Late-Onset Leanness in Mice with Targeted Ablation of Melanin Concentrating Hormone Neurons

Tamar Alon and Jeffrey M. Friedman
Laboratory of Molecular Genetics, The Rockefeller University, New York, New York 10021

The observation that loss of orexin (hypocretin) neurons causes human narcolepsy raises the possibility that other acquired disorders might also result from loss of hypothalamic neurons. To test this possibility for body weight, mice with selective loss of melanin concentrating hormone (MCH) neurons were generated. MCH was chosen to test because induced mutations of the MCH gene in mice cause hypophagia and leanness. Mice with ablation of MCH neurons were generated using toxin (ataxin-3)-mediated ablation strategy. The mice appeared normal but, after 7 weeks, developed reduced body weight, body length, fat mass, lean mass, and leptin levels. Leanness was characterized by hypophagia and increased energy expenditure. To study the role of MCH neurons on obesity secondary to leptin deficiency, we generated mice deficient in both ob gene product (leptin) and MCH neurons. Absence of MCH neurons in ob/ob mice improved obesity, diabetes, and hepatic steatosis, suggesting that MCH neurons are important mediators of the response to leptin deficiency. These data show that loss of MCH neurons can lead to an acquired leanness. This has implications for the pathogenesis of acquired changes of body weight and might be considered in clinical settings characterized by substantial weight changes later in life.

Key words: MCH; ataxin; leanness; ablation; body weight; energy homeostasis

Introduction

In this study, we set out to test whether the loss of a specific hypothalamic cell population could lead to a late onset change in body weight by selectively ablating melanin concentrating hormone (MCH) neurons in the lateral hypothalamus. MCH, a neuropeptide expressed in the lateral hypothalamus and zona incerta, plays an important role in complex motivated behaviors, including the regulation of energy homeostasis (Qu et al., 1996; DiLeone et al., 2003; Verret et al., 2003). The diverse functions of MCH are mediated by two G-protein-coupled receptors, MCH-R1 and MCH-R2 (Chambers et al., 1999; Hill et al., 2001). MCH neuropeptides are derived from a larger prohormone precursor, which also encodes other putative neuropeptides including EI, neuropeptide GE, MCH gene overprinted polypeptide, and antisense RNA overlapping MCH (Bittencourt et al., 1992; Toumaniantz et al., 1996; Borsu et al., 2000). The biological role of these additional neuropeptides is not fully established.

Several studies have established MCH as an important regulator of feeding behavior. It has been shown that the expression level of MCH mRNA increases in leptin-deficient obese (ob/ob) mice as well as in response to fasting in wild-type mice (Qu et al., 1996). It is also been shown that intracerebroventricular administration of MCH causes a rapid increase in food intake (hypophagia) and obesity (Gomori et al., 2003). Genetic models further confirm the role of MCH in energy homeostasis. Mice lacking the MCH1-R are lean, hyperactive, hyperphagic, and hypermetabolic (Chen et al., 2002). The ablation of the prepro-MCH gene lead to leanness in wild-type mice and an attenuation of the obese phenotype in leptin-deficient ob/ob mice (Shimada et al., 1998; Segal-Lieberman et al., 2003), whereas the mice that overexpressed the prepro-MCH gene exhibited obesity and insulin resistance (Ludwig et al., 2001).

The cellular and axonal distribution of the MCH system is also consistent with a role in energy homeostasis regulation. MCH is predominantly expressed in the lateral hypothalamic area, a region implicated in feeding behavior (Guan et al., 2002). MCH neurons make monosynaptic projections throughout the brain to a wide variety of structures that are known to participate in feeding (Zamir et al., 1986). MCH receptors are also expressed in the hypothalamic areas implicated in the control of energy homeostasis such as the ventromedial, dorsomedial, and arcuate nuclei (Kokkotou et al., 2001).

Although mutations in the MCH gene alter body weight, these studies have not allowed an assessment of the broader role of MCH neurons. In addition to MCH, the neurons also express classical neurotransmitters and potentially other neuropeptides. To ablate these neurons, we used a toxin-mediated genetic cell ablation strategy using a truncated Machado-Joseph disease gene product (ataxin-3) under the control of an MCH promoter. The ataxin-3 transgene has been used to induce apoptosis in cells in vitro and for the ablation of orexin neurons in vivo (Yoshizawa et
al., 2000; Hará et al., 2001). As described below, the ablation of MCH neurons results in significant weight loss developing later in life.

Materials and Methods

Animal care. All mice were weaned at 3–4 weeks of age, genotyped, and housed according to gender. Mice were exposed to a standard 12 h light/dark cycle in a temperature-controlled room and fed ad libitum. C57BL/6J (+/+) signal and FVB/n mice were purchased from The Jackson Laboratory (Bar Harbor, ME). Genotypes were determined by the PCR amplification of genomic DNA. All procedures were in accordance with the guidelines set forth by The Rockefeller University Laboratory Animal Research Center.

Generation of MCH/ataxin-3 transgenic mice. The MCH/ataxin-3 transgenic mice were created using bacterial artificial chromosome (BAC) transgenic technology (Yang et al., 1997). The MCH/ataxin-3 construct contained an ataxin-3 transgene under the control of a mouse MCH promoter and the N terminus of the MCH gene. Dr. M. Yanagisawa (University of Texas, Dallas, TX) provided us with an ataxin-3 transgenic cDNA. The ataxin-3 transgene is a cDNA fragment of human ataxin-3 gene with 77 polyglutimine repeats that encodes from amino acid 286 to the C terminus. The human ataxin-3 gene was isolated from a patient with Machado-Joseph disease (Paulson et al., 1997). The N terminus of the ataxin-3 transgene contained hemagglutinin (HA) epitope. The C terminus of the transgene contained a myelocytomatosis oncogene (MYC) epitope (Invitrogen, San Diego, CA) that was followed by a murine protamine 1-gene fragment (mPrm1). The mPrm1 fragment (which included part of exon 1, intron 1, and exon 2) provided an intron and polyadenylation signal (Hara et al., 2001). The ataxin-3 transgene was introduced to the ATG site of the MCH BAC using a homologous recombination-based method. The modified MCH BAC contains 76 kb of a 5′-MCH sequence, followed by the ataxin-3 transgene. BAC filters (BAC mouse II) were obtained from Genome Systems (St. Louis, MO). We cloned the MCH/ataxin-3 construct into the shuttle vector PSV1 for the BAC modification. The PSV1 shuttle vector has a temperature-sensitive origin of replication, recombination RecA gene (which introduces recombination in a recombination-deficient Escherichia coli host), and tetracycline resistant gene (Tet+). The shuttle vector has 76 kb upstream and 97 kb downstream arms of MCH sequence flanking the ataxin-3 on both sides. The shuttle vector was transformed into a DH10B E. coli host, which harbored the MCH BAC. After two homologous recombination events, the modified MCH BAC was selected based on temperature and antibiotic sensitivity. The homologous recombination of the modified BAC was confirmed through Southern blot analysis. The transgenic founder lines were generated by microinjecting BAC DNA purified with a cesium chloride gradient into the pronucleus of fertilized oocytes from an FVB/n strain using the Gene Expression Vessel System Atlas BAC Transgenic Project at the Rockefeller University transgenic facility. Founder animals were mated with FVB/n mice to generate F1 mice, which were used for the characterization. The presence and copy number of the transgene was determined by Southern blot and PCR analysis of the tail genomic DNA. Five transgenic lines were established. To explore the role of MCH neurons in an ob/ob phenotype, we generated a mouse model deficient in both leptin and MCH neurons. We intercrossed C57BL/6/J ob/+ (The Jackson Laboratory) and FVB/n MCH/ataxin-3 mice, which generated MCH/Ataxin-3/ob/+ (F1) mice with a mixed C57BL/6J and FVB/N genetic background. We also intercrossed MCH/Ataxin-3/ob/+ mice to generate MCH/ataxin-3/ob/ob (F2) mice. The ob gene was genotyped by PCR as described previously (Erickson et al., 1996).

Immunohistochemistry. Animals were anesthetized with isoflurane and perfused with 4% paraformaldehyde (PFA) in a PBS solution. Brains were postfixed in a 4% PFA solution for 24 h. Brain sections (50 μm) were obtained using a vibratome. Sections were incubated with a blocking solution (2% normal goat serum, 3% bovine serum, and 0.25% triton in PBS) for 1 h at room temperature. Then, the sections were incubated overnight at 4°C with primary antibodies. For the detection of MCH neurons, we used rabbit anti-MCH antibodies provided to us by Dr. T. Maratos-Flier (Beth Israel Deaconess Medical Center, Boston, MA). For the detection of orexin neurons, we used rabbit anti-Hypocretin antibodies purchased from Oncogene Research Products (Cambridge, MA). For the detection of glia cells, we used rabbit anti-glial fibrillary acidic protein (GFAP) antibodies provided to us by Sigma (St. Louis, MO). For the double label immunohistochemistry (IHC), sections were incubated overnight with primary rabbit anti-MCH and goat anti-MYC (purchased from Santa Cruz Biotechnology, Santa Cruz, CA) antibodies in a blocking solution. Sections were rinsed three times with PBS for 10 min each and incubated with a secondary tetramethylrhodamine isothiocyanate-conjugated goat anti-rabbit antibody and FITC-conjugated donkey anti-goat (purchased from Jackson ImmunoResearch, West Grove, PA) in a blocking solution for 1 h at room temperature. The slides were examined under a Zeiss (Thornwood, NY) Axiosplan fluorescence microscope with Zeiss AxioVision software. The software helped operate the microscope and acquire the sectional images.

Northern blot analysis. We used total hypothalamic RNA (10 μg per lane) for the Northern blot analysis. Following the instructions of the manufacturer, we used Trizol (Invitrogen) for RNA extractions. The RNA samples were fractionated by an agarose-formaldehyde gel electrophoresis and then transferred to nylon membranes using capillary transfer (Sambrook et al., 1989). The filters were incubated at 65°C in Ultra-hyb (purchased from Ambion, Austin, TX) with a 1-mega count/min/ml 32P-labeled DNA probe, which was amplified by PCR. Autoradiography was performed on the filters after they were washed with a high-stringency solution.

Body weight, food intake, and body composition analysis. Body weights were measured once a week for group-housed mice on a regular diet. Food intake was measured in single caged mice after a 1 week period of acclimation. Body composition was used to determine total fat, lean, and water mass (Halaas et al., 1995). Carcasses were weighed and then oven dried in a 90°C oven until constant mass was achieved. Body water weight was determined by subtracting the differences in the weight of the carcasses before and after drying. The dried body mass was ground in a blender. Then, duplicate aliquots were extracted with a 3:1 mixture of chloroform to methanol using Soxhlet apparatus.

Metabolic rate and motor activity measurements. Metabolic rates and spontaneous motor activity was measured by using Oxymax and OptoM3 systems purchased from Columbus Instruments (Columbus, OH). Animals were housed in calorimeter chambers individually and subjected to a 12 h light/dark cycle. Food and water were available in the chambers. Recording began 1 d after the mice were introduced to the chambers to give them time to acclimate. Consecutive IR beam breaks were scored as an ambulatory count. Cumulative ambulatory activity counts were monitored every minute for each IR beam break in three positions, specifically, x, y, and z for both horizontal and vertical 24 h periods. Metabolic rates were calculated using a correction for weight.

Subcutaneous leptin treatment. Two weeks before the mice were treated with leptin, they were individually caged and had their food intake measured daily to establish an average baseline. Alzet 2002 mini-osmotic pumps with an exchange rate of 0.5 μl/h (Alza, Cupertino, CA) were filled with either a PBS solution (control group) or leptin (Amen, Thousand Oaks, CA) at a concentration of 500 ng/μl (transgenic and wild-type leptin-treated groups) and implanted subcutaneously while under anesthesia. Food intake and weight were measured on a daily basis. After 9 d, the osmotic pumps were removed, and food intake and body weight were measured daily for an additional 8 d.

Neuroendocrine function. Blood was collected from the mice using tubes containing EDTA. Plasma was collected after centrifugation and then stored at −20°C for additional analysis. Leptin levels were measured using an ELISA kit from R & D Systems (Minneapolis, MN). Insulin levels were determined by using an ELISA kit from Alpco Diagnostics (Windham, NH). Corticosterone was measured using an RIA kit from R & D Systems. Glucose concentrations were measured using a glucometer from Lifescan (Milpitas, CA). Growth hormone (GH) and insulin-like growth factor (IGF) levels were determined by using an ELISA and EIA kit from Diagnostic Systems Laboratories (Webster, TX).

Taqman real-time PCR analysis. To measure hypothalamic neuropeptide expression and Stearoyl-CoA desaturase (SCD-1) mRNA levels, we
used Taqman real-time PCR analysis. Total RNA was extracted using Trizol (Invitrogen). A reverse transcription kit (Applied Biosystems, Foster city CA) was used for the synthesis of cDNA. Neuropeptide levels were quantified using a Taqman real-time PCR system from Applied Biosystems. All expression data were normalized to cyclophilin expression levels using the same sample. Primers and probes were custom purchased from Applied Biosystems, for measuring agouti-related protein (AGRP), neuropeptide Y (NPY), MCH, proopiomelanocortin (POMC), and SCD-1 mRNA levels.

Statistical analysis. All values are reported as a mean ± SEM. Statistical tests were performed using an unpaired Student’s t test. Statistical analysis of more than three groups was analyzed using ANOVA.

Results
The ataxin-3 transgene is selectively expressed in neurons containing MCH
MCH/ataxin-3 BAC transgenic mice were generated by inserting the cytotoxic ataxin-3 transgene in place of the MCH coding sequence in an MCH BAC transgenic mouse (Yang et al., 1997). The modified MCH BAC contained 76 kb of 5’ mouse MCH sequence, followed by an ataxin-3 cassette and 97 kb of a 3’ mouse MCH sequence. The ataxin-3 transgene included a cDNA fragment of human ataxin-3 gene with 77 polyglutamine repeats, an N-terminal HA epitope, and a C-terminal MYC epitope followed by a murine protamine 1-gene fragment (mPrm1) (Fig. 1A). The modified BAC DNA construct was microinjected into the pronucleus of fertilized oocytes from FVB/n strain to generate five transgenic founder lines. Founder animals were mated with wild-type FVB/n mice to generate F1 transgenic mice.

The expression of an ataxin-3 transgene in MCH neurons was assessed using double immunohistochemistry. Brain sections from 4-week-old MCH/ataxin-3 mice were double stained with anti-MYC and anti-MCH antibodies. This analysis showed colocalization of the MYC epitope with the MCH-expressing neurons in the lateral hypothalamus (Fig. 1B). These neurons contained nuclear aggregates that were positively stained by a MYC antibody. The MYC expression pattern was similar in all transgenic lines, and there was no ectopic expression of MYC immunoreactivity. These results confirmed that the ataxin-3 transgene was specifically expressed in the MCH neurons.

Temporal loss of MCH neurons in the hypothalamus of MCH/ataxin-3 transgenic mice
IHC was next performed at different ages to determine whether there was a loss of MCH neurons in the hypothalamus of MCH/ataxin-3 transgenic mice. Brain sections from MCH/ataxin-3 mice were prepared at ages 2, 4, 8, 12, 15, and 24 weeks and immunostained using MCH and MYC antibodies revealing a time-dependent loss of MCH neurons. The expression of ataxin-3 was observed in the brain sections starting with the first postnatal week (data not shown). The ablation of MCH neurons was first evident during the fourth postnatal week. During the eighth postnatal week, the mice had lost an average of 60% of their MCH neurons. By week 15, 65% of the MCH neurons had disappeared (Fig. 2A). All of the transgenic lines had the same temporal pattern of MCH loss. We further confirmed the loss of MCH neurons by measuring the expression of MCH mRNA in the hypothalamus by Northern blot analysis. There was a marked reduction of MCH mRNA levels in the hypothalamus of 8- and 36-week-old MCH/ataxin-3 mice (Fig. 2B). During postnatal week 19, Taqman Real-Time PCR analysis showed that the expression of MCH mRNA decreased by 68 ± 2.8% when compared with wild-type littermates (data not shown). These data indicated that the expression of the ataxin-3 transgene re-

Figure 1. Specific expression of ataxin-3 transgene in MCH-containing neurons in MCH/ataxin-3 transgenic mice. A, Schematic of a modified MCH/ataxin-3 BAC construct. A modified MCH BAC contains 76 kb of a 5’ MCH sequence, followed by an ataxin-3 transgene. The transgene contains HA epitope, a cDNA fragment starting at amino acid 286 and ending at the C terminus of human mutant ataxin-3, and MYC epitope followed by a murine protamine 1-gene fragment (mPrm1). B, Double immunohistochemistry showing colocalization of MYC epitope (green fluorescence) and MCH (red fluorescence) in the lateral hypothalamic area (LH) of a 4-week-old MCH/ataxin-3 transgenic mouse. Bottom, Bright yellow indicates the presence of nuclear aggregates stained by anti-MYC antibody with MCH-containing neurons. Scale bar, 100 μm.
resulted in a time-dependant elimination of a majority of the MCH neurons in the lateral hypothalamus.

**Normal expression of orexin and GFAP in MCH/ataxin-3 transgenic mice**

There were no gross alterations in the appearance of 24-week-old hematoxylin-eosin-stained sections of hypothalamus from MCH/ataxin-3 mice, an age at which there has already been maximal loss of the MCH neurons (data not shown). To rule out the possibility of nonspecific cell loss in other lateral hypothalamic cell populations, we also examined the level of orexin, a marker for another neuronal cell type in the lateral hypothalamus and GFAP, a glial marker. MCH and orexin are expressed in distinct neuronal populations in the lateral hypothalamus, and both have been implicated in the regulation of feeding behavior (Guan et al., 2002). We used orexin antibodies to immunostain the brain sections of 8- and 15-week-old MCH/ataxin-3 mice and their wild-type littermates. The number of orexin-containing neurons was unchanged in MCH/ataxin-3 when compared with wild-type littermates (Fig. 2C). We also used Northern blot analysis to measure the orexin mRNA level in the hypothalamus of 8- and 36-week-old MCH/ataxin-3 mice. We found no change in the expression level of orexin mRNA in the same samples in which MCH mRNA levels were significantly reduced (Fig. 2B). IHC was also performed using antibodies to GFAP on brain sections of 15-week-old MCH/ataxin-3 mice, confirming that there was no evidence of gliosis in the hypothalamus of MCH/ataxin-3 mice (data not shown). These data suggest that the effect of the MCH/ataxin-3 transgene was limited to the loss of MCH neurons.

**Ablation of the MCH neurons in mice leads to a reduction in body weight, fat mass, and body length**

MCH/ataxin-3 mice were born in mendelian ratios. The mice were viable, fertile, and appeared grossly normal after physical inspection. The weight of these animals was no different than that of the controls until 7 weeks of age. At 7 weeks of age, MCH/ataxin-3 mice diverged from control animals and showed a significant reduction in body weight when compared with their wild-type littersmates (20.48 ± 0.5 g vs 22.56 ± 0.5 g, p < 0.05 for males; 17.18 ± 0.56 vs 20.37 ± 0.6 g, p < 0.05 for females). By 22 weeks of age, weight was reduced by 14% in males and 13% in females (Fig. 3A,B). Body composition analysis revealed that MCH-ataxin-3 mice had 60% less fat mass than their wild-type littersmates. The fat mass in 18-week-old MCH/ataxin-3 females was 1.2 ± 0.2 g (with 5.6% total body mass), compared with 3 ± 0.2 g (with 11.9% total body mass) in wild-type littermates (Fig. 3C). MCH/ataxin-3 mice also exhibited a small but significant reduction of body length, which was first apparent at 14 weeks of age. At 20 weeks of age, body length
was reduced by 4.5% in females (9.4 ± 0.08 vs 9 ± 0.01 cm; p < 0.002) and 6% in males (9.86 ± 0.03 vs 9.3 ± 0.01 cm; p < 0.002) (Fig. 3D). Consistent with this, there was a significant decrease in the percentage of lean mass. MCH/ataxin-3 mice had 7% less lean mass than their wild-type littermates. The lean body mass for 13-week-old males was 7.1 ± 0.1 g (with 32% of total weight), whereas it was 7.7 ± 0.1 g (with 30% of total weight) for their wild-type littermates (Fig. 3C). The lean body mass for 13-week-old MCH/ataxin-3 females was 6.1 ± 0.1 g (with 27% of total weight), whereas it was 7.0 ± 0.14 g (with 27% of total weight) for their wild-type littermates (Fig. 3C).

Mice lacking the MCH neurons exhibited a negative energy balance

Food intake, locomotor activity, and metabolic rate were measured over 24 h periods. Total and resting oxygen consumption and carbon dioxide production were measured using the Oxymax system. The food intake of MCH/ataxin-3 mice was not different than that of the controls at 8 weeks of age (Fig. 4A). At 16 weeks of age, female MCH/ataxin-3 mice consumed 13% less food than their wild-type littermates over a 24 h period (4 ± 0.2 vs 4.6 ± 0.2 g/d; p < 0.03) (Fig. 4A). Male mice consumed 6.7% less food than wild-type littermates (4.5 ± 0.05 vs 4.8 ± 0.1 g/d; p < 0.05) (Fig. 4A). We did not see a decrease in food intake when our measurements were corrected for body weight. Female MCH/ataxin-3 mice consumed as much as their wild-type littermates over a 24 h period (0.200886 ± 0.010796 vs 0.204048 ± 0.007986 g/kg/d; p > 0.05). Male mice also consumed as much as their wild-type littermates (0.19 ± 0.006 vs 0.192 ± 0.012 g/kg/d; p > 0.05). The mice had a normal diurnal feeding pattern consuming most of their calories during the dark cycle. The reduction in food intake was observed during both dark and light cycles (data not shown).

The effects of the ablation of MCH neurons on energy expenditure were sexually dimorphic with significant effects evident in males but not females. Male MCH/ataxin-3 mice showed an increase of 27.6% in total and 26.1% in resting VO2 (Fig. 4B) [total 4473.6 ± 300.6 ml/kg/hr in MCH/ataxin-3 mice vs 3507.1 ± 119.6 ml/kg/hr in littermates (p < 0.009); resting VO2, 4009.2 ± 171.8 ml/kg/hr in MCH/ataxin-3 mice vs 3178.6 ± 146.7 ml/kg/hr in littermates (p < 0.004)]. Carbon dioxide production (VCO2) was increased by 20.3% in the male mice (4238.8 ± 251.8 ml/kg/hr in MCH mice and 3524.2 ± 542.8 ml/kg/hr in their wild-type littermates; p < 0.05). Locomotor activity was equivalent in both males and females. Together, these data indicate that hypophagia contributes to the leanness of MCH/ataxin-3 with an added effect in males secondary to an increase in energy expenditure.

Neuroendocrine function

Plasma leptin levels were reduced by 30% in 19-week-old females compared with their wild-type littermates (1.9 ± 0.07 vs 2.7 ± 0.34 ng/ml; p < 0.05; n = 6). However, this difference was no longer evident in 28-week-old females (data not shown). Levels of glucose were not significantly different between 19-week-old females and their wild-type littermates, whereas 28-week-old males had 20% reduction in glucose levels. Female 19-week-old mice had a 20% (0.39 ± 0.01 ng ml⁻¹) reduction in insulin levels when compared with their wild-type littermates (0.49 ± 0.01 ng ml⁻¹; p < 0.05; n = 6). Corticosterone levels were unchanged in 19-week-old mice (data not shown). Our analysis shows that MCH/ataxin-3 mice are hypoleptinemic, which is consistent with their lean phenotype. The reduced insulin levels also suggest that the loss of MCH neurons may lead to a slight increase in the insulin sensitivity of the mouse.

Plasma levels of GH and insulin-like growth factors (IGF-1) were not significantly different between MCH/ataxin-3 mice and their wild-type littermates. We found that IGF-I levels in 13-week-old females and males were 1083 ± 107 and 1331.2 ± 274.8 ng/ml, whereas the levels for their wild-type littermates were 1007 ± 115 and 1156.19 ± 166 ng/ml. We also found that GH levels in 13-week-old females and males were 12.2 ± 1.6 and 14.1 ± 3 ng/ml, whereas their wild-type littermates were 9.6 ± 0.45 and 10.9 ± 1.4 ng/ml. Hypoleptinemia generally leads to an increase in NPY and AGRP RNA levels and a reduction of POMC RNA levels in the hypothalamus (Hillebrand et al., 2002). However, this was not the case in the MCH/ataxin-3 knock-out mice in which there were no significant changes in the expression of the RNAs for
these neuropeptides, in 19-week-old female MCH/ataxin-3 mice (Fig. 5).

MCH/ataxin-3 mice exhibited a normal response to fasting, leptin treatment, and deficiency

We next assessed the response of MCH/ataxin-3 mice to fasting and acute and chronic leptin deficiency, first in studies of food intake and body weight in mice before and after fasting for 24 h. Weight loss after a fast was similar in both groups. When provided with food after 24 h of fasting, younger 19-week-old MCH/ataxin-3 mice initially consumed less food than controls during the first 3 and 6 h (2.1 ± 0.1 vs 2.83 ± 0.2 g; p < 0.007). However, by 24 h, the animals had caught up and food intake was unchanged between the groups after 24 h (6.4 ± 0.08 vs 6.6 ± 0.19 g) (Fig. 6A). This reduction of food intake at early times after fasting was no longer evident in older 24-week-old animals. In both groups, weight gain following a fast was also equivalent. These data demonstrate that although there is an initially sluggish response to fasting, MCH/ataxin-3 mice exhibited a normal compensatory hyperphagic response to fasting.

Leptin has been suggested to modulate the activity of the MCH system (Huang et al., 1999). To investigate the response of MCH/ataxin-3 mice to leptin, the effect of leptin treatment on food intake and body weight in 19-week-old MCH/ataxin-3 male was determined. MCH/ataxin-3 mice responded with equivalent reductions in food intake and body weight as controls (45% reduction of food intake at day 2 and a 13% decrease of body weight at day 9) (Fig. 6B, C). In addition, using a protocol that leads to acute leptin deficiency, the hyperphagic response to leptin withdrawal was also normal in the MCH/ataxin-3 animals. In these experiments, we treated the mice with high dose subcutaneous leptin followed by abrupt leptin withdrawal (Montez et al., 2004). This protocol leads to a state of acute leptin deficiency that is followed by a robust and predictable period of hyperphagia. Four days after leptin withdrawal, MCH/ataxin-3 mice showed a 35.8 ± 4.5% increase in food intake, whereas wild-type mice consumed 33.4 ± 5.2% more food when compared with the baseline (Fig. 6B). The baseline was the average food consumption for 24 h measured over a 3 day period before leptin treatment. Weight gain was also equivalent in both groups when compared with the baseline (Fig. 6C). The baseline was the body weight measured for 2 d before the leptin treatment.

The loss of MCH neurons lead to the attenuation of the ob/ob phenotype

We next assessed whether, similar to the effects of an MCH mutation, the ablation of MCH neurons could also attenuate an obese phenotype. Despite the fact that the biologic response to starvation, leptin treatment, and acute leptin deficiency was unchanged, the ablation of MCH neurons did lead to a significant alteration in the response to congenital leptin deficiency in ob/ob mice. C57BL/6J ob/+ and MCH/ataxin-3 mice were intercrossed to generate MCH/ataxin-3/ob/ob mice. The MCH/ataxin-3/ob/ob mice showed a significant reduction in body weight when compared with ob/ob mice. At 24–28 weeks of age, weight was
reduced by 13% in females (60 ± 2.1 g vs 68 ± 1.3 g; p < 0.004) (Fig. 7A). Mice lacking both leptin and MCH neurons also exhibited a significant reduction in plasma glucose concentration (226 ± 46 vs 414 ± 61 mg/dl; p < 0.02) (Fig. 7B) and no change in insulin levels. There was also a marked improvement in hepatomegaly and steatosis in the ob/ob mice lacking MCH neurons (average liver weight, 2.6 g for MCH/ataxin-3 ob/ob mice vs 4.4 g in ob/ob animals vs 1 g for wild-type mice (Fig. 7C). In histologic sections, the steatosis characteristic of ob/ob mice was normalized by ablation of MCH neurons (Fig. 7D). In addition, the hepatic SCD-1 gene expression was decreased by 34% in MCH/ataxin-3/ob/ob mice when compared with ob/ob (data not shown).

There was also an improvement in fertility for both MCH/ataxin-3/ob/ob and ob/ob control mice. All mice had a mixed C57BL/6J and FVB/N genetic background. Fertile male mice were comprised of 12.5% of the ob/ob and 12.5% of the MCH/ob/ob, whereas female fertile mice were comprised of 33% of the ob/ob and 30% of the MCH/ob/ob. The fertile females produced normal size litters but were unable to nurture their pups through weaning. These data indicate that the ablation of MCH neurons can also reduce an already developed obese phenotype in adult animals.

Discussion
In this study, we generated mice with selective loss of MCH neurons using a toxin (ataxin-3)-mediated ablation strategy. We showed that the selective loss of the MCH neurons leads to an acquired, late onset reduction of body weight. These results suggest the possibility that late onset changes in body weight could in some cases be the result of the loss of specific neural populations in the hypothalamus.

Obesity is a complex multifactorial condition influenced by genetic and environmental factors. Genetic factors account for a substantial fraction of the variance in body weight. However, in some instances, substantial changes in body weight, in either direction, develop later in life. Although it is possible that these alterations have a genetic component, these changes could also be caused by environmental factors acting on susceptible genetic factors. One theoretical possibility for changes in weight later in life is that key neurons that regulate food intake and body weight are lost. In humans, it is known that hypothalamic lesions resulting from tumors, sarcoid, or viral infections

Figure 6. MCH/ataxin-3 mice exhibited a normal response to fasting, leptin treatment, and leptin deficiency. A, Cumulative food intake in response to 24 h starvation in 19-week-old male MCH/ataxin-3 mice (open squares; n = 5) and their wild-type littermates (wt; filled diamonds; n = 6). *p < 0.03, **p < 0.007. B, C, Food intake (B) and body weight (C) in response to 9 d of leptin treatment followed by 8 d of leptin withdrawal in 19-week-old male MCH/ataxin-3 mice (open squares; n = 6) and their wild-type littermates (filled diamonds; n = 6) compared with PBS control mice (filled triangles; n = 4). Food intake and body weight are represented as a percentage of the baseline. The food intake baseline was an average of the food consumption for 24 h measured over a 3 d period before leptin treatment. The body weight baseline was body weight measured 2 d before leptin treatment. Tg, Transgenic.

Figure 7. The loss of MCH neurons in ob/ob mice improved obesity, diabetes, and hepatic abnormalities. A, Body weight in 24- to 28-week-old female MCH/ataxin-3/ob/ob (filled bar; n = 7) and ob/ob control (downward diagonal filled bar; n = 10) mice. *p < 0.004. B, Glucose levels in 24- to 28-week-old female MCH/ataxin-3/ob/ob (filled bar; n = 10), ob/ob control mice (downward diagonal filled bar; n = 8), and wild-type mice (wt; dot-filled bar; n = 6). *p < 0.03, **p < 0.002. C, Photographs of freshly dissected livers from 24-week-old female MCH/ataxin-3/ob/ob (center), ob/ob control (left), and wild-type (right) mice. D, Hematoxylin and eosin staining of liver sections from female MCH/ataxin-3/ob/ob (right) and ob/ob control (left) (females, n = 3; males, n = 2). Scale bar, 20 μm.
However, these lesions are generally associated with easily discernible histologic changes. It is also possible that acquired changes in weight could result from the selective loss of specific classes of neurons without their being any evident gross histologic changes. This theoretical possibility is illustrated by the recent realization that sporadic cases of narcolepsy are the result of the spontaneous loss of orexin neurons in the lateral hypothalamus later in life (Hara et al., 2001). Although the pathogenesis of the neural loss for this condition is unknown, it has been suggested that it could be attributed to undefined environmental factors that act on a susceptible genetic background. One of the predisposing factors is associated with HLA, suggesting that an autoimmune component might be the cause of the selective destruction of orexin neurons (Carlander et al., 2001; Smith et al., 2004). These seminal findings in narcolepsy research suggest that the loss of different neuronal populations in hypothalamus could lead to physiologic alterations in other biologic functions controlled by this brain region. Additional studies will be necessary to determine the contributions of environmental factors and their interaction with genetic risk factors in the pathogenesis of weight changes later in life.

In this study, we ablated MCH neurons to test the possibility that specific neural loss could lead to late onset alterations in the homeostatic control of energy balance. MCH-expressing neurons are localized exclusively in the lateral hypothalamus in which they are found in a nonoverlapping distribution with orexin (Guan et al., 2002). MCH is a potent orexigenic mediator and a key central regulator of energy homeostasis (Qu et al., 1996). Mutations in the MCH gene reduce food intake and body weight, making the expressing neurons attractive candidates for testing the possibility that the loss of this population might be associated with the development of an extremely lean phenotype later in life without changing viability. Here, we show that mice lacking MCH neurons are viable but develop a late-onset nutritional disorder characterized by extreme leanness.

Although traditional methods, such as diphtheria toxin gene product and herpes simplex virus 1 thymidine kinase, have been used in the past for cell ablation, they are not very effective when ablating neural postmitotic cells. Therefore, MCH neurons were ablated using a toxin-mediated cell ablation strategy using the human ataxin-3 gene. This mutant protein has an expanded number of polyglutamine repeats, which leads to the toxic cell death of neurons that express it (Paulson et al., 1997; Beivinio and Loll, 2001). The ataxin-3 transgene has been used before for the ablation of orexin neurons in vivo (Hara et al., 2001). We generated MCH/ataxin-3 transgenic mice by introducing an ataxin-3 cassette into the initiation codon of the MCH gene via homologous recombination in a bacterial strain carrying an MCH bacterial artificial chromosome (Yang et al., 1997). MCH cell-specific expression of the ataxin-3 transgene specifically in MCH neurons was confirmed in multiple different transgenic lines suggesting that the 76 kb of 5′ and 97 kb of 3′ sequences in this BAC are sufficient to confer correct expression of the MCH gene. This BAC can now be used to direct the expression of other proteins or reporters to these neurons with a high likelihood of correct expression.

The phenotype of the MCH/ataxin-3 mice was remarkable similar to that of mice with induced mutations of the MCH gene (Shimada et al., 1998). In both cases, the mice exhibited reduced body weight, fat mass, and leptin levels with reduced food intake and an increase of energy expenditure. It has also been shown that the ablation of MCH1-R (the receptor for MCH) lead to an increase in locomotor activity (Chen et al., 2002). If anything, it is likely that the MCH/ataxin-3 mice have slightly less severe phenotypes, because the neuronal ablation was not complete. Although the MCH neurons are likely to express classical neurotransmitters and potentially other neuropeptides, the similarity of the phenotypes suggest that MCH itself is a major mediator of the physiologic actions of these neurons. The only notable difference between the MCH knock-out mice and the MCH/ataxin-3 mice was in the linear growth in which the ablation of MCH neurons resulted in a mild decrease in skeletal growth and lean mass. It is possible that this is the result of actions of other neuropeptide systems in these neurons on the somatotropic axis, including GH, IGF-I, IGF binding proteins, and IGF receptors, which is regulated by nutritional state and endocrine mechanisms (Bornfeldt et al., 1989; Renaville et al., 2002). However, our data showed that the levels of GH and IGF-I were not significantly different between MCH/ataxin-3 mice and their wild-type littermates. Additional studies are needed to elucidate the mechanisms leading to the decreased skeletal growth.

Numerous previous studies have confirmed that leptin acts as a positive regulator of POMC and CART while acting as a negative regulator of AGRP and NPY (Elias et al., 1998; Hillebrand et al., 2002). Consistent with this, POMC gene expression is reduced in the hypothalamus of MCH knock-out mice. In contrast, the levels of POMC and AGRP RNA were not altered in the hypothalami of the MCH/ataxin-3 mice despite the fact that the leptin levels were reduced (Shimada et al., 1998). This result could mean that MCH neurons normally modulate POMC gene expression in an MCH-independent manner. Bidirectional communication has been demonstrated between MCH-expressing neurons in the lateral hypothalamus and the arcuate-expressing neurons, such as POMC, NPY, and AGRP (Elias et al., 1998).

Both the MCH knock-out and ablation of MCH neurons significantly reduce the obesity associated with leptin mutations (Segal-Lieberman et al., 2003). Thus, MCH neurons play an important role in mediating the orexigenic response to chronic leptin deficiency. In contrast, the ablation of these neurons does not appreciably alter the biologic response to fasting or acute leptin deficiency (induced using a leptin withdrawal protocol), both of which are normal in the MCH/ataxin-3 mice. The basis for this difference is not clear. It may be that acute changes in food intake or body weight activate multiple, potentially independent pathways, whereas chronic leptin deficiency activates fewer such pathways. It is noteworthy, however, that neither the ablation of MCH neurons nor MCH mutations completely normalize the weight of ob mice, suggesting that other pathways also play a role in the response to chronic leptin deficiency (Erickson et al., 1996).

The toxin-mediated ablation approach has also been used in the ablation of orexin neurons (Hara et al., 2001). Both MCH and orexin are expressed in separate neuronal populations within the lateral hypothalamus area and have been implicated in feeding regulation (Hara et al., 2001; Guan et al., 2002). Orexin neurons have been linked to arousal and sleep regulation. The roles of MCH in feeding regulation are different from those of orexin. Mice lacking orexin neurons exhibited a late-onset obesity despite their reduced food intake. Their obesity was characterized by reduced energy expenditure because of a decrease in motor activity and metabolic rate. Although mice lacking MCH neurons, exhibited a late onset leanness because of hypophagia and an increased metabolic rate. These differences suggest that MCH and orexin may play different roles in energy metabolism regulation.

In summary, MCH/ataxin-3 mice develop a late onset syndrome characterized by leanness, hypophagia, and, in males, increased energy expenditure without any obvious changes in the
Bevvino AE, Loll PJ (2001) An expanded glutamine repeat destabilizes na-
possibly other homeostatic systems later in life.
results suggest that the use of immunohistochemistry to assess
arcuate POMC neurons might result in obesity. Overall, these
raising the possibility that selective loss of arcuate NPY/AGRP
possibility can be easily tested. The loss of other neuronal popu-
could be associated with the loss of the MCH cell population, this
overly speculative to suggest that weight loss associated with an-
classes of neurons, undetectable using standard methods, can
These results thus raise the possibility that selective loss of specific
histologic appearance of the hypothalamus. Indeed, the
basis for the phenotype in these animals would not be evident
gross histologic appearance of the hypothalamus. Indeed, the
results suggest that the use of immunohistochemistry to assess

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