# Selective deletion of leptin receptor in neurons leads to obesity

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Animals with mutations in the leptin receptor (ObR) exhibit an obese phenotype that is indistinguishable from that of leptin deficient *ob/ob* mice. ObR is expressed in many tissues, including brain, and the relative importance of leptin's effects on central versus peripheral sites has not been resolved. To address this, we generated mice with neuron-specific (ObR<sup>SynI</sup>KO) and hepatocyte-specific (ObR<sup>Alb</sup>KO) disruption of ObR. Among the ObR<sup>SynI</sup>KO mice, the extent of obesity was negatively correlated with the level of ObR in hypothalamus and those animals with the lowest levels of ObR exhibited an obese phenotype. The obese mice with low levels of hypothalamic ObR also show elevated plasma levels of leptin, glucose, insulin, and corticosterone. The hypothalamic levels of agouti-related protein and neuropeptide Y RNA are increased in these mice. These data indicate that leptin has direct effects on neurons and that a significant proportion, or perhaps the majority, of its weight-reducing effects are the result of its actions on brain. To explore possible direct effects of leptin on a peripheral tissue, we also characterized ObRAIbKO mice. These mice weigh the same as controls and have no alterations in body composition. Moreover, while *db/db* mice and ObR<sup>SynI</sup>KO mice have enlarged fatty livers, ObR<sup>Alb</sup>KO mice do not. In summary, these data suggest that the brain is a direct target for the weight-reducing and neuroendocrine effects of leptin and that the liver abnormalities of *db/db* mice are secondary to defective leptin signaling in the brain.

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# Introduction

Leptin is an adipocyte hormone that functions as an afferent signal in a negative feedback loop that regulates energy balance (1-3). Mice with mutations in leptin (*ob/ob*) or its receptor (*db/db*) are hyperphagic and severely obese (4-6). These mutant mice also manifest a large number of metabolic and endocrine abnormalities including diabetes, hypercortisolemia, infertility, and cold intolerance (7,8). These *ob/ob* and *db/db* mice also have enlarged, steatotic livers (9, 10). The fatty liver is in part a result of an increased rate of hepatic lipogenesis, which is also thought to contribute to the development of obesity (11).

Treatment of *ob/ob* mice with leptin reduces food intake and body weight and corrects the metabolic and endocrine defects associated with leptin deficiency. Leptin treatment also normalizes the hepatomegaly and associated elevations in hepatic glycogen and lipid (12). Infusions of leptin into wild-type mice in physiological amounts results in a dose-dependent reduction in food intake and body weight (13–15). This metabolic response to leptin in both *ob/ob* and wild-type mice is novel and is not solely a consequence of its anorectic effects (16). Intracerebroventricular (ICV) leptin has similar effects, but at much

lower doses, suggesting that leptin has direct effects on brain (15, 17).

The leptin receptor, which has five splice variants (ObRa-e), is broadly expressed, however, and the relative importance of leptin's effects on brain versus peripheral sites is unclear (18, 19). This distinction is of importance because leptin deficiency is associated with myriad abnormalities in many tissues including liver. Although leptin has been suggested to act directly to activate signal transduction and deplete triglycerides, the contribution of these actions to body weight homeostasis, deposition of lipid in peripheral sites, and neuroendocrine function in vivo is untested (20, 21). To determine the role of leptin action in the central nervous system and the periphery, we are systematically deleting ObR in a tissue-specific fashion using the Cre-loxP system. Here we present data from mice with either neuronal or hepatocyte-specific deletions of ObR.

# Methods

ObR floxed mice. A 12-kb genomic clone containing the first coding exon of ObR was isolated. A single loxP element was PCR-amplified from the plasmid pNeoTKLox. The oligonucleotide primers were engineered to introduce a BamHI-NcoI double restriction site at one end and a BamHI site at the other end. After digestion with BamHI, the product was ligated into the BamHI site of the genomic clone, resulting in the clone BNLOXB with a single LoxP site upstream of the first exon. A loxP-Neo-HSV-TK-LoxP element was isolated by digesting pNeoTKLox with BstXI and Sall. The fragment was blunt-ended and cloned into BglII digested blunt-ended

BNLOXB. This clone, denoted ObR-lox-SS1-lox, was double digested with AfIII and KpnI, blunt-ended, and re-ligated to generate the targeting vector. The excised 500-bp AfIII-KpnI fragment was used as probe 1. The targeting vector was linearized with SalI and electroporated into 129/SV embryonic stem (ES) cells. Cells were selected with G418, and surviving clones were screened for homologous recombination. Positive clones were transiently transfected with the Cre recombinase expressing plasmid, p0G231 (kindly provided by S. O'Gorman) (22). Cells were then subjected to ganciclovir selection, and surviving clones were checked by Southern blotting using probe 2, which is within exon 2, to confirm the deletion of the Neo-HSV-TK cassette (type II deletion). ES cells with the correct genotype (ObRflox/+) were injected into C57BL/6 blastocysts, and resulting chimeras were bred with C57BL/6 mice to obtain germline transmission. ObRflox/+ mice were crossed to generate the line of ObRfloxed mice, denoted ObR<sup>flox/flox</sup>.

*ObR-null mice. ObR*<sup>flox/+</sup> mice were crossed with transgenic adenovirus EIIA Cre mice (kindly provided by H. Westphal) (23). Given that Cre is expressed early in embryogenesis in this line, progeny were screened by Southern blotting with probe 2 for germline Cre-mediated deletion of the first coding exon (type I deletion). These mice, with the genotype  $ObR^{A/+}$ , were crossed to produce homozygous  $ObR^{A/A}$  mice, designated ObR-null mice.

Conditional deletion of ObR. Neuronspecific deletion of ObR was achieved using synapsinI-Cre transgenic mice (SynI-Cre(+)) (kindly provided by J. Marth) (24, 25). Hepatocyte-specific deletion was achieved using Albumincre transgenic mice (Alb-Cre(+)). The Alb-Cre transgene (see Figure 5a) was constructed using plasmid NB, which contains 2 kb of the Albumin promoter enhancer (kindly provided by R. Palmiter) (26). A 2-kb fragment containing the Cre-recombinase gene and a nuclear localization and polyadenylation signal was excised from a separate plasmid by digesting with BglII and blunt-ending and then digesting with KpnI. This fragment was cloned into KpnI-EcoRV digested plasmid NB downstream of the albumin promoter

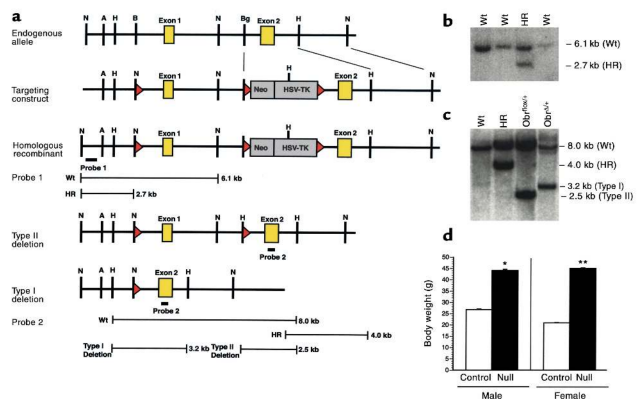
enhancer. The resulting plasmid was digested with NotI and KpnI to release the 4-kb transgene, which was gel purified and injected into fertilized eggs from  $C57/BL6 \times CBA$  (F1) mice to produce transgenic mice. The presence of the transgene in both (SynI-Cre(+)) and (Alb-Cre(+)) mice was verified by PCR and Southern blotting. Tissue-specific knockout mice were generated by two successive crosses (see Figure 2a). First,  $ObR^{\Delta/+}$  mice were crossed with either synapsinI-Cre transgenic mice (SynI-Cre(+)) or albumin-cre transgenic mice (Alb-Cre(+)) to generate ObR $^{\Delta/+}$ , SynI-Cre(+) mice or  $ObR^{\Delta/+}$ , Alb-Cre(+) mice. These mice were then mated to *ObR<sup>flox/flox</sup>* mice to generate mice with the genotype ObR<sup>∆/flox</sup>, SynI-Cre(+) or  $ObR^{\Delta/flox}$ , Alb-Cre(+), hereafter designated ObR<sup>SynI</sup>KO and ObR<sup>Alb</sup>KO, respectively. Mice with the following genotypes were also generated: (a)  $ObR^{\Delta/flox}$ , SynI-Cre(-) and  $ObR^{\Delta/flox}$ , Alb-Cre(-) (heterozygotes); (b) ObR-<sup>flox/+</sup>, SynI-Cre(+) and ObR<sup>flox/+</sup>, Alb-Cre(+) (referred to as wild-type); and (c) ObR<sup>flox/+</sup>, SynI-Cre(-) and ObR<sup>flox/+</sup>, Alb-Cre(-) (referred to as wild-type). All animals were housed under controlled temperature (23°C) and lighting (12 hours of light; 12 hours of dark) with free access to food and water. All procedures were in accordance with the guidelines of the Rockefeller University Laboratory Animal Research Center.

Assay of Cre specificity. Genomic DNA was prepared from multiple tissues from ObR<sup>flox/+</sup> SynI-Cre(+) and ObR<sup>flox/+</sup> Alb-Cre(+) mice, and 200 ng was PCRamplified using primers flanking the first coding exon of ObR. A schematic of the primer locations is shown in Figure 2b. In tissues expressing Cre recombinase, exon 1 is excised and a single loxP site remains, generating an ObR<sup>∆</sup> allele. Although primers 1 and 3 can amplify a product from the  $ObR^{\Delta}$  allele, the primers are separated by too great a distance for amplification to occur in the ObR<sup>flox</sup> or ObR<sup>+</sup> (wild-type) alleles. As a control, primers 1 and 2 were used to amplify both the ObR<sup>flox</sup> and ObR<sup>+</sup> alleles from all tissues. The ObR<sup>+</sup> allele produces a slightly smaller product owing to the absence of loxP sequence.

Assay of Cre efficiency. To determine the degree of Cre-mediated recombination, ObR expression levels from all

ObR<sup>SynI</sup>KO and ObR<sup>Alb</sup>KO mice and a sample of mice with all other possible genotypes were quantitated using Taqman real-time PCR (Perkin Elmer Applied Biosystems, Foster City, California, USA). Total RNA was isolated from ObR<sup>SynI</sup>KO and ObR<sup>Alb</sup>KO mice and reverse transcribed into cDNA using random hexamers with the Reverse Transcription Reagents from Roche Molecular Systems (Branchburg, New Jersey, USA). Expression levels were determined using 25 ng of each cDNA sample assayed in duplicate and amplified with the ABI Prism 7700 Sequence Detection System (Perkin Elmer Applied Biosystems). The location of primers and fluorescent probes is indicated in Figure 3b. As amplification occurs, the probe is cleaved, resulting in a signal from a reporter dye that is directly related to the amount of amplicon. Primers were derived from sequences in the 5' untranslated region (fwd) and the second coding exon (rev). The fluorescent probe, labeled with the FAM dye, contains sequence located within the first coding exon. When this exon is deleted, the probe cannot be cleaved, and no signal is generated. As a control for input amount, each cDNA sample was also amplified using primers and a probe, labeled with the VIC dye, for cyclophilin. Data were analyzed with the ABI Sequence Detector software (Perkin Elmer Applied Biosystems). Every set of reactions contained a set of four serial twofold dilutions of the same liver cDNA source, which was used to generate a standard curve for both ObR and cyclophilin. Amounts of each transcript were calculated from the standard curve, and levels of ObR were corrected for levels of cyclophilin (ObR level/cyclophilin level).

Body composition analysis. Body composition was analyzed as described previously (13). Carcasses were weighed and then oven dried in a 90°C oven until weight was constant. The total body water was calculated as the difference between the weights before and after drying. The carcass was then homogenized in a blender, and duplicate aliquots were extracted with a Soxhlet apparatus using a 3:1 mixture of chloroform/methanol. The extracted homogenate was dried overnight and weighed to calculate fat mass and lean mass.



# Figure 1

LoxP targeting of the ObR locus. Gene targeting was used to insert loxP sites on either side of the first coding exon of ObR. (**a**) Restriction maps (from top to bottom) of the genomic locus, targeting vector, homologous recombinant, and the type II and type I deletion alleles. Probe 1, located outside the targeting construct, was used to screen for homologous recombinants. Probe 2, located in the second coding exon, distinguishes the endogenous allele, homologous recombinants, and type I (deletion of the first coding exon) and II (deletion of the targeting cassette leaving loxP sites on either side of the first coding exon) deletions. A, AfIII; B, BamHI; Bg, BgIII; H, HindIII; N, Ncol. (**b**) Southern blot analysis of Ncol-digested genomic DNA from ES cell clones using probe 1. The endogenous allele (Wt) and homologous recombinants (HR) migrated at the predicted sizes. (**c**) Southern blot analysis of Hind-III digested genomic DNA using probe 2. The endogenous allele (Wt), homologous recombinant (HR), and type II deletion were detected from ES cell DNA. ObR<sup>flox/+</sup> mice were generated from the type II deletion and crossed to adenovirus EIIA-Cre mice. The type I deletion was detected in tail DNA from progeny derived from this cross. All alleles migrated at the predicted sizes. (**d**) Body weight at 8 weeks of age of ObR<sup>Δ/Δ</sup> and littermate control (ObR<sup>Δ/+</sup>) mice. Data represent the mean ± SEM of at least nine animals of each genotype and gender. \**P* < 0.02; \*\**P* < 0.001; unpaired Student's *t* test.

Neuropeptide expression levels. Neuropeptide levels were quantitated by Taqman real-time PCR from individual hypothalami. Expression levels were measured in the obese ObR<sup>SynI</sup>KO mice, in heterozygote controls, and in female ObR-null mice. Primers and probes were made for agouti-related protein (AGRP), neuropeptide Y (NPY), melanin concentrating hormone (MCH), proopiomelanocortin (POMC), and cocaine and amphetamine regulated transcript (CART).

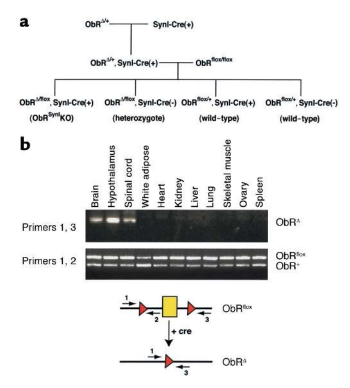
*Neuroendocrine function*. After sacrifice, mice were exsanguinated and blood was collected on EDTA. Plasma was collected after centrifugation and used for all assays. Leptin levels were determined using an ELISA kit from R&D Systems Inc. (Minneapolis, Minnesota, USA). Insulin levels were determined using an ELISA kit from Alpco Diagnostics (Windham, New Hampshire, USA). Glucose levels were determined using Trinder reagents from Sigma Chemical Co. (St. Louis, Missouri, USA). Triglyceride levels were determined using an enzymatic kit from Wako Chemical USA Inc. (Richmond, Virginia, USA). Corticosterone, estradiol, testosterone, and thyroxine levels were determined using RIA kits from ICN Radiochemicals Inc. (Costa Mesa, California, USA).

Liver triglyceride quantitation. A total of 40–100 mg of liver from each mouse was homogenized in 4 ml of chloroform-methanol (2:1). A total of 0.8 ml of 50 mM NaCl was added to each sample. Samples were then centrifuged for 5 minutes at 1,300 g. The lower phase was removed, and duplicate 50- $\mu$ l aliquots were evaporated under N<sub>2</sub> gas and then assayed for triglycerides using the Trinder reagent from Sigma Chemical Co. Values are expressed as milligrams of triglyceride per gram of liver.

Statistical analysis. Data are expressed as means  $\pm$  SE. Unless otherwise indicated, significance was evaluated using the unpaired Student's *t* test. Significance of correlation coefficients was evaluated using the *t* test for correlation.

# Results

We have used the Cre-loxP system to generate mice with tissue-specific deletions of ObR (27). The construction of the targeting plasmid is described above (Figure 1a). The first coding exon of ObR was flanked by loxP sites. This exon contains the signal sequence, and deletion of it by Cre-mediated recombination was predicted to inacti-



# Figure 2

Cre-mediated recombination specifically in the brain of ObR<sup>Syn1</sup>KO mice. (**a**) The breeding strategy that was used to generate ObR<sup>Syn1</sup>KO mice and littermate controls. ObR<sup>Alb</sup>KO mice (described below) were generated using the same strategy. (**b**) Genomic DNA was prepared from several tissues from ObR<sup>flox/+</sup>, Syn1-Cre(+) mice and were PCR-amplified using primers flanking the first coding exon of ObR. The locations of the primers are shown on the schematic below the gel. In tissues expressing the Cre recombinase, exon 1 is excised and a single loxP site remains, generating an ObR<sup>4</sup> allele. While primers 1 and 3 can amplify a product from the ObR<sup>4</sup> allele, in the ObR<sup>flox</sup> or ObR<sup>+</sup> (wild-type) DNA, the primers are separated by a distance too great for amplification to occur. As a control, primers 1 and 2 were used to amplify both the ObR<sup>flox</sup> allele and the ObR<sup>+</sup> alleles from all tissues. The ObR<sup>+</sup> allele produces a slightly smaller product due to the absence of loxP sequences.

vate all splice variants. After transfection into mouse ES cells, three of 72 G418-resistant clones had undergone homologous recombination (Figure 1b). The targeting cassette was deleted by transiently transfecting one of the homologous recombinant ES cell lines with a plasmid expressing Cre recombinase and screening for clones with a type II deletion (Figure 1a). Clones with the correct genotype, ObRflox/+, were confirmed by Southern blotting and injected into C57BL/6 blastocysts (Figure 1c). Breeding of chimeras confirmed that animals with a germline insertion of loxP sites into ObR had been generated. Mice homozygous for the floxed allele, ObRflox/flox, were generated by crossing *ObR<sup>flox/+</sup>* animals.

*ObR*<sup>flox/flox</sup> mice were viable, fertile, and indistinguishable from wild-type or *ObR*<sup>flox/+</sup> mice, indicating that the insertion of loxP sites into the introns flanking the first coding exon did not interfere with ObR function. To determine whether Cre-mediated deletion of the first coding exon results in a null allele, *ObR<sup>flox/+</sup>* mice were crossed with transgenic mice expressing Cre at an early embryonic stage (23). Mice with the genotype  $ObR^{\Delta/+}$ , heterozygous for a germline deletion of the first coding exon (type I deletion), were generated and mated to produce homozygous  $ObR^{\Delta/\Delta}$  mice, also referred to as ObR-null mice (Figure 1c). ObRnull mice are massively obese and indistinguishable from *db*<sup>3J</sup>/*db*<sup>3J</sup> mice, which are also null for all ObR isoforms (Figure 1d) (28). These data confirm that the floxed allele is wild-type and that the deleted allele is null, thus validating the targeting strategy.

Neuron-specific ObR-knockout mice, designated ObR<sup>SynI</sup>KO, were generated by two successive crosses (Figure 2a, and see Methods). These mice carry one floxed and one null allele so that Cremediated inactivation of only one allele, rather than two, is sufficient to delete ObR in a given cell. The ObR<sup>Syn1</sup>KO mice were compared to littermate controls with the genotype ObR<sup> $\Delta$ /flox</sup>, SynI-Cre(–), also referred to as heterozygotes. These mice were used as controls because some phenotypic changes have been observed in db/+ mice (29). Data were also obtained from lean littermates with the genotypes ObR<sup>flox/+</sup>, SynI-Cre(+) or ObR<sup>flox/+</sup>, SynI-Cre(–), both of which are referred to as wild-type. Genotypes of all mice were determined using PCR and Southern blotting.

In the synapsinI-Cre transgenic line, expression of Cre recombinase is controlled by the rat synapsin I promoter. This promoter has been shown to drive Cre expression specifically in neurons (30). Crossing SynI-Cre(+) mice to LacZ indicator transgenic strains showed that Cre activity was first detectable at embryonic day (E) 12.5 and restricted to brain, spinal cord, and the dorsal root ganglion and was absent from astrocytes and glia (25). To confirm the tissue specificity of Cre-mediated recombination, a qualitative PCR assay was developed that was capable of detecting recombination between the lox sites flanking the first coding exon of ObR in genomic DNA. In ObR<sup>flox/+</sup>, synapsinI-Cre(+) mice, Cre-mediated recombination was restricted to brain, hypothalamus, and spinal cord (Figure 2b). As previously indicated by reporter gene expression in SynI-CAT transgenic mice, low level recombination was also seen in testis (data not shown).

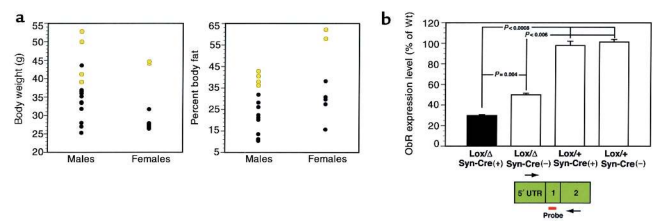
The weight of the ObR<sup>SynI</sup>KO mice was compared to that of heterozygous mice at 4 months of age. Although the weights of ObR<sup>SynI</sup>KO mice, on average, were no different than heterozygotes, in both sexes a subset of the ObR<sup>SynI</sup>KO mice had an increased body weight (Figure 3a). The percent body fat of these mice (Figure 3a, yellow) was more than 2.5 SD greater than the average percent body fat of heterozygotes (Figure 3a). This suggested the possibility that the extent of the knockout of ObR was variable and that the most obese ObR<sup>SynI</sup>KO animals had the lowest levels of ObR. To assess the efficiency of Cre-mediated recombination, real-time PCR assays (Taqman) were performed to quantitate ObR RNA using cDNA prepared from individual hypothalami (31). Previous data have suggested that in neurons, ObR is expressed at highest levels in hypothalamus and at much lower levels in other brain regions. This assay detected all forms of the leptin receptor. ObR<sup>flox/+</sup>, SynI-Cre(+) and ObR<sup>flox/+</sup>, SynI-Cre(-) mice had equivalent levels of ObR expression, and thus both genotypes were designated as wild-type (Figure 3b). As expected, ObR<sup>Δ/flox</sup>, SynI-Cre(-) mice, which have one allele inactivated in the germ line, have 50% as much RNA as wild-type mice (P < 0.006). The average level of ObR RNA was significantly lower in ObR<sup>SynI</sup>KO mice than in ObR<sup>∆/flox</sup>, SynI-Cre(-) heterozygote littermates (30% vs. 50% of wild-type, respectively; P = 0.004). Moreover, although many of the ObR<sup>SynI</sup>KO mice had levels of ObR RNA that were indistinguishable from heterozygotes, a subset of the ObR<sup>SynI</sup>KO mice had markedly reduced levels of ObR. These results confirmed that Cremediated recombination is variable, and indicated that in many animals there is no evident recombination.

We next considered whether the most obese ObR<sup>SynI</sup>KO (i.e., percent body fat 2.5 SD > heterozygotes) animals had lower levels of ObR RNA than the lighter ones. Analysis of the data indicated that percent body fat and body weight were significantly increased in each of six animals in which ObR RNA was less than 15% that of wild-type (Figure 3a, yellow) (P < 0.005 for percent body fat; P < 0.05 at all ages greater than 5 weeks for body weight). This confirmed that a significant deletion of ObR is associated with an obese phenotype. Furthermore, in those cases where the deletion is most extreme, a severely obese phenotype is evident. Two ObR<sup>SynI</sup>KO females and one ObR<sup>SynI</sup>KO male had a greater than 97% reduction of ObR RNA, and each weighed more than 50 g at sacrifice, weights that approach, but do not equal, those of *db/db* mice. In these studies, as well as studies of the ObRAlbKO mice (see below), there were no lean mice with hypothalamic ObR RNA levels less than 15% that of wild-type, and there were no obese mice (obese defined as percent body fat > 2.5 SD above heterozygotes: 30.5% for males, 47.8% for females) with RNA levels greater than 15% that of wild-type. These data suggest that a near-normal body weight can be maintained until markedly reduced levels of ObR in hypothalamus are evident. In all following studies, the phenotype of the animals with less than 15% wild-type levels of hypothalamic ObR RNA was compared to heterozygotes. These mice are referred to as obese ObR<sup>SynI</sup>KO mice.

The weights of the obese ObR<sup>SynI</sup>KO mice were significantly increased relative to heterozygotes at all time points greater than 5 weeks of age (P < 0.05).

At 5 months of age, males and females weighed 35% and 66% more than heterozygotes, respectively (Figure 4a). The increase in body weight was associated with increased adipose tissue mass and, as is also the case in ob/ob and db/db mice, decreased lean body mass (Table 1). At sacrifice, the percent body fat of ObR<sup>SynI</sup>KO mice was 39% in males and 60% in females, compared with 15% in male heterozygotes and 26% in female heterozygotes.

The obese ObR<sup>SynI</sup>KO mice were also analyzed with respect to a number of other abnormalities associated with the *db/db* mutation (Table 1). Plasma leptin concentrations were elevated 5.8-fold in obese ObR<sup>SynI</sup>KO males (P < 0.05) and 8.3-fold in obese ObRSynIKO females (P < 0.0001) and were highly correlated with percent body fat (r = 0.82, P < 0.01, t test for correlation). In addition, plasma insulin was increased sixfold in obese ObR<sup>SynI</sup>KO males (P < 0.05) and 16.1-fold in ObRSynIKO females. Glucose was increased 2.3-fold in  $ObR^{SynI}KO$  females (P < 0.005), and was unchanged in ObRSynIKO males. Plasma corticosterone was elevated 1.8-fold in ObR<sup>SynI</sup>KO males and 2.4-fold in ObR<sup>SynI</sup>KO females (P < 0.05). The elevation in plasma glucose and the greater elevation in corticosterone specific to females could be due to the females analyzed having a lower level of ObR RNA and being more obese. Of



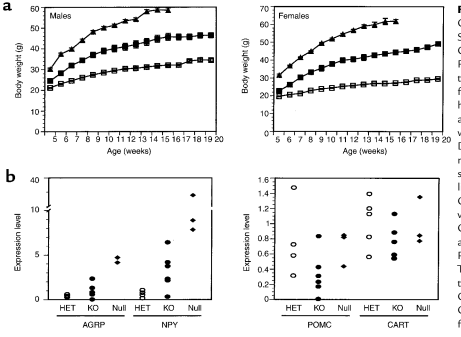
#### Figure 3

ObR RNA levels. (**a**) The distribution of body weight and percent body fat for male and female ObR<sup>SynI</sup>KO mice at 16 weeks of age. Those mice with less than 15% of ObR RNA (see **b**) are shown in yellow. (**b**) Expression levels were determined using Taqman real-time PCR with the ABI Prism 7700 Sequence Detection System. The locations of primers and fluorescent probe are as indicated. Primers were derived from sequences in the 5' untranslated region and the second coding exon. The fluorescent probe is located within the first coding exon 1. When this exon is deleted, no signal is generated. As a control for input amount, each cDNA sample was also amplified using primers and a probe for cyclophilin. Data were analyzed with ABI Sequence Detector software, and the levels of ObR were normalized to cyclophilin. Levels are represented as the percentage of the levels in ObR<sup>flox/+</sup>, SynI-Cre(+) and ObR<sup>flox/+</sup>, SynI-Cre(-) wild-type mice. Data represent the mean  $\pm$  SEM. At least 13 animals were analyzed for each genotype. *P* values for comparisons between genotypes are indicated.

note, an increased plasma corticosterone is unique to *ob/ob* and *db/db* mice, whereas other forms of rodent obesity are not generally associated with elevated corticosterone (32). Mutations in leptin or its receptor are also associated with infertility, and leptin has been shown to influence the hypothalamicpituitary-gonadal axis (33). Consistent with this, obese female ObR<sup>SynI</sup>KO mice had 43% reductions in plasma estradiol. Male ObR<sup>SynI</sup>KO mice, however, did not have reduced levels of plasma testosterone. This sexual dimorphism is consistent with the fact that leptin deficiency has a more profound effect on female reproduction than male reproduction (34). Finally, plasma thyroxine and triglycerides were not significantly different between mutant and control mice (data not shown).

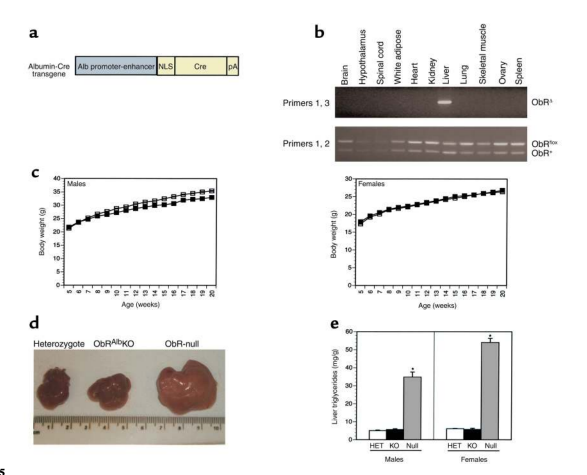
Despite the marked obesity in this subset of  $ObR^{SynI}KO$  mice, these animals still weighed less than mice with germline deletions of ObR (Figure 4a; P < 0.05 at all ages). These data suggest either that the knockout was incomplete even in the most obese animals or that leptin also acts at peripheral sites to reduce weight. To further assess the extent of the knockout in brain, levels of a number of hypothalamic neuropeptides were examined using Taqman assays (Figure 4b). Defective leptin signaling is associated with increased levels of hypothalamic AGRP, NPY, and MCH RNA and reduced levels of POMC and CART RNA (35-40). Comparisons of heterozygotes to obese ObR<sup>SynI</sup>KO mice showed that AGRP and NPY levels were increased, whereas POMC and CART levels showed a trend toward being reduced. Although the reduction in POMC and CART RNA levels is not significant, the levels of these RNAs are only modestly decreased in ob/ob and *db/db* hypothalamus. MCH levels were indistinguishable between heterozygous mice, ObR<sup>SynI</sup>KO mice, and ObR-null mice. The reported 80% increase in MCH RNA in *ob/ob* mice is also more modest than the reported increases in AGRP and NPY RNA (38). Although the levels of AGRP and NPY were increased in the obese ObR<sup>SynI</sup>KO mice, they were less elevated than in null mice. This suggests that, despite the low levels of ObR RNA in these mice, some ObR-expressing neurons remained.

The data here indicate that leptin has direct effects on brain but leave open the possibility that leptin also has direct effects on peripheral tissues. To assess a possible role of ObR in liver, a peripheral tissue that has been suggested to be a direct target of leptin action, mice with a hepatocyte-specific knockout of ObR were generated (ObR<sup>Alb</sup>KO). These mice were generated using the same breeding strategy as described for ObR<sup>SynI</sup>KO mice (Figure 2a). The albumin promoter was used to direct hepatocyte-specific expression of Cre recombinase (Figure 5a) (26). Genomic PCR performed on multiple tissues from ObR<sup>flox/+</sup> albumin-Cre(+) mice confirmed that Cre-mediated recombination was restricted to liver (Figure 5b). In contrast to ObR<sup>SynI</sup>KO mice, ObRAlbKO mice did not manifest an increased weight even in those animals in which the levels of ObR RNA in liver (as measured using the aforementioned Taqman assay) were reduced to less than 30% of wild-type levels (9 of 12 males and 11 of 16 females analyzed) (Figure 5c). Because the liver is composed of a number of different cell types, and 80% of the mass is accounted for by hepatocytes, this likely represents a nearcomplete deletion in hepatocytes (41, 42). Inspection of the weight distributions showed that no ObRAlbKO mice manifested an increased weight. Body composition was analyzed to determine whether hepatocyte-specific deletion of ObR had a subtle effect on body partitioning in ObRAlbKO mice. Water mass, lean mass, and fat mass were all unchanged in these animals (Table 1). In both sexes, plasma leptin levels were within the normal range as were insulin, glucose, and corticosterone. Although *db/db* and *ObR<sup>SymI</sup>*KO mice have enlarged fatty livers characteristic of leptin-deficient mice, the livers of ObRAlbKO mice were grossly normal (Figure 5d). Wet liver weight and liver weight as a percentage of body weight were also



#### Figure 4

Obesity in a subset of ObR<sup>SynI</sup>KO mice. Significant obesity was evident in those ObR<sup>SynI</sup>KO mice that had 15% or less ObR RNA in the hypothalamus than did wildtype mice. (a) Weight curves of male and female ObR<sup>SynI</sup>KO mice (filled squares), heterozygote littermates (open squares), and ObR-null mice (filled triangles). Mice were weighed weekly from 5 weeks of age. Data represent the mean ± SEM of four male and two female ObR<sup>SynI</sup>KO mice, seven male and nine female heterozygote littermates, and six male and five female ObR-null mice. For both sexes, ObR<sup>SynI</sup>KO versus heterozygotes, *P* < 0.05 at all ages; ObR<sup>SynI</sup>KO versus ObR-null, P < 0.05 at all ages. (b) Expression levels of AGRP, NPY, POMC, and CART were determined by Tagman and are expressed as normalized to cyclophilin. Levels were measured in ObR<sup>SynI</sup>KO, heterozygotes (HET), and ObR-nulls. (n = 5 for heterozygotes, n = 6for knockouts, and n = 3 for null mice.)



# Figure 5

Normal body weight and liver phenotype in ObR<sup>Alb</sup>KO mice. (**a**) Schematic of *Albumin*-Cre transgene. (**b**) Genomic DNA was prepared from several tissues from ObR<sup>flox/+</sup>, Alb-Cre(+) mice and was PCR-amplified using primers flanking the first coding exon of ObR as described in Figure 2b. (**c**) Weight curves of male and female ObR<sup>Alb</sup>KO mice (filled squares) and heterozygote littermates (open squares). Mice were weighed weekly from 5 weeks of age. Data represent the mean  $\pm$  SEM of 9 male and 11 female ObR<sup>Alb</sup>KO mice and 23 male and 23 female heterozygote littermates. For both sexes, ObR<sup>Alb</sup>KO versus heterozygotes, *P* values were not significant at all ages. (**d**) Photographs of freshly dissected livers from representative mice with the given genotypes. (**e**) Liver triglycerides (milligrams of triglyceride per grams of liver) were determined for ObR<sup>Alb</sup>KO mice with less than 30% of wild-type ObR RNA and heterozygote controls. Data represent the mean  $\pm$  SEM of six male and five female ObR<sup>Alb</sup>KO mice, five male and five female heterozygote controls, and six male and three female ObR-null mice. For both sexes, \**P* < 0.05 for ObR<sup>Alb</sup>KO and heterozygotes relative to ObR null.

unchanged between ObRAlbKO mice and heterozygotes (data not shown). Quantitation of liver triglycerides showed no difference between ObR<sup>Alb</sup>KO mice and heterozygote controls, whereas levels in ObR-null mice were significantly elevated relative to both of these groups (P < 0.05) (Figure 5e). These data indicate that ObR expressed in hepatocytes is not likely to play a significant role in body weight homeostasis or the hepatic steatosis associated with the *db* mutation, and that the liver abnormalities observed in ob/ob and db/db mice are secondary to defective leptin signaling in the brain.

# Discussion

In this study, two possible sites of leptin action were evaluated by analyzing

the phenotype of mice with neuron and hepatocyte-specific knockouts of the leptin receptor. These data provide direct genetic evidence that leptin signals via direct effects on neurons. Separate data from studies of dbKls/dbKls mice indicate that the specific absence of the ObRb form of the receptor results in a phenotype indistinguishable from leptin deficiency, demonstrating that ObRb is absolutely required for the weight-reducing effects of the hormone (4-6). ObRb is the only isoform that has all of the motifs necessary for signal transduction and is expressed at high levels in brain and a number of other tissues.

These findings suggest that leptin exerts its weight-reducing and some, or perhaps all, of its neuroendocrine effects via interactions with the ObRb form of the receptor in specific classes of neurons in brain. This is consistent with the high potency of leptin-administered ICV and the anatomic distribution of ObRb (see below) (15, 17). This conclusion is also consistent with data from experiments in which an NSE-ObRb transgene was able to partially complement the phenotype of db/db mice (43). However, in this study, ObRb was overexpressed 30-fold in all neurons, with some leakiness in peripheral tissues.

The phenotype of the obese  $ObR^{Syml}KO$ , while significant, did not reach that of ObR-null mice. In these animals, however, hypothalamic AGRP and NPY levels, while increased, were still not as high as in null mice, suggesting that there were still some residual

### Table 1

Body composition and neuroendocrine function in ObR<sup>SynI</sup>KO and ObR<sup>Alb</sup>KO mice<sup>A</sup>

|                        | ObR <sup>SynI</sup> KO | ObR <sup>SynI</sup> Het       | <b>O</b> bR <sup>AIb</sup> KO | ObR <sup>Alb</sup> Het |
|------------------------|------------------------|-------------------------------|-------------------------------|------------------------|
| Percent fat mass       |                        |                               |                               |                        |
| Males                  | 39.4 ± 0.7 (4)         | 15.4 ± 0.8 (8) <sup>B</sup>   | 14.0 ± 1.5 (4)                | 13.2 ± 0.5 (5)         |
| emales                 | 60.1 ± 1.5 (2)         | $25.8 \pm 0.9 (10)^{B}$       | $15.4 \pm 0.9$ (7)            | 15.6 ± 1.1 (5)         |
| Percent lean mass      |                        |                               |                               |                        |
| /ales                  | 19.7 ± 0.6 (4)         | 25.7 ± 0.2 (8) <sup>B</sup>   | 30.2 ± 0.8 (4)                | 30.2 ± 0.8 (5)         |
| emales                 | 12.7 ± 1.2 (2)         | $22.6 \pm 0.3 (10)^{B}$       | $28.0 \pm 0.7 (4)$            | 27.5 ± 0.7 (5)         |
| Percent water mass     |                        |                               |                               |                        |
| Males                  | $40.9 \pm 0.3$ (4)     | 59.0 ± 0.6 (8) <sup>B</sup>   | 55.8 ± 1.3 (4)                | 56.6 ± 0.6 (5)         |
| emales                 | 27.2 ± 0.3 (2)         | $51.6 \pm 0.6 (10)^{B}$       | 56.6 ± 0.9 (4)                | 56.9 ± 0.7 (5)         |
| _eptin (ng/ml)         |                        |                               |                               |                        |
| Aales                  | 73.8 ± 8.5 (4)         | 12.7 ± 0.4 (10) <sup>C</sup>  | 13.1 ± 0.8 (9)                | 13.9 ± 0.7 (15)        |
| emales                 | 125.0 ± 0.7 (2)        | $15.1 \pm 0.8 (12)^{B}$       | 12.7 ± 0.9 (6)                | 6.7 ± 0.2 (12)         |
| nsulin (ng/ml)         |                        |                               |                               |                        |
| Aales                  | 17.1 ± 2.1 (4)         | 2.8 ± 0.2 (8) <sup>C</sup>    | $1.9 \pm 0.1 (9)$             | 1.6 ± 0.1 (6)          |
| emales                 | 40.1 ± 6.5 (2)         | $2.5 \pm 0.2$ (8)             | $0.8 \pm 0.1$ (4)             | $0.7 \pm 0.1 (5)$      |
| Glucose (mg/dl)        |                        |                               |                               |                        |
| Aales                  | 158.3 ± 9.3 (4)        | 165.3 ± 4.9 (10)              | 80.3 ± 2.3 (7)                | 89.1 ± 2.5 (8)         |
| emales                 | 282.6 ± 32.3 (2)       | 123.1 ± 4.7 (8) <sup>B</sup>  | 87.3 ± 3.9 (5)                | 91.9 ± 2.0 (9)         |
| Corticosterone (ng/ml) |                        |                               |                               |                        |
| Aales                  | 238.5 ± 17.9 (4)       | 134.9 ± 40.2 (3)              | 54.0 ± 4.2 (6)                | 63.4 ± 2.7 (11)        |
| emales                 | 469.4 ± 32.7 (2)       | 195.9 ± 16.1 (8) <sup>C</sup> | 208.8 ± 25.1 (5)              | 108.5 ± 9.3 (9)        |
| Estradiol (pg/ml)      |                        |                               |                               |                        |
| Females                | $3.9 \pm 0.3 (2)$      | $6.8 \pm 0.8$ (3)             | ND                            | ND                     |

<sup>A</sup>After sacrifice, the subset of significantly obese ObR<sup>SynI</sup>KO mice and the subset of ObR<sup>Alb</sup>KO mice with less than 30% ObR RNA as well as heterozygote littermates were exsanguinated, and plasma was collected for the following assays. All plasma was collected from mice sacrificed at 1200 hours under free feeding conditions. Average ages for ObR<sup>SynI</sup>KO and ObR<sup>Alb</sup>KO mice sampled here are 26 weeks and 23 weeks. All data represent the mean  $\pm$  SEM, with the sample size indicated in parentheses. <sup>B</sup>P < 0.005 versus knockout; <sup>C</sup>P < 0.05 versus knockout in an unpaired Student's *t* test. ND, not determined.

ObR-expressing cells. This leaves open the possibility that a truly complete brain-specific knockout (i.e., deletion of ObR on every neuron) could recapitulate the obese phenotype of ObR-null mice. The generation of such a knockout might require the simultaneous use of multiple different neuron-specific Cre-expressing lines, now under way.

In some studies, the expression of ObRb in brain was reported to be restricted to the hypothalamus, possibly implicating the hypothalamus as the principal target of leptin's weight-reducing effects. In hypothalamus, ObRb is enriched in the arcuate, ventromedial, dorsomedial, and paraventricular hypothalamic nuclei, all of which are known, based on studies of animals with stereotactic lesions, to play a role in regulating food intake and body weight (19, 44–46). However, in other studies, ObRb expression has also been reported in other brain sites including the brainstem, thalamus, and cerebellum. The possibility that one or more of these

extrahypothalamic brain sites play important roles in the regulation of body weight can be tested using regionspecific or neuron-specific promoters to drive Cre expression in localized brain areas and specific cell types. In the hypothalamus, ObRb is expressed in functionally distinct classes of neurons, the best characterized being neurons that coexpress either NPY or  $\alpha$ MSH (47, 48). Both of these neuropeptides exert potent effects on food intake and body weight. Experiments to test the effects of ObR deletion in these two types of neurons may allow an analysis of the functional importance of these two neuropeptides as downstream effectors of leptin signaling and could establish the extent to which these two cell types by themselves can fully account for the neural response to leptin.

A number of studies have suggested that leptin also has direct effects on liver (20, 21). The data presented here indicate that hepatocyte-specific deletion does not have any observable effects on body weight or body composition and does not result in any gross liver pathology. Furthermore  $ObR^{Alb}KO$  mice have equivalent amounts of liver triglycerides as do controls, indicating that the hepatic phenotype of db/db mice is secondary to defective leptin signaling in brain. These findings do not exclude the possibility that hepatocyte ObR by itself serves functions, such as modulating leptin turnover.

Although these findings suggest that neurons are required for leptin's effects on body weight, they do not exclude the possibility that leptin also directly acts on other peripheral targets. In addition to regulating food intake, body weight, and the neuroendocrine axis, leptin can also modulate the immune system, bone metabolism, and angiogenesis (49–51). Indeed, ObRb is expressed in a number of peripheral tissues and direct effects of leptin on T cells, macrophages, pancreatic  $\beta$ -cells, and other cell types have been reported (20, 21, 52). Other forms of ObR (ObRa, c, d, e) may play a role in transport of leptin across the blood brain barrier and choroid plexus, sites that express high levels of ObR. The function of ObR at these and other sites can now be assessed by crossing ObR<sup>flox/flox</sup> mice to transgenic lines that express Cre in a tissue-specific fashion. Such studies should enable an assessment of the relative role of direct effects of leptin on these tissues versus indirect effects via signaling in brain. In conclusion, these findings indicate that the brain is a direct target for the weight-reducing and neuroendocrine effects of leptin and that the liver pathology of db/db mice is secondary to defective leptin signaling in neurons.

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