Direct Effects of Leptin on Brown and White Adipose Tissue

Catherine A. Siegrist-Kaiser,* Véronique Pauli,* Cristiana E. Juge-Aubry,* Olivier Boss,† Agnès Pernin,* William W. Chin,* Isabelle Cusin,† François Rohner-Jeanreanu,‡ Albert G. Burger,* Jürgen Zapf,* and Christoph A. Meier*‡‡†‡

*Unité de Thyroïde, Division d’Endocrinologie et Diabétologie, Hôpital Universitaire de Genève, Hôpital Universitaire de Genève, CH-1211 Geneva 14, Switzerland; ‡Biochimie Médicale, Centre Médical Universitaire, Université de Genève, CH-1211 Geneva 14, Switzerland; †Laboratoires des Recherches Métaboliques, Faculté de Médecine, Université de Genève, CH-1211 Geneva 14, Switzerland; ‡Division of Endocrinology and Diabetes, University Hospital Zürich, CH-8091 Zurich, Switzerland; ‡‡Clinique de Médecine II, Département de Médecine, Hôpital Universitaire de Genève, CH-1211 Geneva 14, Switzerland; and †Endocrine Division, Massachusetts General Hospital, Boston, Massachusetts 02114

Abstract

Leptin is thought to exert its actions on energy homeostasis through the long form of the leptin receptor (OB-Rb), which is present in the hypothalamus and in certain peripheral organs, including adipose tissue. In this study, we examined whether leptin has direct effects on the function of brown and white adipose tissue (BAT and WAT, respectively) at the metabolic and molecular levels. The chronic peripheral intravenous administration of leptin in vivo for 4 d resulted in a 1.6-fold increase in the in vivo glucose utilization index of BAT, whereas no significant change was found after intracerebroventricular administration compared with pair-fed control rats, compatible with a direct effect of leptin on BAT. The effect of leptin on WAT fat pads from lean Zucker Fa/fa rats was assessed ex vivo, where a 9- and 16-fold increase in the rate of lipolysis was observed after 2 h of exposure to 0.1 and 10 nM leptin, respectively. In contrast, no increase in lipolysis was observed in the fat pads from obese fa/fa rats, which harbor an inactivating mutation in the OB-Rb. At the level of gene expression, leptin treatment for 24 h increased malic enzyme and lipoprotein lipase RNA 1.8±0.17 and 1.9±0.14-fold, respectively, while aP2 mRNA levels were unaltered in primary cultures of brown adipocytes from lean Fa/fa rats. Importantly, however, no significant effect of leptin was observed on these genes in brown adipocytes from obese fa/fa animals. The presence of OB-Rb receptors in adipose tissue was substantiated by the detection of its transcripts by RT-PCR, and leptin treatment in vivo and in vitro activated the specific STATs implicated in the signaling pathway of the OB-Rb.

Taken together, our data strongly suggest that leptin has direct effects on BAT and WAT, resulting in the activation of the Jak/STAT pathway and the increased expression of certain target genes, which may partially account for the observed increase in glucose utilization and lipolysis in leptin-treated adipose tissue. (J. Clin. Invest. 1997. 100:2858–2864.)

Key words: leptin • adipocytes • lipolysis • Jak/STAT

Introduction

Adipocyte differentiation and metabolism are regulated by a variety of endocrine and paracrine factors, including certain cytokines, such as IL-6 and TNF-α (1–3). The ob gene product (leptin) was initially cloned in ob/ob mice and shown to be a 167-amino acid cytokine-like peptide, secreted by adipocytes in a regulated fashion (4, 5). Subsequent physiological studies revealed that leptin was able to inhibit food intake and enhance energy expenditure by acting at the hypothalamic level, providing a feedback signal reflecting the nutritional status of the periphery (6–8). The gene encoding a leptin receptor (OB-R) was cloned from hypothalamic cDNA, and several differentially expressed splice variants were detected in various other tissues (9–12). While most of these variants are currently thought to be involved in leptin binding and transport, the intracellular domain of the long form of the OB-R (OB-Rb) is homologous to the IL-6/gp130 receptor family (13). Several studies performed in vivo and in vitro demonstrated the capability of the OB-Rb to activate the Janus kinase (Jak)/signal transducers and activators of transcription (STAT) pathway, resulting in the nuclear translocation of specific STATs, such as STAT 1, 3, 5, and 6 (12–16). Although the OB-Rb is highly expressed in the hypothalamus and the lymphoid system, the presence of small amounts of its mRNA was also demonstrated in other organs, including pancreatic β cells and brown and white adipose tissue (BAT and WAT, respectively) (12, 17, 18). Recently, direct functional effects of leptin on some of these nonhypothalamic target organs were described, such as on hematopoietic cells, the endocrine pancreas, pituitary, and ovary (17, 19–23). However, no data are currently available on potential direct paracrine or autocrine effects of leptin on adipose tissue, although several studies have provided some evidence compatible with such an hypothesis (24–28).

In this study, we examined the effect of leptin on brown and white adipocytes in vivo, ex vivo, and in primary cultures to demonstrate direct actions of leptin on metabolism and gene expression. In addition, we provide evidence that leptin administration in vivo and in vitro activates the Jak/STAT pathway in adipocytes, resulting in the nuclear translocation of STAT1.

1. Abbreviations used in this paper: BAT, brown adipose tissue; Jak, Janus kinase; OB-R, leptin receptor; PPAR, peroxisome proliferator-activated receptor; RT-PCR, reverse transcription PCR; STAT, signal transducers and activators of transcription; UCP, uncoupling protein; WAT, white adipose tissue.
Methods

**Measurement of glucose utilization in BAT in vivo.** Freely moving adult male rats were infused for 4 d with vehicle (0.1 M Tris-HCl, pH 9.0) or leptin (12 μg/d) administered into the lateral cerebral ventricle. Another group of rats was given either vehicle (Ringer solution) or 1 mg/d of recombinant mouse leptin (Novartis, Basel, Switzerland) for 4 d into the femoral vein via a custom-made device including a swivel mounted on a counterbalanced arm, supporting the catheters and allowing the animals to move freely. To eliminate the confounding variable of leptin-induced hypophagia, all vehicle-infused rats were pair-fed to the amount of food consumed by leptin-infused animals. At the end of the treatment period, the animals were anesthetized with 60 mg/kg sodium pentobarbital intraperitoneally and prepared for euglycemic-hyperinsulinemic clamps associated with the labeled 2-deoxyglucose technique to measure the insulin-stimulated glucose utilization index in BAT in vivo. This technique was described previously and validated (29–31).

**Measurement of lipolysis in WAT fat pads ex vivo.** Epidyymal fat pads were removed and pooled from Sprague-Dawley (140–160 g) or lean (Fa/fa, 170–190 g) or obese (fa/fa, 205–230 g) Zucker rats, and the amount of glycerol released into the medium was determined as described (32). Briefly, after a preincubation period of 1 h, the pooled fat pads were incubated at 37°C for different time intervals in the presence of Krebs-Ringer-Hepes buffer containing diazylized human serum albumin (30 mg/ml), 2 mg/ml glucose, as well as epinephrine (0.5–5,000 nM) or leptin (0.1–1,000 nM), as appropriate. After the incubation period, glycerol release into the medium was determined and the results were expressed as micromoles of glycerol released per 100 mg of adipose tissue per hour. The fat pads from the leptin-resistant obese fa/fa rats served as negative controls for these experiments.

**Primary culture of BAT and WAT.** 3-wk-old male lean Zucker Fa/fa rats were killed and interscapular BAT as well as inguinal WAT were immediately removed. For both tissues, precursors were isolated as described elsewhere and suspended in a (1:1 vol/vol) mixture of Dulbecco’s modified Eagle’s medium and Ham’s F-12, supplemented with 16 μM bina, 18 μM pantethenic acid, 5 mM glutamine, 16 mM glucose, 15 mM Hepes, penicillin 50 IU/ml, streptomyycin 50 μg/ml, 100 μg ascorbate, and 10% fetal calf serum (complete medium) (33). The medium was changed on day 1 and replaced with a differentiation medium containing 510 mM insulin, 100 mM dexamethasone, 0.2 mM 3,5,3’-triiodothyronine, and 10 μg/ml transferrin in complete medium (34). The medium was changed every 2 d until the cells were fully differentiated after 14 d, as ascertained by the presence of lipid accumulation and the expression of uncoupling protein-1 (UCP-1) mRNA. The brown adipocytes from the leptin-resistant obese fa/fa rats served as negative controls for these experiments.

**Analysis of gene expression in brown adipocytes.** Fully differentiated brown adipocytes were treated for 24 h with 1 μM BRL49653 (kind gift of Dr. W. Wahl, Lausanne, Switzerland), 625 nM leptin, or a combination of both. Total RNA was isolated by guanidinium isothiocyanate extraction (35). Cells were lysed in a 4 M guanidinium thiocyanate-containing buffer containing 5 mM sodium citrate, pH 7.0, 0.5% sacrosyl, and 0.1 M β-mercaptoethanol and ultracentrifuged overnight through a CsCl step gradient. RNA (15 μg/sample) was denatured and subjected to electrophoresis on a 1% agarose-formaldehyde gel before blotting and ultraviolet cross-linking onto a nylon membrane (NYTRAN 13 N; Schleicher & Schuell, Dassel, Germany). The probes for malic enzyme (EcorI insert from mPOE6, kindly provided by Dr. V. Nikodem, Bethesda, MD), lipoprotein lipase (EcorI insert from gPgem2, kindly provided by Dr. M. Schotz, Los Angeles, CA), and aP2 (PstI insert from pAL422, kindly provided by Dr. D. Lane, Baltimore, MD) were labeled with 50 μCi [α-32P]dCTP (Rediprime random primer labeling kit; Amersham, Buchs, Switzerland) to a specific activity of at least 2 × 10⁹ cpm/μg (36, 37). Hybridization was carried out for 2 h with 2 ng/ml of labeled probe, using the Rapid-Hyb buffer (Amersham). Membranes were washed, with the last step being 0.1× SSC, 0.1% SDS at 65°C for 15 min. The autoradiograms were quantified by laser scanning densitometry (Molecular Dynamics, Sunnyvale, CA). The results were corrected for 18S and 28S ribosomal RNA on the gel as quantitated by scanning densitometry.

**Reverse transcription PCR (RT-PCR) assay for the detection of OB-Rb RNA.** RNA was prepared from the hypothalamus, WAT, and BAT, as well as cultured brown and white adipocytes as described above. 5 μg of total RNA was used for reverse transcription with Moloney murine leukemia virus reverse transcriptase (2 × 40 min, 37°C), primed by oligo-dT16. Primers for PCR amplification were synthesized according to the published sequence for the long form of the rat OB-R (OB-Rb, GenBank accession No. D85558), and the cDNA was amplified using the sense primer OB1F (nucleotides 2541–2564: 5’-TAGTGTCCATTGACGGATATTT-3’) and the reverse primer OB1R (nucleotides 2895–2913: 5’-CCAGGAGATGACTGTGT-3’). All PCR reactions were carried out in standard buffer [10 mM KCl, 20 mM Tris-HCl, pH 8.0, 1% Triton X-100, 6 mM (NH₄)₂SO₄, 0.2 mM dNTP, 2 mM MgCl₂, 0.5 μM of each primer, 2.5 U/100 μl of native Pfu (Stratagene, Basel, Switzerland)], for > 40 cycles with the following temperatures: denaturation 30 s at 92°C, annealing 30 s at 50°C, and elongation for 90 s at 72°C. The PCR products were analyzed on a 2% agarose gel and stained with ethidium bromide. The PCR products obtained from WAT and BAT and culture were sequenced by direct sequencing using standard protocols for the ABI373A automated sequencer with primers OB1F and OB1R.

**Preparation of nuclear extracts.** Lean male Fa/fa Zucker rats were injected intravenously with either vehicle or 500 μg of leptin in Tris 0.1 M, pH 8.1. After 60 and 90 min, the animals were killed and BAT was removed. Nuclear extracts were prepared as described elsewhere (15). Briefly, tissues were homogenized in 10 mM KCl, 1.5 mM MgCl₂, 10 mM Hepes, pH 7.9, 1 mM DTT, 1 mM Na₂VO₄, 10 μg/ml leupeptin, 0.3 μg/ml aprotinin, 1 μg/ml pepstatin A, and 1 mM PMSF. The homogenate was centrifuged at 2,000 g for 10 min and the pellet was washed with the same buffer. The washed pellet was resuspended in a buffer containing 420 mM NaCl, 10 mM KCl, 20 mM Hepes, pH 7.9, 20% glycerol, 1 mM DTT, 1 mM Na₂VO₄, and the protease inhibitors specified above. Nuclei were extracted for 30 min at 4°C on a shaking rotor and centrifuged for 10 min at 16,000 g. Supernatants were diluted in a buffer containing 10 mM KCl, 20 mM Hepes, pH 7.9, 20% glycerol, 1 mM DTT, 1 mM Na₂VO₄, and the protease inhibitors specified above. After additional centrifugation, the supernatants were concentrated on a Microcon-50 ultrafiltration column (Amicon, Wallisellen, Switzerland).

Nuclear extracts from primary brown or white adipocyte cultures were prepared as follows (38). Cells were cultured in 6-cm petri dishes as described above. The cells were placed in serum-free medium for 6 h before stimulation with 0.2 μM leptin for 15 or 30 min as indicated. Cells were washed with 2 ml of cold PBS and scraped off the dish in 1 ml of PBS. Two plates were pooled for nuclear protein extraction. Cells were pelleted by centrifugation at 5,200 g for 15 s. The pellet was resuspended in 400 μl of hypotonic buffer (10 mM Hepes, pH 7.9, 10 mM KCl, 0.1 mM EDTA, 0.1 mM EGTA, 1 mM DTT, 1 mM PMSF, and 0.5 mM Na₂VO₄) by gentle pipetting. The cells were left to swell for 15 min on ice, after which 25 μl of 1% NP-40 was added and the tubes were vigorously vortexed. After spinning at 15,800 g for 30 s, the pellet was resuspended by mild vortexing in 50 μl of cold hypotonic buffer (20 mM Hepes, pH 7.9, 0.4 M NaCl, 1 mM EDTA, 1 mM EGTA, 20% glycerol, 1 mM DTT, 1 mM PMSF, and 0.5 mM Na₂VO₄). After 15 min of incubation, the tubes were centrifuged for 5 min at 15,800 g at 4°C. Supernatants were aliquoted and stored at −70°C. The protein concentration in the nuclear extracts was determined by the Bradford assay (BioRad, Glattbrugg, Switzerland).

**Activation of STATs in vivo and in vitro (EMSA).** A double-stranded oligonucleotide (Microsynth, Balgach, Switzerland) containing a binding site (M67-SIE) for STAT1, 3, and 4, was labeled by fill-in with Klenow DNA polymerase in the presence of [α-32P]dATP (39). 3 μg of nuclear extracts from the primary cell cultures, or 5 μg of
nuclear extracts from BAT, was incubated with 25,000 cpm of the labeled M67-SIE, and 2 μg of poly [d(I-C)] in EMSA binding buffer (50 mM KCl, 20 mM Hepes, 20% glycerol, 0.05% NP-40, 10 mM β-mercaptoethanol, pH 7.5) with a final volume of 25 μl. Incubation was performed at room temperature for 20 min and the samples were analyzed on a 5% polyacrylamide gel in 0.5 x TBE (44.5 mM Tris, 44.5 mM boric acid, 1 mM EDTA, pH 8) for 75 min at 300 V. Gels were dried and then autoradiographed.

For the supershift assay, either 0.3 or 0.75 μg of a mouse monoclonal IgG1 anti-STAT1 antibody (Santa Cruz Biotechnology, Basel, Switzerland) or of an anti-STAT3 antibody, respectively, was added to the EMSA incubation mixture for 15 min before the addition of the labeled oligonucleotide.

Statistical analysis. Results are expressed as the mean±SEM and the unpaired Student’s t test was used to assess statistical significance. P > 0.05 was considered not significant (NS).

Others. All experimental animal protocols were approved by the Review Board for Animal Experimentation of the Canton of Geneva, Switzerland.

Results

Leptin enhances glucose utilization of BAT in vivo. To assess direct peripheral effects of leptin on BAT, male rats were given leptin either intracerebroventricularly or intravenously for 4 d, and euglycemic-hyperinsulinemic clamps were performed to determine the insulin-stimulated glucose utilization index. As shown in Fig. 1A, the glucose utilization index in BAT remained unchanged in rats treated with leptin intracerebroventricularly, as compared with pair-fed controls (195±23 vs. 174±11 ng/min/mg tissue, NS). In contrast, when leptin was administered intravenously, the insulin-stimulated glucose utilization index increased 1.6-fold to 320±45 ng/min/mg tissue, compared with 196±14 ng/min/mg tissue in pair-fed controls (P < 0.05). Hence, the peripheral, but not central, administration of leptin enhances glucose utilization in BAT compared with pair-fed control rats, compatible with a direct peripheral effect of leptin on this tissue.

Figure 1. Effects of leptin on adipocyte metabolism. (A) Leptin enhances the insulin-stimulated glucose utilization index in vivo compared with pair-fed control rats. Rats were injected for 4 d with vehicle, or leptin intravenously (1 mg/d) or intracerebroventricularly (i.c.v.) (12 μg/d), while the control animals were pair-fed to the controls during the entire experiment. At the end of the treatment period, the rats were used for euglycemic-hyperinsulinemic clamps, where the uptake of 2-deoxyglucose into brown adipose was measured (n = 5–6 per group). (B) Concentration-dependent increase in the basal lipolytic rate of ex vivo white fat pads from Sprague-Dawley rats in response to leptin and epinephrine. White fat pads from four animals were pooled and incubated as described in the text. After 1 h of incubation in the presence of either vehicle, leptin or epinephrine, the amount of glycerol released into the medium was measured (32) (n = 2; where no error bars are visible the SEM is < 10%). (C) Time-dependent increase in the lipolytic rate of ex vivo white fat pads from four Sprague-Dawley rats in response to vehicle (basal), leptin (10 nM), or epinephrine (5 nM). The experimental methods are identical to those in B (n = 2; where no error bars are visible the SEM is < 10%). (D) Leptin stimulates lipolysis in ex vivo white fat pads from lean Zucker Fa/fa rats, but not in WAT from obese fa/fa animals, while the white fat pads from both animals respond to epinephrine. The experimental methods are identical to those in B (n = 2; where no error bars are visible the SEM is < 10%).
**Leptin increases lipolysis in WAT fat pads ex vivo.** The effect of leptin on the basal lipolytic rate of epididymal fat pads was assessed ex vivo. As shown in Fig. 1, B and C, leptin increases glycerol release in a dose- and time-dependent manner in fat pads from Sprague-Dawley rats, where leptin is a strong stimulator of lipolysis. On a molar basis, epinephrine stimulated lipolysis about fourfold more potently than leptin. To ascertain that this effect is indeed mediated by leptin and the OB-Rb, a similar experiment was performed in lean (Fa/fa) and obese (fa/fa) Zucker rats. As shown in Fig. 1 D, epinephrine was able to enhance the glycerol release in the fat pads from Fa/fa and fa/fa animals, as expected, whereas only the leptin-sensitive Fa/fa animals responded to leptin in a dose-dependent manner.

**Effects of leptin treatment on gene expression in brown adipocytes.** Primary cultures of in vitro differentiated brown adipocytes from genetically lean Fa/fa or obese fa/fa rats were treated for 24 h with either 1 μM of the thiazolidinedione BRL49653, 625 nM of leptin, or both. Analysis of the extracted total RNA by Northern blotting after treatment with BRL49653 revealed as expected an about twofold increase in the mRNA for malic enzyme, lipoprotein lipase, and the fatty acid–binding protein, aP2, in brown adipocytes derived from lean rats (Fig. 2) (40–42). Interestingly, treatment with leptin resulted in a similar increase in malic enzyme and lipoprotein lipase mRNA (1.8±0.17 and 1.9±0.14-fold, respectively; P < 0.01 compared with control), while the expression of the aP2 gene remained unaltered. The combined treatment with leptin and BRL49653 resulted in a partially additive effect on malic enzyme expression (3.1±0.22-fold, P < 0.05 compared with BRL alone). In contrast, the brown adipocyte cultures from obese Zucker fa/fa rats harboring a partially inactivating mutation in the OB-Rb did not respond to leptin (0.73-fold stimulation), although the thiazolidinedione BRL49653 induced gene expression as expected (2.2-fold) (43–45).

**Adipose tissue expresses OB-Rb.** The presence of small amounts of OB-Rb in adipose tissue has been reported previously (12). Using total RNA extracted from rat hypothalamus, BAT, and WAT, as well as from primary brown and white adipocyte cultures, we performed RT-PCR to demonstrate the presence of OB-Rb transcripts. The amplified products derived from native and cultured adipocytes were sequenced, confirming the presence of the correct PCR product (data not shown). With this assay, OB-Rb mRNA could be detected in tissue samples as well as cultures derived from BAT (Fig. 3). Lean Fa/fa rats as well as obese fa/fa animals expressed OB-Rb as expected, since the fa missense mutation alters intracellular trafficking, rather than expression of the OB-R (43). Similar results were obtained for native and cultured WAT (data not shown).

**Leptin activates STATs in brown and white adipocytes.** To determine whether leptin is capable of activating the Jak/STAT pathway in adipocytes, as previously reported for the hypothalamus, we examined whether leptin would induce nuclear STAT translocation in vivo and in cultured primary adipocytes (15). Nuclear extracts were prepared from BAT 60 and 90 min after the intravenous injection of Fa/fa rats with 500 μg of leptin. The extracts were assayed for the presence of specific STATs by means of an EMSA. Leptin strongly induced the translocation of STAT1, as demonstrated by binding to the M67-SIE DNA element and supershifting with an anti-STAT1, but not anti-STAT3, antibody (Fig. 4 A). In contrast,
and in parallel with the data on glucose utilization (Fig. 1A), no peripheral activation of STATs was observed after the administration of 12 μg of leptin i.c.v. (data not shown).

To determine whether these effects of leptin were direct, primary brown and white adipocyte cultures were treated with leptin and the nuclear extracts assayed for STAT activity. Leptin-inducible binding of STATs in the nuclear extracts from brown and white adipocytes increased 1.8- and 2.7-fold, respectively, after 15 min on the M67-SIF element, which binds STAT1 and 3 (Fig. 4B). The shift in mobility of the band observed after treatment of white adipocytes with leptin suggests the possible transition from hetero- to homodimeric complexes or vice versa. However, no activation of STATs was observed in primary cultures of BAT from obese fa/fa rats, compatible with the genetic leptin resistance of these animals (Fig. 4B, right-most lane). Interestingly, even basal STAT activity was absent in the fa/fa adipocytes, suggesting either that the basal signal observed in Fa/fa cells is due to the paracrine effects of leptin secreted in culture, or, alternatively, due to a general desensitization of STAT signaling in fa/fa adipocytes.

Time course experiments performed in WAT demonstrated a rapid increase in STAT translocation at 15 min (1.7-fold), which increased further after 30 min (3.1-fold; Fig. 4C).

Discussion

In this paper, we demonstrate that leptin has direct effects on BAT and WAT at a metabolic and molecular level. This conclusion is supported by several independent lines of evidence: (a) peripheral, but not central, administration of leptin increases the insulin-stimulated utilization of glucose in BAT when compared with pair-fed control rats; (b) leptin at concentrations as low as 0.1 nM stimulates basal lipolysis in white fat pads ex vivo in a time- and dose-dependent manner, and this effect is absent in fat pads from obese fa/fa rats which are known to be deficient in functional OB-R; (c) leptin induces the expression of malic enzyme and lipoprotein lipase in primary cultures of brown adipocytes; (d) BAT, WAT, and primary adipocyte cultures all express OB-Rb mRNA; and (e) treatment of brown and white adipocytes with leptin in vivo or in vitro induces the nuclear translocation of STATs, compatible with the currently known mechanism of action of leptin through the OB-R. Taken together, our results suggest that
leptin has direct auto- or paracrine effects on adipocytes, which possibly contribute to the weight- and fat-reducing activity of leptin.

While the hypothalamus was the first target organ described for leptin, recent evidence at the level of mRNA and cellular function suggests that peripheral organs also express OB-Rb and are responsive to leptin (12, 17, 22). Specifically, several recent reports demonstrated an inhibitory effect of leptin on the release of insulin from pancreatic β cells, and another paper reported decreased synergism of insulin-like growth factor I on the follicle-stimulating hormone–induced estradiol synthesis when leptin was present (17, 19, 20). In addition, leptin was shown to increase the pituitary secretion of luteinizing hormone, independent of any confounding hypothalamic effects (22). We now show that leptin increases glucose utilization in BAT in vivo and the rate of lipolysis in white fat pads ex vivo. This latter result is compatible with the recent observations of Shimabukuro et al. that leptin treatment in vivo dramatically reduces tissue triglyceride stores compared with pair-fed controls, already suggesting a role for leptin beyond its appetite-reducing properties (25, 26). In addition, the same group has demonstrated increased lipolysis and the increased expression of lipolytic enzymes in primary cultures of pancreatic islets, analogous to our findings in white fat pads (28). However, other groups have failed to detect a significant effect of leptin on basal lipolysis in cultured adipocytes in vitro (27). Our positive results in this study are likely due to the use of intact fat pads ex vivo, since the model of isolated adipocytes used in the negative reports was shown previously to be less responsive to other lipolytic stimuli, such as epinephrine (Zapf, J., unpublished data). The observation that already 0.1 nM of leptin is able to stimulate basal lipolysis and that fat pads from Zucker fa/ fa rats, which are known to harbor a partially inactivating mutation in the OB-Rb gene, are unresponsive to leptin suggests that this effect is specific and of potential physiological relevance. However, the possibility cannot be excluded that the adipocytes from fa/ fa animals have acquired postreceptor defects related to their obese and insulin-resistant state, contributing to their unresponsiveness to the lipolytic action of leptin. Nevertheless, the observation that leptin fails to activate STATs in brown adipocytes derived from the leptin-resistant fa/ fa animals is compatible with the concept that the observed direct effects of leptin might be mediated through the OB-R.

Leptin also enhances the mRNA expression of malic enzyme and lipoprotein lipase, but not of the fatty acid–binding protein aP2, in brown adipocytes from lean Fa/ fa rats, but not in brown adipocytes derived from obese Zucker fa/ fa rats. The only gene shown previously to be regulated directly by leptin in a preadipocyte cell line is the acetyl-CoA carboxylase gene, which is suppressed in the presence of leptin (24). Since the malic enzyme and lipoprotein lipase genes are both regulated by the peroxisome proliferator-activated receptors (PPARs), it is conceivable that leptin activates this nuclear receptor pathway. However, the fact that leptin does not significantly increase aP2 mRNA, which represents another PPAR-regulated gene, suggests that any putative functional interaction between the OB-R and PPAR signaling pathways is more complex. Since leptin not only decreases food intake, but also increases energy expenditure, we examined its effect on the expression of UCP-1 in brown adipocyte cultures. However, we were unable to detect any chances in the levels of UCP-1 mRNA after leptin treatment, and hence it remains to be seen whether the direct effects of leptin on BAT are implicated in regulating thermogenesis (data not shown).

The levels of OB-Rb in all target organs other than the hypothalamus and the hematopoietic system are very low, requiring RT-PCR or RNase protection assays for the detection of transcripts. In this study, we were able to detect OB-Rb mRNA in BAT, WAT, and primary adipocyte cultures, which is compatible with previous reports demonstrating the presence of OB-Rb mRNA in adipose tissue by RNase protection (12, 18). However, other than the hypothalamus, no target tissue for leptin was hitherto shown to respond by nuclear STAT translocation (15). The present experiments demonstrate that the peripheral, but not central, administration of leptin in vivo activates STAT1 in BAT. Experiments with primary cultures of brown and white adipocytes suggest that this effect of leptin on adipocytes is indeed direct and does not involve the hypothalamus. However, the exact nature of the STATs required for transducing the leptin signal is still controversial and may depend on the cellular context and the concomitant presence of other stimuli in vivo. For example, a previous in vivo study implicated only STAT3 in hypothalamic leptin signaling, whereas experiments involving transfected cells have provided evidence for the activation of either STAT3, 5, 6, STAT1, 3, 5, or only STAT1 and 3, depending on the cellular model (12–15).

In summary, we have shown that physiological levels of leptin have direct and specific effect on the metabolism and gene expression of brown and white adipocytes. These observations suggest that leptin may exert its weight-reducing action not only through an endocrine, hypothalamic mode of action, but also through an auto- or paracrine pathway.

Acknowledgments

We are grateful to Dr. D. Lane for providing the aP2 probe, to Dr. M. Schotz for the lipoprotein lipase probe, and to D. V. Nikodem for the malic enzyme cDNA. Leptin was kindly provided by Novartis. The expert technical assistance of Mrs. Mireille de Meyer and Mrs. Marjana Paponja is gratefully acknowledged.

C.A. Meier is supported by a Fellowship from the Howard Hughes Medical Institute. The work was supported by the Swiss National Science Foundation (grants 32-37536.93 to A.G. Burger and C.A. Meier, and 32-046808.96 to J. Zapf) and also in part through a grant-in-aid from Boehringer-Ingelheim International Inc. (to C.A. Meier).

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