, Li-Cor). Afterwards, membranes were stripped (NewBlot PVDF stripping buffer, Li-Cor), blocked, and incubated overnight at 4 °C with phospho-Akt (Ser473) antibody (Cell Signaling Technology, Danvers, MA, USA; Catalogue #4060, 1:2000 dilution), Tris-buffered saline with Tween (TBST) was used to wash the membranes and the ensuing incubation with secondary antibody (IRDye 800CW goat anti-rabbit IgG, Li-Cor; Catalogue #925-32211, 1:10000 dilution) lasted 1 h at room temperature. Membranes were washed before imaging (Odyssey Classic Infrared Imaging System, Li-Cor). Afterwards, membranes were stripped (NewBlot PVDF stripping buffer, Li-Cor), blocked, and incubated overnight at 4 °C with an Akt antibody that detects total Akt protein (Cell Signaling Technology; Catalogue #2920, 1:2000 dilution). The rest of the steps taken for imaging have been described above, except the secondary antibody was IRDye 680RD goat anti-mouse IgG (Li-Cor; Catalogue #925-68070, 1:10000 dilution). ImageJ (National Institutes of Health, Bethesda, MD, USA) was used to quantify bands; quantification of bands for insulin- or vehicle-injected mice were normalized to bands of AdipoqCre<sup>Lepr<sup>fl/fl</sup></sup> mice that received the same type of injection. Liver samples from these mice were also used to determine hepatic glycogen content according to the manufacturer’s instructions (BioVision, Milpitas, CA, USA).

**Endpoint PCR and isolation of adipocytes from WAT.** AdipoqCre<sup>Lepr<sup>lox/lox</sup></sup> and AdipoqCre<sup>Lepr<sup>fl/fl</sup></sup> mice were euthanized and tissues were immediately collected, placed in liquid nitrogen, and stored at −80 °C. Subcutaneous WAT (scWAT) was obtained from the inguinal region and brown adipose tissue (BAT) was obtained from the interscapular region. Subsequently, DNA was extracted from WAT, BAT, adipocyte fractions of WAT, hypothalamus, liver, skeletal muscle (gastrocnemius), heart, and pancreas and endpoint PCR was carried out using primers previously described to determine Lepr recombination. Adipocyte fractions were isolated from WAT based on Ruan et al. Briefly, freshly obtained scWAT and perigonadal WAT (pgWAT) samples were minced in collagenase solution, which consisted of 2 mg/mL collagenase Type I ( Worthington, Lakewood, NJ, USA) dissolved in a working solution (pH = 7.4) of Krebs-Ringer-Phosphate-HEPES buffer, 200 μM adenosine (Sigma, St. Louis, MO, USA), and 2.5% bovine serum albumin (Millipore, Billerica, MA, USA). Afterwards, samples were incubated at 37 °C until most of the adipose tissue pieces were not visible, which took approximately 40 min. Digestion was stopped by washing three times with working solution. The floating cells (adipocytes) were allowed to rise to the top for 1–2 minutes and adipocytes together with the mucosa were scraped using a glass microscope slide. Collected tissues were immediately placed in liquid nitrogen, stored at −80 °C, and subsequently DNA was extracted.

**qPCR.** We developed an assay to assess the extent of excision of Lepr<sup>fl/fl</sup>, that is, Lepr knockdown in DNA isolated from pgWAT, BAT, and gastrointestinal tissues (AdipoqCre<sup>Lepr<sup>lox/lox</sup></sup> vs. AdipoqCre<sup>Lepr<sup>fl/fl</sup></sup> mice). The intron 17 probe is an internal reference gene and the intron 16 probe anneals to the intact Lepr gene, but not the excised Lepr gene. The LeprInT17-FAM probe is 5′ 3′-FAM-TAGGGCGGA/GATCTCACACATACCAGATCC3 and the LeprInT16-HEX probe is 5′ 3′-5HEX/AGGAACTTCG/ZEN/TGAACACGCAAATGT/3IABkFQ and the LeprInt17 probe is 5′ 3′-GCCACCACACTTAGCTCCAATA3. Primers used are as follows: LeprInT17-F, 5′-CTTTCCAGATATGCTGTAGA3; LeprInT17-R, 5′-GCACCACTTACTGCTCAAAT3; LeprInT16-F, 5′-GATCTCACACATACAGATC3; LeprInT16-R, 5′-ATTTGATTCCACAAAGTGTTCC3. A master mix was created using SsoAdvanced universal probes supermix (Bio-Rad). The Pfaffl equation, which takes into account the different efficiencies of the two primer sets, was used to calculate the extent of recombination. Another assay was created to determine the extent of excision of Lepr<sup>lox/lox</sup>, that is, Lepr reconstitution in DNA isolated from pgWAT, scWAT, and BAT of AdipoqCre<sup>Lepr<sup>lox/lox</sup></sup> mice. Two primer sets were used. The first primer set quantifies Lepr<sup>fl/fl</sup> in a sample and is as follows: 5′-GGAAGATGCTGACTCTAGATAATGAG3 and 5′-CCAACATTGCATTCTATTTATGTTTCGAG3. The second primer set quantifies the recombined Lepr<sup>lox/lox</sup> allele.
Mann-Whitney differed over time for each parameter (GraphPad Prism 7). The with Tukey’s post-hoc also performed. When comparing a parameter among more than 2 groups at a given timepoint, one-way ANOVA correlation, or Spearman correlation if large deviations from normality existed in one or both genotypes, were U from normality in one or both genotypes, in which case the Mann-Whitney genotypes were performed using unpaired t-tests, unless distributions of a given parameter deviated substantially testing, as appropriate. Comparisons between two ANOVA with Tukey’s post-hoc testing, as appropriate. Comparisons between two groups at a given timepoint, one-way ANOVA with Tukey’s post-hoc test was used. Repeated measures two-way ANOVA, with group as the between-subject variable and time as the within-subject variable, was performed with our research question being whether groups differed over time for each parameter (GraphPad Prism 7). The post-hoc analysis of repeated measures two-way ANOVA was done when the main effect of group and/or an interaction was statistically significant and on each such occasion, groups were compared at each timepoint with either Bonferroni (2 groups) or Tukey (more than 2 groups) adjustment for multiple comparisons. Where repeated measures two-way ANOVA could not be done due to missing data points, an alternative analysis was done using R software and is described in Supplementary Information.

Results

Lepr recombination. Lepr knockdown was observed in WAT, adipocytes isolated from WAT, and BAT of AdipoqCre Lep"flox/flox mice, aged 20–28 weeks, were euthanized after a ~4–8 h fast. The ex vivo lipolysis assay protocol was performed as described by Sakaguchi et al., except that ~20–35 mg of each adipose tissue depot were used and the dose of isoprenaline (Sigma) was 0.1 μM. The FFA and glycerol concentrations in the media were divided by the tissue weight. Lepr in these media was assayed with a mouse leptin ELISA (Crystal Chem) and the results were divided by tissue weight. We have previously described the hepatic lipid extraction protocol in Huynh et al. lipid content was divided by weight of liver tissue sample.

Ex vivo lipolysis assay, ex vivo leptin secretion, and hepatic lipid extraction. For the ex vivo lipolysis assay, AdipoqCre Lep"flox/flox and AdipoqCre Lep"flox/flox mice, aged 20–28 weeks, were euthanized after a ~4–8 h fast. The ex vivo lipolysis assay protocol was performed as described by Sakaguchi et al., except that ~20–35 mg of each adipose tissue depot were used and the dose of isoprenaline (Sigma) was 0.1 μM. The FFA and glycerol concentrations in the media were divided by the tissue weight. Lepr in these media was assayed with a mouse leptin ELISA (Crystal Chem) and the results were divided by tissue weight. We have previously described the hepatic lipid extraction protocol in Huynh et al.; lipid content was divided by weight of liver tissue sample.

Statistical analyses. Data are presented as mean ± SEM. Statistical analyses consisted of unpaired t-test (or Mann-Whitney U test), Pearson correlation (or Spearman correlation), one-way ANOVA with Tukey’s post-hoc test, or repeated measures two-way ANOVA with post-hoc testing, as appropriate. Comparisons between two genotypes were performed using unpaired t-tests, unless distributions of a given parameter deviated substantially from normality in one or both genotypes, in which case the Mann-Whitney U test was used (SPSS 25). Pearson correlation, or Spearman correlation if large deviations from normality existed in one or both genotypes, were also performed. When comparing a parameter among more than 2 groups at a given timepoint, one-way ANOVA with Tukey’s post-hoc test was used. Repeated measures two-way ANOVA, with group as the between-subject variable and time as the within-subject variable, was performed with our research question being whether groups differed over time for each parameter (GraphPad Prism 7). The post-hoc analysis of repeated measures two-way ANOVA was done when the main effect of group and/or an interaction was statistically significant and on each such occasion, groups were compared at each timepoint with either Bonferroni (2 groups) or Tukey (more than 2 groups) adjustment for multiple comparisons. Where repeated measures two-way ANOVA could not be done due to missing data points, an alternative analysis was done using R software and is described in Supplementary Information.

ANOVA, with body weight as co-variate, was performed for heat and food intake (SPSS 25). Significance was achieved when p < 0.05.

Mice with adipose tissue-specific Lepr knockdown. Body weight was lower in AdipoqCre Lep"flox/flox (knockdown) vs. AdipoqCre Lep"flox/flox (Flx control) male mice starting at 18 weeks of age and lower in female knockdown mice vs. female Flx controls at 24 and 26 weeks of age (p < 0.05; Fig. 2A,C). Blood glucose concentrations did not differ between genotypes at different ages for each sex (Fig. 2B,D). These mice subsequently underwent DEXA and metabolic cage assessments. In male knockdown mice, lean tissue mass, fat tissue mass, sum of lean and fat tissue, and percent fat were significantly lower compared to controls (p < 0.05; Fig. 2E–H). In females, differences in these parameters only reached statistical significance for percent fat, which was lower in the knockdown mice (p < 0.05). Respiratory exchange ratio (RER), heat, activity, and food intake were similar between genotypes, for each sex (Fig. 2I–N).

Various plasma analyses of these mice were measured at 6 and 16 weeks of age (Fig. 3). At 6 weeks of age, plasma leptin levels did not significantly differ between male knockdown and control mice, but female...
Figure 1. (A–E) Endpoint PCR indicating the location on the gel of Lepr\textsuperscript{fllox} (1369 bp) and the allele resulting from Cre-induced excision, Lepr\textsuperscript{Δ17} (952 bp), in white adipose tissue (WAT), brown adipose tissue (BAT), adipocytes isolated from WAT, and non-adipose tissues in male and female AdipoqCre\textsuperscript{+} Lepr\textsuperscript{fllox/flox} (+) and AdipoqCre\textsuperscript{−} Lepr\textsuperscript{fllox/flox} (−) mice. Results in (A–E) are from 6 separate gels and the full-length blots/gels are presented in Supplementary Fig. S1. In A, results are from one single gel; in B, results are from the second gel that has been cropped and the white space indicates cropping; in C, results are from the third gel; in D and E, results are from the fourth gel that has been cropped and the white space indicates cropping, and from the fifth and sixth gels. (F–I) Extent of Lepr recombination in AdipoqCre\textsuperscript{+} Lepr\textsuperscript{fllox/fllox} and AdipoqCre\textsuperscript{+} Lepr\textsuperscript{loxTB/loxTB} mice determined by qPCR. pg, perigonadal; sc, subcutaneous; HY, hypothalamus; L, liver; SM, skeletal muscle (gastrocnemius); HE, heart; P, pancreas; D, duodenum mucosa; J, jejunum mucosa; I, ileum mucosa; C, colon mucosa; S, stomach.
Figure 2. Body weight and blood glucose of male (A,B) and female (C,D) AdipoqCreLepr^floxflox and AdipoqCreLepr^lox/lox mice after a 4h fast. Body composition (E–H) and metabolic cage parameters (I–N) of AdipoqCreLepr^floxflox and AdipoqCreLepr^lox/lox mice; male mice were 28–30 weeks old and female mice were 32 weeks old. For males, n = 15 for AdipoqCreLepr^floxflox and n = 13 for AdipoqCreLepr^lox/lox mice. For females, n = 11 for AdipoqCreLepr^floxflox and n = 13 for AdipoqCreLepr^lox/lox mice. In (A–D), repeated measures two-way ANOVA with Bonferroni post-hoc test was performed. For heat and food intake, an ANCOVA, using the non-fasting body weight obtained on the day metabolic cage experiments were started as co-variate, was performed and estimated marginal means ± SEM are presented. For other parameters, an unpaired t-test was performed, except for Females in (E,F,H), where the Mann-Whitney U test was used. *p < 0.05 vs. AdipoqCreLepr^floxflox of same sex. In (D), there is a main effect of time (p < 0.05). L, lean; F, fat.
knockdown mice had lower leptin concentrations than female controls (Fig. 3A,B). At 16 weeks of age, although plasma leptin concentrations did not differ between knockdown and control mice, the relationship between body weight and plasma leptin was weaker in knockdown mice than controls. Among males, Spearman correlation ($\rho$) was 0.854 ($p < 0.05$) for controls and 0.625 ($p < 0.05$) for knockdown mice. Among females, Spearman correlation was 0.827 ($p < 0.05$) for controls, whereas $\rho$ was not significant in knockdown mice. At 6 weeks of age, plasma insulin levels did not significantly differ between genotypes in males, but female knockdown mice had lower insulin concentrations than controls. At 16 weeks of age, insulin was significantly lower in knockdown mice vs. control mice in males and females (Fig. 3C,D). Plasma FFAs in males and females, and plasma glycerol and TGs in males were not significantly different between genotypes (Fig. 3E–H).

Glucose tolerance assessed by IPGTT was similar between male knockdown mice and controls (Fig. 4A). At 16 weeks of age, the blood glucose excursion during the OGTT was similar between male knockdown and control mice, but the knockdown mice had a lower insulin response ($p < 0.05$ at 7 and 15 min; Fig. 4B). Similar results were obtained in IPGTTs and OGTTs for females (Fig. 4C,D). Neither male nor female knockdown mice had statistically significant differences in insulin sensitivity, assessed by ITT, compared to their controls at different ages (Fig. 4E–H). Moreover, insulin was injected via the portal vein to assess phosphorylated Akt in the liver, a marker of hepatic insulin sensitivity. The ratio of insulin-stimulated phospho-Akt (Ser473) to total Akt in the liver was similar between Lepr knockdown and Flox control mice (Fig. 5A). The ratio was also similar between genotypes in vehicle-injected mice (Fig. 5A). During fasting, hepatic glycogen content...
Figure 4. Metabolic tests in male and female AdipoqCre^{-}Lepr^fl/fl and AdipoqCre^{+}Lepr^fl/fl. (A) IPGTT (1.5 g/kg) in males; n = 10 AdipoqCre^{-}Lepr^fl/fl and n = 14 AdipoqCre^{+}Lepr^fl/fl. (B) Glucose (left) and insulin (right) during OGTT (1.5 g/kg) in males; n = 10 AdipoqCre^{-}Lepr^fl/fl and n = 14 AdipoqCre^{+}Lepr^fl/fl. (C) IPGTT (1.5 g/kg) in females; n = 14 AdipoqCre^{-}Lepr^fl/fl and n = 19 AdipoqCre^{+}Lepr^fl/fl. (D) Glucose (left) and insulin (right) during OGTT (1.5 g/kg) in females; n = 6 AdipoqCre^{-}Lepr^fl/fl and n = 7 AdipoqCre^{+}Lepr^fl/fl. (E) ITT (0.65 U/kg) in males; n = 10 AdipoqCre^{-}Lepr^fl/fl and n = 14 AdipoqCre^{+}Lepr^fl/fl. (F) ITT (0.65 U/kg) in females; n = 14 AdipoqCre^{-}Lepr^fl/fl and n = 18 AdipoqCre^{+}Lepr^fl/fl. (G) ITT (0.75 U/kg) in males; n = 10 AdipoqCre^{-}Lepr^fl/fl and n = 13 AdipoqCre^{+}Lepr^fl/fl. (H) ITT (0.75 U/kg) in females; n = 6 AdipoqCre^{-}Lepr^fl/fl and n = 9 AdipoqCre^{+}Lepr^fl/fl. Repeated measures two-way ANOVA with Bonferroni post-hoc test was performed for parameters measured over time. An unpaired t-test was performed for each area under the curve (AUC) calculation, except for glucose concentration in (D), where Mann-Whitney U test was used. *p < 0.05 vs. AdipoqCre^{-}Lepr^fl/fl. In (A–H), there is a main effect of time (p < 0.05).
Figure 5. Effect of intraportal insulin or vehicle on the ratio of phospho-Akt (Ser473) to total Akt in the liver of female AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup> mice (A). For vehicle injection, n = 3 per genotype, and for insulin injection, n = 4 per genotype. For each type of injection, quantification was normalized to bands of AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> mice. Results for each protein are from 2 separate blots and Supplementary Fig. S4 contains the full-length blots. Representative blots for each protein are from the same samples in the first blot. Blood glucose and plasma FFA concentrations during fasting-refeeding experiments at different ages in male and female AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup> mice (B–I). For younger males, n = 3 per genotype. For older males, n = 4 for AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and n = 5 for AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup>. For younger females, n = 4 for AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and n = 6 for AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup>. For older females, n = 6 for AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and n = 9 for AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup>. FFA and glycerol concentrations in media collected in ex vivo lipolysis assay under basal (J–M) and isoprenaline-stimulated conditions (N,O). In (J,K), n = 6 for AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and n = 7 for AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup>. In (L,M), n = 8 for AdipoqCre<sup>−</sup>Lepr<sup>lox/lox</sup> and n = 9 for AdipoqCre<sup>+</sup>Lepr<sup>lox/lox</sup>.
In (N,O), n = 6 per genotype for males and for females, n = 6 for AdipoqCre-LeprloxTB/flox and n = 8 for AdipoqCre-LeprloxTBfloxFlox in (A), the Mann-Whitney U test was performed for vehicle injection and an unpaired t-test was performed for insulin injection. In (B–I), repeated measures two-way ANOVA with Bonferroni post-hoc test was performed and there is a main effect of time (p < 0.05). In (J–O), an unpaired t-test was used to compare the two genotypes for each tissue and parameter. *p < 0.05 vs. AdipoqCre-LeprloxTBfloxFlox. F, fast; R, refeed; Iso, isoprenaline; pgW AT, perigonadal white adipose tissue; scW AT, subcutaneous white adipose tissue.

was similar between Lepr knockdown and Flox control mice (2.29 ± 0.64 and 2.82 ± 0.41 mg per g liver, respectively; not statistically significant).

We performed fasting-refeeding experiments in both sexes at different ages (Fig. 5B–I) and found that blood glucose did not differ between knockdown and control mice, but younger female knockdown mice had higher plasma FFA concentrations after a prolonged fast (16 h) compared to controls (Fig. 5G). The rate of lipolysis, which is based on the amount of FFA and glycerol released ex vivo by different adipose tissue depots, was not altered in male and female knockdown mice compared to their controls (Fig. 5J–M). The response of pgWAT to isoprenaline, a β3-receptor adrenergic receptor agonist, was also comparable between knockdown mice and controls in both sexes (Fig. 5N,O). Leptin secretion from pgWAT and scWAT samples was similar between Lepr knockdown and Flox controls, among males and females (Supplementary Fig. S5).

To assess the contribution of adipose tissue leptin signalling to the glucose-lowering effect of exogenous leptin during insulin deficiency, male mice were rendered diabetic with STZ and subsequently treated with murine leptin delivered via mini-osmotic pumps for 8 days. While all mice had similar hyperglycemia following STZ injections, leptin therapy induced more rapid normalization in blood glucose levels in the knockdown mice (Supplementary Fig. S6). STZ + leptin groups had a progressive decrease in body weight vs. Sham controls (Supplementary Fig. S6). Mice became hypoleptinemic following STZ administration and leptin therapy increased plasma leptin levels to a greater extent in knockdown vs. Flox controls (p < 0.05, Supplementary Fig. S6). The elevated plasma leptin concentrations in the Lepr knockdown STZ + leptin groups vs. Flox control STZ + leptin group were associated with increased concentrations of leptin receptor in plasma (p < 0.05, Supplementary Fig. S6). STZ-injected groups had similar insulin deficiency (Supplementary Fig. S6). To test if the duration of leptin therapy affected the difference in plasma leptin concentrations between genotypes among STZ-injected mice, we treated diabetic mice with a single injection of murine leptin and monitored their blood glucose and plasma leptin over 6 hours, a protocol similar to that used by Burnett et al. in non-diabetic mice22. The single leptin injection did not lower blood glucose differentially in diabetic Lepr knockdown vs. Flox control mice and plasma leptin levels were similar between genotypes at all timepoints (Supplementary Fig. S7). The half-life of leptin, with 30 min post-injection as the starting time for the calculation (time of highest average leptin concentration), was 53 ± 6 min for Lepr knockdown and 57 ± 12 min for Flox control mice, a difference that was not statistically significant.

**Mice with adipose tissue-specific Lepr reconstitution.** As an alternative approach to understand the consequences of leptin signalling in adipose tissue, we investigated the effect of reconstituting Lepr expression specifically in adipose tissues of LeprloxTB/loxTB mice, in which global Lepr expression is inhibited due to a transcriptional block. These mice have a similar phenotype to db/db mice including obesity and hyperglycemia23. Male LeprloxTB/loxTB mice grow faster and reach a higher maximum body weight compared to controls (Fig. 6A and Supplementary Fig. S8), and interestingly, mice with Lepr expression selectively in adipose tissues (AdipoqCre-LeprloxTBfloxflox) have a higher maximum body weight than mice lacking Lepr expression globally (AdipoqCre-LeprloxTBfloxflox) (Fig. 6A). No differences in body composition, assessed by DEXA, were detected in male AdipoqCre-LeprloxTBfloxflox vs. AdipoqCre-LeprloxTBfloxflox mice, but as expected LeprloxTB/loxTB mice had higher body fat content compared to male controls (AdipoqCre-Lepr+/- and AdipoqCre-Lepr+/+ mice) (Table 1). Although initially hyperglycemic, male AdipoqCre-LeprloxTBfloxflox mice had similar blood glucose levels to controls starting at 10 weeks of age, but this normalization was delayed in male AdipoqCre-LeprloxTBfloxflox mice (Fig. 6B). Although LeprloxTB/loxTB females were obese compared to controls, the maximum body weight of female AdipoqCre-LeprloxTBfloxflox mice did not differ from that of AdipoqCre-LeprloxTBfloxflox mice (Fig. 6C and Supplementary Fig. S8). Body composition was similar between female AdipoqCre-LeprloxTBfloxflox and AdipoqCre-LeprloxTBfloxflox mice (Table 1). At 6 weeks of age, LeprloxTB/loxTB females were hyperglycemic (Fig. 6D). AdipoqCre-LeprloxTBfloxflox female mice reached normal glucose levels by 10 weeks but both AdipoqCre-LeprloxTBfloxflox and AdipoqCre-LeprloxTBfloxflox female mice had slightly elevated blood glucose (p < 0.05 vs. controls) at 14 and 16 weeks of age. The improvement in glycemia among AdipoqCre-LeprloxTBfloxflox mice compared to AdipoqCre-LeprloxTBfloxflox mice at 10 weeks of age was not associated with statistically significant differences in plasma insulin concentrations between these two genotypes; they were both hyperinsulinemic (Fig. 6E). At an older age, mice with adipose tissue Lepr reconstitution had higher plasma insulin concentrations than mice with global inhibition of Lepr during the OGTT, despite having similar glucose concentrations throughout the test (Supplementary Fig. S9). These results are complementary to those obtained during the OGTT in the adipose tissue Lepr knockdown colony. Mice with adipose tissue Lepr reconstitution also had higher blood glucose at 120 min of the IPGTT compared to mice with global inhibition of Lepr and differences in the plasma insulin profile throughout IPGTT were not statistically significant (Supplementary Fig. S9).

Lepr knockdown mice and Flox controls on HFD were similarly heavier relative to mice on LFD (Fig. 6F) and transient hyperglycemia was observed in both genotypes (Fig. 6G). Interestingly, the magnitude of reduced weight gain in male mice with disruption of Leprb in adipose tissue was similar to that of the increased weight gain in male mice with reconstitution of Leprb in adipose tissue.
Figure 6. Body weight, blood glucose, and plasma insulin concentrations at 10 weeks old of male and female mice from the ATLeprEXP colony (A–E) and of male mice from the AdipoqKO colony on a high (HFD) or low fat diet (LFD) (F,G), obtained after a 4 h fast. In (A,B,E), n = 10 for AdipoqCre+ LeprloxTB/loxTB, except at 16 weeks of age, when n = 9; n = 11 for AdipoqCre+ LeprloxTB/loxTB, except at 16 weeks of age, when n = 9; n = 16 for AdipoqCre+ LeprloxTB/loxTB; n = 9 for AdipoqCre+ LeprloxTB/loxTB. In (C,D,E), n = 8 for AdipoqCre+ LeprloxTB/loxTB; n = 8 for AdipoqCre+ LeprloxTB/loxTB; n = 7 for AdipoqCre+ LeprloxTB/loxTB. In (F,G), Pre-diet refers to the day before the start of diet; for AdipoqCre+ LeprloxTB/loxTB, n = 10 on HFD and n = 7 on LFD, and for AdipoqCre+ LeprloxTB/loxTB, n = 8 on HFD and n = 4 on LFD. In (H), RT-qPCR for Fasn, Pnpla2, and Scl2a4 was done in perigonadal white adipose tissue and for Ucp1 in brown adipose tissue; all samples were from male mice, 17–20 weeks old, from the ATLeprEXP colony. In (H), n = 8 for AdipoqCre+ LeprloxTB/loxTB, except for Ucp1 where n = 9; n = 10 for AdipoqCre+ LeprloxTB/loxTB, except for Ucp1 where n = 11; n = 14 for AdipoqCre+ LeprloxTB/loxTB, except for Ucp1 where n = 15; and n = 7 for AdipoqCre+ LeprloxTB/loxTB, except for Ucp1 where n = 8. For body weights of males and females from ATLeprEXP colony (A,C), only statistical significance of maximum body weight (asymptote)}
is indicated in the figure. In (F,G), repeated measures two-way ANOVA with Tukey post-hoc test was performed and there is a main effect of time (p < 0.05). In (E), for each sex, and in (H), for each parameter, one-way ANOVA with Tukey post-hoc was carried out. *p < 0.05, AdipoqCre + LeprloxTB/loxTB vs AdipoqCre − LeprloxTB/loxTB; †p < 0.05, AdipoqCre + LeprloxTB/loxTB and AdipoqCre − LeprloxTB/loxTB vs AdipoqCre + LeprμT/μT and AdipoqCre − LeprμT/μT; ‡p < 0.05, AdipoqCre + LeprloxTB/loxTB vs AdipoqCre + LeprμT/μT and AdipoqCre − LeprμT/μT; ¶p < 0.05, AdipoqCre + LeprloxTB/loxTB vs AdipoqCre − LeprμT/μT; ††p < 0.05, AdipoqCre + LeprloxTB/loxTB vs AdipoqCre − LeprμT/μT; ‖p < 0.05, AdipoqCre + LeprloxTB/loxTB vs AdipoqCre − LeprμT/μT; ¶¶p < 0.05, HFD groups vs. LFD groups.

Expression of enzymes involved in lipid synthesis (Fasn) and lipolysis (Pnpla2), and of the glucose transporter GLUT4 (Slc2a4) was also assessed in pgWAT of male mice from the ATLeprEXP colony (Fig. 6G). Their expression was decreased in LeprloxTB/loxTB mice compared to controls (p < 0.05), but statistically significant differences were not detected between AdipoqCre + LeprloxTB/loxTB and AdipoqCre − LeprloxTB/loxTB mice. The expression of Ucp1 in BAT of males was similar among the 4 genotypes (Fig. 6H).

Levels of total triglycerides and cholesterol in the liver were higher in male LeprloxTB/loxTB mice compared to controls (p < 0.05), but were similar between AdipoqCre + LeprloxTB/loxTB and AdipoqCre − LeprloxTB/loxTB mice (Fig. 7A,B). LeprloxTB/loxTB male and female mice were hyperleptinemic (p < 0.05 vs. controls; Fig. 7C) and hyperinsulinemic (for males, p < 0.05 vs. controls; for females, p < 0.05 for AdipoqCre + LeprloxTB/loxTB vs. controls) (Fig. 7D). Plasma resistin was similar between groups (Fig. 7E). At 12 weeks of age, plasma FFA concentrations were not significantly different among males, but female LeprloxTB/loxTB mice had higher FFA concentrations compared to controls (p < 0.05; Fig. 7F). Plasma glycerol concentrations were higher in male AdipoqCre + LeprloxTB/loxTB mice compared to AdipoqCre + LeprμT/μT mice (p < 0.05), but no differences were found among female mice (Fig. 7G). Plasma triglycerides were similar among the 4 genotypes, regardless of sex (Fig. 7H). At a later age, in males, plasma FFA and 4 triglycerides were similar among the 4 genotypes, but plasma glycerol levels were 87% higher in male AdipoqCre + LeprloxTB/loxTB vs. AdipoqCre + LeprμT/μT mice (p < 0.05 for AdipoqCre + LeprloxTB/loxTB vs. controls; Fig. 7I–K).

Discussion
We found that Cre-mediated excision of Lepr in adipose tissues modestly decreased body weight and diminished the insulin response during an oral glucose challenge in male and female mice. In the setting of insulin-deficiency, male mice with Lepr knockdown also had an accelerated glucose-lowering response to prolonged leptin administration compared to controls that was associated with increased concentrations of plasma leptin and leptin receptor. The half-life of leptin, calculated after a single injection of leptin, was similar between diabetic Lepr knockdown and Flox control mice. In complementary studies, reconstitution of Lepr selectively in adipose tissues of LeprloxTB/loxTB mice, which are obese and hyperglycemic, resulted in lower concentrations of blood glucose in both male and female mice at younger ages, and a greater maximum body weight was observed in male mice. Mice with reconstitution of Lepr in adipose tissues were also characterized by elevated plasma insulin concentrations during the OGTT and increased blood glucose concentrations during the IPGTT compared to mice with global inhibition of Lepr expression.

Cre-mediated recombination of Lepr was similar between males and females and in the Lepr knockdown and reconstitution studies. We observed relatively modest recombination using AdipoqCre mice, unlike most other reports. Cre efficiency can vary depending upon the floxed line, but our results were comparable between LeprloxTB/loxTB and LeprloxTBloxTB and mtmG reporter line. Cre is expected to be expressed only in mature adipocytes and adipose tissue also contains immature adipocytes so perhaps the ratio of these cells, and ultimately the activity of the adiponectin promoter, is different in our lines compared to those that others have used.

Adiponectin and Lepr are both expressed in intestinal epithelial cell-specific Leprb knockout mice and achieved robust recombination in intestinal mucosa. A key finding in the paper was that following 12 weeks on high fat diet, intestinal epithelial cell-specific Leprb knockout mice were ~5 g lighter compared to Flox controls as a result of impaired nutrient absorption. Although after 12 weeks on high fat diet our mice with adipose tissue Lepr knockdown were ~5 g lighter vs Flox controls, we did not find evidence of Lepr recombination in intestinal mucosa. Therefore, we do not believe that Lepr recombination in the intestines is a major cause of the phenotype we observed. However, plasma leptin was lower in female adipose tissue Lepr knockdown mice at 6 weeks old, and at 16 weeks old there was a trend, albeit not statistically significant, for plasma leptin to be lower in male and female adipose tissue Lepr knockdown mice compared to controls. Therefore, it is possible that small changes in circulating leptin in mice with adipose tissue Lepr knockdown impaired absorption of nutrients, thereby reducing weight gain.

The results we obtained by knockdown of Lepr in adipose tissues contrast with results using antisense RNA to downregulate expression of leptin receptors in WAT. In the study by Huan et al., male and female mice were characterized by obesity, glucose intolerance, and insulin resistance, although obesity and glucose intolerance appeared at a later age in females. We suggest four possible explanations for the different findings. First, we targeted Leprb while Huan et al. targeted all isoforms of Lepr, although downregulation was only reported for the short isoforms. The short isoforms predominate in WAT, but Leprb is considered to be responsible for the majority of the effects of leptin. Second, we targeted BAT and WAT, whereas Huan et al. only targeted WAT. Ultimately, the phenotype of our mice is the net effect of alterations in Lepr signalling in BAT and WAT. However, we did not detect differences in the histology of BAT or in the ex vivo release of FFA and glycerol from BAT in Lepr knockdown mice vs. controls. Moreover, Ucp1 expression was not altered in BAT of males with Lepr reconstitution. Third, while we used Cre-lox methodology, Huan et al. downregulated the expression of Lepr isoforms using
antisense RNA expressed postnatally under the control of the PEPCK promoter. This knockdown approach has been suggested to induce obesity, regardless of the target RNA14. Finally, we can not discount the possibility that developmental effects in either model contribute to the observed differences.

Circulating hyperleptinemia was associated with the obese phenotype obtained by downregulation of leptin receptors in WAT using antisense RNA13. Hyperleptinemia was also observed upon reduction of Leprb in multiple peripheral tissues using a tamoxifen-inducible model, despite the mice being of normal body weight11. The differences in circulating levels of leptin in our mice with adipose tissue-specific knockdown of Lepr compared to control mice were not statistically significant, except in females at 6 weeks of age, when it was lower in Lepr knockout mice. Moreover, using an ex vivo assay, leptin secretion by WAT samples, which was in the range of what has been reported11,37, was similar between Lepr knockdown and Flox control mice, in males and females. These findings do not support a negative feedback loop of leptin secretion via Leprb at the level of adipose tissues. However, we achieved modest recombination in adipose tissues and the short isoforms of Lepr may modulate these findings partially acts to trigger its glucose-lowering effects in insulin-deficient diabetes, it remains unclear if the CNS pathways involved are the same as for centrally administered leptin44–48. Perry et al.49 demonstrated that peripheral

Table 1. Body composition in mice from ATLexpERX colony. For males, n = 8 for AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup>, n = 10 for AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup>, n = 16 for AdipoqCre<sup>−</sup> Lepr<sup>+/+</sup>, n = 7 for AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup>. For females, n = 15 for AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup>, n = 8 for AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup>, n = 8 for AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup>, n = 7 for AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup>, <sup>†</sup>p < 0.05, AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> vs AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> and AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> vs AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup>; <sup>‡</sup>p < 0.05, AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> vs AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> and AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> vs AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup>. For each parameter within each sex, one-way ANOVA followed by Tukey's test was performed. Age = 17–19 weeks of age.

### Table 1. Body composition in mice from ATLEXP colony.

| Sex | Genotype          | Lean mass (g) | Fat mass (g) | Sum of lean and fat mass (g) | Percent fat |
|-----|-------------------|---------------|--------------|----------------------------|-------------|
| M   | AdipoqCre<sup>+</sup> Lepr<sup>loxTB/loxTB</sup> | 25.9 ± 0.6 | 36.9 ± 0.8 | 62.8 ± 1.0 | 58.8 ± 0.7 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> | 25.2 ± 0.5 | 33.7 ± 1.4 | 58.7 ± 1.7 | 57.0 ± 1.1 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> | 23.2 ± 0.4 | 8.3 ± 0.7 | 31.5 ± 0.9 | 25.9 ± 1.4 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> | 24.1 ± 0.7 | 9.8 ± 1.2 | 33.8 ± 1.6 | 28.4 ± 2.5 |
| F   | AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> | 24.2 ± 1.4 | 39.1 ± 1.3 | 63.2 ± 2.1 | 62.0 ± 1.4 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>loxTB/loxTB</sup> | 25.3 ± 0.5 | 39.1 ± 0.7 | 64.3 ± 0.9 | 60.7 ± 0.5 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> | 18.7 ± 0.6 | 6.7 ± 1.0 | 25.4 ± 1.5 | 25.5 ± 2.4 |
|     | AdipoqCre<sup>−</sup> Lepr<sup>−/−</sup> | 19.0 ± 0.7 | 7.6 ± 0.8 | 26.7 ± 1.4 | 28.1 ± 1.8 |

Although most studies indicate that the CNS is the key tissue upon which peripherally administered leptin initially acts to trigger its glucose-lowering effects in insulin-deficient diabetes, it remains unclear if the CNS pathways involved are the same as for centrally administered leptin44–48.
Figure 7. Hepatic (A, B) and plasma (C–K) parameters of mice from the ATLeprEXP colony. Liver samples used for the analysis in (A, B) were obtained from male mice at the time of euthanasia (17–20 weeks old). In (A, B), n = 10 for AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup>, n = 9 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>, n = 16 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, and n = 8 for AdipoqCre<sup>−</sup>Lepr<sup>+/+</sup>. Plasma obtained from cardiac blood collected at the time of euthanasia (17–20 weeks old) in (C–E) and (I–K), and from saphenous vein blood at 12 weeks old (F–H). For males in (C–E) and (I–K), n = 9 for AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup>, n = 8 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>, n = 14 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, n = 7 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, except in (I–K), where n = 10 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>. For females in (C–E), n = 11 for AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup>, n = 4 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>, n = 4 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, and n = 5 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>. In (F–H), for males, n = 11 for AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup>, n = 10 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>, n = 16 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, n = 9 for AdipoqCre<sup>−</sup>Lepr<sup>+/+</sup>; for females, n = 15 for AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup>, n = 8 for AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup>, n = 8 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>, n = 7 for AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>. For each sex, one-way ANOVA followed by Tukey’s test was performed at each timepoint. *p < 0.05 vs. AdipoqCre<sup>−</sup>Lepr<sup>loxTB/loxTB</sup> and AdipoqCre<sup>−</sup>Lepr<sup>+/+</sup>. †p < 0.05 vs. AdipoqCre<sup>+</sup>Lepr<sup>loxTB/loxTB</sup> and AdipoqCre<sup>+</sup>Lepr<sup>+/+</sup>.
leptin diminishes hypothalamic-pituitary-adrenal (HPA) axis activity to reduce gluconeogenesis and glycemia, but this mechanism is specific to conditions of severe insulin deficiency because an increase in the magnitude of residual levels in insulin-deficient diabetes diminishes the ability of leptin to lower circulating glucose.52,53

Herein, prolonged (8 days) leptin therapy resulted in a faster decrease in blood glucose in diabetic mice with adipose tissue-specific Lepr knockdown vs. controls and this was associated with significantly higher leptin levels following leptin therapy. Since endogenous plasma leptin levels were similar between genotypes in mice with normal levels of insulin (ie. not injected with STZ), leptin clearance may be impaired in insulin-deficient Lepr knockdown mice upon administration of prolonged exogenous leptin. The binding of circulating leptin to its leptin receptor diminishes the clearance of leptin52 and hence, the elevated concentration of plasma leptin receptor we observed in diabetic Lepr knockdown mice may contribute to the increased circulating leptin concentration in Lepr knockdown mice following 8 days of leptin therapy. However, the differential effect of exogenous leptin on plasma leptin concentrations in diabetic Lepr knockdown and Flox controls depends on the duration of leptin therapy. Following an acute injection of leptin, STZ-injected mice of both genotypes had similar plasma leptin profiles and leptin half-lives. After the single leptin injection, blood glucose profiles were similar between the two groups and, as expected based on our previous studies, blood glucose was not normalized.34

POMC neuron-specific expression of Lepr reduces obesity in LeprloxTB/loxTB males starting at 13 weeks of age, plateauing at a ~10 g differential at 15–20 weeks of age.6 In our study, the magnitude of the difference in body weight between male mice with adipose tissue-specific Lepr reconstitution (AdipoqCre− LeprloxTB/loxTB) and loxTB control mice (AdipoqCre+ LeprloxTB/loxTB) was similar, ~6 g at 16 weeks of age, but Lepr reconstitution did not reduce obesity in our study. Our results indicate that obesity can be modestly prevented or augmented if Lepr expression is reduced or reconstituted, respectively, in adipose tissues of males. There were no differences in body weight between female mice with reconstitution of Lepr specifically in POMC neurons (POMC-Cre+ LeprloxTB/loxTB) and loxTB control mice (POMC-Cre− LeprloxTB/loxTB)8. In our study, we also did not find an effect on body weight in females since the maximal body weight of female mice with adipose tissue-specific reconstitution of Lepr did not differ from that of female loxTB control mice.

Mice with Lepr reconstitution had lower blood glucose concentrations at certain ages, compared to AdipoqCre− LeprloxTB/loxTB mice. The transient alterations in blood glucose concentration of LeprloxTB/loxTB male and female mice observed in our study are consistent with what has been reported in db/db mice, where blood glucose concentrations are inversely associated with plasma insulin concentrations.35 The improved blood glucose concentrations in mice with adipose tissue Lepr reconstitution at younger ages was not due to elevations in plasma insulin and at older ages, mice with Lepr reconstitution were mildly glucose intolerant.

High fat diet diminishes Leprb expression in VAT and overexpression of Leprb in VAT and BAT, under the control of an aP2 promoter, inhibits the body weight gain in high fat diet-fed mice.36 In contrast, we found that disruption of Lepr signalling in fat reduced body weight gain, and reconstitution of leptin signalling in fat in male mice lacking leptin signalling elsewhere increased weight gain. Although it is difficult to reconcile these results, the aP2-Cre model can promote recombination in both adipose and non-adipose tissues.6

In conclusion, our results indicate that peripheral leptin signalling in adipose tissues affects body weight regulation and glucose metabolism. Mice with adipose tissue-specific Lepr knockdown were modestly lighter, while reconstitution of Lepr in adipose tissues made male LeprloxTB/loxTB mice, which were obese, attain an even greater maximal body weight. In response to an oral glucose challenge, while mice with adipose tissue Lepr knockdown had a blunted plasma insulin profile, mice with adipose tissue Lepr reconstitution had higher plasma insulin concentrations compared to controls. Although mice with Lepr reconstitution, which were hyperleptinemic and hyperinsulinemic, had lower blood glucose compared to AdipoqCre− LeprloxTB/loxTB mice at certain younger ages, they were mildly glucose intolerant at older ages. In insulin-deficient diabetes, prolonged exogenous leptin therapy induced a faster reduction in blood glucose, possibly as a result of diminished leptin clearance. Additional research is warranted to further investigate the underlying mechanisms by which adipose tissue leptin signalling affects body weight and glucose homeostasis.

Data Availability

The datasets generated during and/or analysed during the current study are available from the corresponding author on reasonable request.

References
1. Scott, M. M., Williams, K. W., Rossi, J., Lee, C. E. & Elmquist, J. K. Leptin receptor expression in hindbrain Glp-1 neurons regulates food intake and energy balance in mice. Journal of Clinical Investigation 121, 2413–2421 (2011).
2. Dodd, G. T. et al. The thermogenic effect of leptin is dependent on a distinct population of prolactin-releasing peptide neurons in the dorsomedial hypothalamus. Cell Metab 20, 639–669 (2014).
3. Minokoshi, Y., Haque, M. S. & Shimazu, T. Microinjection of leptin into the ventromedial hypothalamus increases glucose uptake in peripheral tissues in rats. Diabetes 48, 287–291 (1999).
4. Bonzon-Kulichenko, E. et al. Central leptin regulates total ceramide content and sterol regulatory element binding protein-1C proteolytic maturation in rat white adipose tissue. Endocrinology 150, 169–178 (2009).
5. Balthasar, N. et al. Leptin receptor signaling in POMC neurons is required for normal body weight homeostasis. Neuron 42, 983–991 (2004).
6. van de Wall, E. et al. Collective and individual functions of leptin receptor modulated neurons controlling metabolism and ingestion. Endocrinology 149, 1773–1785 (2008).
7. Shi, H. et al. Sexually different actions of leptin in proopiomelanocortin neurons to regulate glucose homeostasis. Am J Physiol Endocrinol Metab 294, E630–639 (2008).
8. Berglund, E. D. et al. Direct leptin action on POMC neurons regulates glucose homeostasis and hepatic insulin sensitivity in mice. Journal of Clinical Investigation 122, 1000–1009 (2012).
9. Bates, S. H. & Myers, M. G. Jr. The role of leptin receptor signaling in feeding and neuroendocrine function. Trends Endocrinol Metab 14, 447–452 (2003).
10. Moon, H. S. et al. Leptin's role in lipodystrophic and nonlipodystrophic insulin-resistant and diabetic individuals. Endocr Rev 34, 377–412 (2013).

11. Guo, K. et al. Disruption of peripheral leptin signaling in mice results in hyperleptinemia without associated metabolic abnormalities. Endocrinology 148, 3987–3997 (2007).

12. Huynh, F. K. et al. Disruption of hepatic leptin signaling protects mice from age- and diet-related glucose intolerance. Diabetes 59, 3032–3040 (2010).

13. Huan, J. N. et al. Adipocyte-selective reduction of the leptin receptors induced by antisense RNA leads to increased adiposity, dyslipidemia, and insulin resistance. J Biol Chem 278, 45638–45650 (2003).

14. de Luca, C. et al. Complete rescue of obesity, diabetes, and infertility in db/db mice by neuron-specific LEPR-B transgenes. Journal of Clinical Investigation 115, 3484–3493 (2005).

15. Wang, M. Y., Orci, L., Ravazzola, M. & Unger, R. H. Fat storage in adipocytes requires inactivation of leptin's paracrine activity: implications for treatment of human obesity. Proc Natl Acad Sci USA 102, 18011–18016 (2005).

16. Lee, K. Y. et al. Lessons on conditional gene targeting in mouse adipose tissue. Diabetes 62, 864–874 (2013).

17. Eguchi, J. et al. Transcriptional control of adipose lipid handling by IRF4. Cell Metab 13, 249–259 (2011).

18. McMinn, J. E. et al. An allelic series for the leptin receptor gene generated by CRE and FLPe recombinase. Mamm Genome 15, 677–685 (2004).

19. Tuduri, E. et al. Partial ablation of leptin signaling in mouse pancreatic alpha-cells does not alter either glucose or lipid homeostasis. Am J Physiol Endocrinol Metab 306, E748–755 (2014).

20. Muzumdar, M. D., Tasic, B., Miyamichi, K., Li, L. & Luo, L. A global double-fluorescent Cre reporter mouse. Genesis 45, 593–605 (2007).

21. Denroche, H. C. et al. Leptin induces fasting hypoglycaemia in a mouse model of diabetes through the depletion of glycerol. Diabetologia 58, 1100–1108 (2015).

22. Burnett, L. C., Skowronska, A. A., Rausch, R., LeDuc, C. A. & Leibel, R. L. Determination of the half-life of circulating leptin in the mouse. Int J Obes (Lond) 41, 355–359 (2017).

23. Denroche, H. C. et al. Leptin therapy reverses hyperglycemia in mice with streptozotocin-induced diabetes, independent of hepatic leptin signaling. Diabetes 60, 1414–1423 (2011).

24. Ayala, I. E. et al. Standard operating procedures for describing and performing metabolic tests of glucose homeostasis in mice. Dis Model Mech 3, 525–534 (2010).

25. Lee, K. T., Karunakaran, S., Ho, M. M. & Clee, S. M. PWD/Ph and WSB/Ej mice are resistant to diet-induced obesity but have normal insulin secretion. Endocrinology 152, 3005–3017 (2011).

26. Ramey, G., Fayé, A., Durel, B. & Viollet, B. & Vaulont, S. Iron overload in Hepc1−/− mice is not impairing glucose homeostasis. PLoS ONE 8, e59756 (2013).

27. Sebastian, D. et al. Mitofusin 2 (Mfn2) links mitochondrial and endoplasmic reticulum function with insulin signaling and is essential for normal glucose homeostasis. Proc Natl Acad Sci USA 109, 5523–5528 (2012).

28. Aljanabi, S. M. & Martinez, I. Universal and rapid salt-extraction of high quality genomic DNA for PCR-based techniques. Nucleic Acids Research 25, 4692–4693 (1997).

29. Buan, H. Zarnowski, M. J., Cushman, S. W. & Lodish, H. F. Standard isolation of primary adipose cells from mouse epidydimal fat pads induces inflammatory mediators and down-regulates adipocyte genes. J Biol Chem 278, 47585–47593 (2003).

30. Iqbal, J. et al. An intrinsic gut leptin-melanocortin pathway modulates intestinal microsomal triglyceride transfer protein and lipid absorption. J Lipid Res 51, 1929–1942 (2010).

31. Tannriver, A., et al. Intestinal deletion of leptin signaling alters activity of nutrient transporters and delayed the onset of obesity in mice. Faseb J 28, 4100–4110 (2014).

32. Higurashi, T. et al. Conditional knockout of the leptin receptor in the colonic epithelium revealed the local effects of leptin receptor signaling in the progression of colonic tumours in mice. Carcinogenesis 35, 2134–2141 (2014).

33. Su, X. et al. Expression of FABP4, adipin and adiponectin in Paneth cells is modulated by gut Lactobacillus. J Endocrinol 217, 1984–1993 (2017).

34. Sakaguchi, M. et al. Adipocyte dynamics and reversible metabolic syndrome in mice with an inducible adipocyte-specific deletion of the insulin receptor. Cell Metabolism 25, 448–462 (2017).

35. Davis, R. C. et al. Early hepatic insulin resistance precedes the onset of diabetes in obese C57BLKS-db/db mice. Diabetes 59, 1616–1625 (2010).

36. Berry, R. & Rodeheffer, M. S. Characterization of the adipocyte cellular lineage in vivo. Nature Cell Biology 15, 302–308 (2013).

37. Lee, M. J. & Fried, S. K. Multilevel regulation of leptin storage, turnover, and secretion by feeding and insulin in rat adipose tissue. J Lipid Res 47, 1984–1993 (2006).

38. Li, Z., Ceccarini, G., Eisenstein, M., Tan, K. & Friedman, J. M. Phenotypic effects of an induced mutation of the ObRa isoform of the leptin receptor. Mol Metab 2, 364–375 (2013).

39. Yang, W. H. et al. Leptin induces IL-6 expression through OBR receptor signaling pathway in human synovial fibroblasts. PLoS ONE 8, e75551 (2013).

40. Dunnmore, S. J. & Brown, J. E. The role of adipokines in beta-cell failure of type 2 diabetes. J Endocrinol 216, T37–45 (2013).

41. Best, J. D. et al. Role of glucose effectiveness in the determination of glucose tolerance. Diabetes Care 19, 1018–1030 (1996).

42. Pacini, G., Thomaseth, K. & Ahren, B. Contribution to glucose tolerance of insulin-independent vs. insulin-dependent mechanisms in mice. Am J Physiol Endocrinol Metab 281, E693–703 (2001).

43. Harris, R. B. Direct and indirect effects of leptin on adipocyte metabolism. Biochim Biophys Acta 1842, 414–423 (2014).

44. Perry, R. J., Petersen, K. F. & Shulman, G. I. Pleotropic effects of leptin to reverse insulin resistance and diabetic ketoacidosis. Diabetologia 59, 933–937 (2016).

45. Meek, T. H. & Morton, G. J. The role of leptin in diabetes: metabolic effects. Diabetologia 59, 928–932 (2016).

46. Fujikawa, T. et al. Leptin engages a hypothalamic neurocircuitry to permit survival in the absence of insulin. Cell Metab 18, 431–444 (2013).

47. Xu, Y., Chang, J. T., Myers, M. G. Jr., Xu, Y. & Tong, Q. Euglycemia Restoration by Central Leptin in Type 1 Diabetes Requires STAT3 Signaling but Not Fast-Acting Neurotransmitter Release. Diabetes 65, 1040–1049 (2016).

48. Xu, L. et al. Genetic identification of leptin neural circuits in energy and glucose homeostases. Nature 556, 505–509 (2018).

49. Perry, R. J. et al. Leptin reverses diabetes by suppression of the hypothalamic–pituitary–adrenal axis. Nat Med 20, 759–763 (2014).

50. Morton, G. J., Meek, T. H., Matsen, M. E. & Schwartz, M. W. Evidence against hypothalamic–pituitary–adrenal axis suppression in the antidiabetic action of leptin. Journal of Clinical Investigation 125, 4587–4591 (2015).

51. Perry, R. J. et al. Mechanism for leptin’s acute insulin-independent effect to reverse diabetic ketoacidosis. Journal of Clinical Investigation 127, 657–669 (2017).

52. Huang, L., Wang, Z. & Li, C. Modulation of circulating leptin levels by its soluble receptor. J Biol Chem 276, 6343–6349 (2001).
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Author Contributions
S.P. and T.J.K. designed the study. S.P., S.M.O., T.D.W., V.S., J.S.Y., R.K.B., M.M., M.M.G., A.A., S.K. and S.D.C. performed the research. S.P., C.E., M.M., S.K. and S.M.C. analyzed the data. S.P. and T.J.K. wrote the article.

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