Insulin-Dependent Leptin Expression in Breast Cancer Cells

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Abstract

Pathologic conditions associated with hyperinsulinemia, such as obesity, metabolic syndrome, and diabetes, seem to increase the risk of breast cancer. Here, we studied molecular mechanisms by which insulin activates the expression of leptin, an obesity hormone that has been shown to promote breast cancer progression in an autocrine or paracrine way. Using MDA-MB-231 breast cancer cells, we found that (a) insulin stimulated leptin mRNA and protein expression, which was associated with increased activation of the leptin gene promoter; (b) insulin increased nuclear accumulation of transcription factors hypoxia inducible factor (HIF)-1α and Sp1 and their loading on the leptin promoter; (c) small interfering RNA (siRNA)-mediated knockdown of either HIF-1α or Sp1 significantly down-regulated insulin-induced leptin mRNA and protein expression; further inhibition of leptin expression was observed under the combined HIF-1α and Sp1 siRNA treatment; (d) inhibition of extracellular signal-regulated kinase (ERK)1/2 and phosphatidylinositol-3-OH kinase (PI-3K) pathways significantly, albeit partially, decreased insulin-dependent leptin mRNA and protein expression, which coincided with reduced association of HIF-1α and/or Sp1 with specific leptin promoter regions; and (e) inhibition of ERK1/2 reduced recruitment of both HIF-1α and Sp1 to the leptin promoter, whereas down-regulation of PI-3K influenced only HIF-1α binding. In summary, our data suggest that hyperinsulinemia could induce breast cancer progression through leptin-dependent mechanisms. In MDA-MB-231 cells, this process requires Sp1- and HIF-1α–mediated leptin gene transcription and is partially regulated by the PI-3K and ERK1/2 pathways. [Cancer Res 2008;68(12):4919–27]

Introduction

Recent epidemiologic reports suggested that general obesity is associated with an increased risk of developing breast cancer in postmenopausal women, whereas central obesity can increase breast cancer risk in both premenopausal and postmenopausal populations (1, 2). The molecular mechanisms underlying the obesity-breast cancer link are not clear, but it is thought that one important factor might be excess exposure of mammary epithelium to various bioactive substances (for instance, estrogens and growth factors) synthesized by the adipose tissue (2–5). The principal hormone produced by adipocytes is leptin (6). Leptin, the product of the (ob) gene, is best known as a regulator of food intake and energy expenditure via hypothalamic-mediated effects (6). However, leptin also controls other processes such as reproduction, lactation, hematopoiesis, immune responses, cell differentiation, proliferation, survival, and angiogenesis (6). Furthermore, there is evidence that leptin can be involved in neoplastic processes (7). Data obtained using breast cancer cellular and animal models suggested that leptin can stimulate cell growth, survival, and transformation, and interfere with the action of antioxidants (2, 3, 5, 7–9).

Recently, we and others reported that leptin and its receptor (ObR) are overexpressed in breast cancer relative to noncancer mammary epithelium (10–12), which indicates that leptin can influence breast cancer cells not only by endocrine and/or paracrine actions but also through autocrine pathways. We also showed that overexpression of leptin and ObR mRNA in breast cancer could be induced by high doses of insulin (10). Interestingly, although insulin was a major stimulant of leptin in less differentiated estrogen receptor-α (ER)-negative MDA-MB-231 cells, it only modestly up-regulated leptin mRNA in less aggressive ER-positive MCF-7 cells (10). Because hyperinsulinemia, one of the characteristics of obesity, diabetes, and metabolic syndrome, has been associated with increased breast cancer risk (4, 13, 14), we set out to investigate the mechanism by which insulin up-regulates leptin expression in ER-negative breast cancer cells.

The human leptin gene promoter (from –2931 to +1) contains multiple transcription regulatory elements that can be activated by insulin. These include seven Sp1 binding sites (GC boxes) and eight binding sites for hypoxia inducible factor (HIF; ref. 15). Sp1 is a ubiquitous nuclear factor that mediates the effects of insulin in different cell types, regulating such processes as glucose metabolism, lipid biosynthesis (16–18). The abundance of nuclear Sp1 has been shown to be increased by insulin and can be posttranslationally regulated by phosphatidylinositol-3-OH kinase (PI-3K) and extracellular signal-regulated kinase (ERK)1/2 pathways in some cell models (19–21).

HIF is a transcription factor of major importance in cellular response to oxygen deficiency, as it acts as a master regulator of genes involved in tissue reoxygenation (22). In addition, HIF is known to facilitate cancer progression by promoting tumor neoangiogenesis, cell motility, and invasion (23). HIF is a heterodimer consisting of a constitutively expressed HIF-1β subunit and an oxygen-regulated, unstable HIF-1α subunit. HIF interactions with DNA are mediated through Hypoxia-Responsive Elements (HRE; 5′-RCGTG-3′; ref. 22). HIF activation involves HIF-1α stabilization, which occurs either due to hypoxia or under normoxia upon activation of different signaling pathways, for instance, the insulin-responsive PI-3K/mammalian target of...
rapamycin and ERK1/2 cascades (24, 25). Importantly, ERK1/2 may also phosphorylate and activate the histone acetyltransferase p300 (26–28), a common coactivator of Sp1 and HIF (28–30), leading to increased transcriptional activity of both factors. HIF and Sp1 have been implicated in the regulation of the leptin promoter in noncancer tissues (17, 31), according to our recent work, HIF can induce leptin gene in breast cancer cells (32).

Here, we show that in MDA-MB-231 breast cancer cells, insulin-induced up-regulation of leptin expression is regulated by PI-3K and ERK1/2 and depends on Sp1 and HIF-1α interactions with specific regions in the leptin promoter.

Materials and Methods

Cell culture and treatments. MDA-MB-231 cells were grown in DMEM: F12 (Cellgro) containing 5% calf serum. The cells were synchronized in phenol red–free serum-free medium for 24 h and then stimulated with F12 (Cellgro) containing 5% calf serum. The cultures were synchronized in phenol red–free serum-free medium for 24 h and then stimulated with 340 nmol/L insulin (Sigma) for 4 or 16 h, depending on the experiment.

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Western blotting. The cells were stimulated with 340 nmol/L insulin for 4 h and then lysed to obtain cytoplasmic and nuclear proteins, as described before (34). The expression of proteins was analyzed in 70 µg of cytoplasmic or nuclear cell lysates. The following antibodies (Ab) were used for Western blotting (WB): HIF-1α monoclonal (m)Ab (R&D systems); Sp1 (PEP2) polyclonal (p)Ab (Santa Cruz Biotechnology); anti–α-tubulin H235 (Santa Cruz); and nucleolin mAb (Santa Cruz). The proteins were separated on a 4% to 15% polyacrylamide gel, immunoblotted, and quantified as described before (34).

Luciferase reporter assays. Luciferase assays were performed as described before (34). In brief, MDA-MB-231 cells grown in 6-well plates were transfected for 6 h with 0.5 µg DNA mixture per well using Fugene 6 (Roche). The transfection mixtures contained 0.3 µg of either the leptin promoter reporter plasmid pGL3-Ob1 encoding the firefly luciferase (Luc) cDNA under leptin promoter (from -2924 to +31; ref. 35) or the empty vector pGL3. To control transfection efficiency, the cells were cotransfected with 100 ng of pRl-TK-Luc, a plasmid encoding Renilla luciferase (Rl Luc; Promega). Upon transfection, the cells were treated with 340 nmol/L insulin for 4 h, or left untreated. Luciferase activity (Luc and Rl Luc) in cell lysates was measured using Dual Luciferase Assay System (Promega) following the manufacturer’s instructions and quantified as described previously (34). All experiments were repeated at least four times.

Chromatin immunoprecipitation. We followed chromatin immunoprecipitation (ChIP) methodology described in detail previously (34). MDA-MB-231 cells were treated with 340 nmol/L insulin and/or 50 µmol/L LY, 5 µmol/L UO for 4 h, or left untreated. For knockdown experiments, the cells were transfected with HIF-1α siRNA and/or Sp1 siRNA (Applied Biosystems), as described below. Total cellular RNA was isolated using TRIzol reagent (Invitrogen). Three micrograms of total RNA were reverse transcribed using the Taq Man RT kit (Applied Biosystems) according to vendor’s instructions. Two microliters of the RT products were used to amplify leptin sequences using the HS01748771_m1 Lept Taq Man kit (Applied Biosystems). To normalize quantitative real-time PCR (QRT-PCR) reactions, parallel Taq Man assays were run on each sample for α-actin. Changes in the leptin mRNA content relative to α-actin mRNA were determined using a threshold cycle (CT) method (ABI User Bulletin no. 2) to calculate changes in CT and, ultimately, fold and percent change. An average CT value for each RNA was obtained for replicate reactions.

Knockdown of HIF-1α and Sp1 using small interfering RNA. HIF-1α and Sp1 knockdowns were achieved with specific small interfering RNA (siRNAs) purchased from Ambion. HIF siRNA and/or Sp1 siRNA (150 nmol/L) was mixed with the transfection agent RNAiFect (Qiangen; siRNA to RNAiFect ratio, 1:3) and incubated for 15 min at room temperature. Then, the mixture was transfected into MDA-MB-231 cells for 6 h. After that, the cells were placed in fresh medium for 24 h. Next, the cells were treated with 340 nmol/L insulin for 4 h and leptin RNA and protein levels were assessed respectively by QRT-PCR and immunofluorescence/deconvoluted microscopy. In parallel, to examine the efficacy of siRNA treatments, HIF-1α and Sp1 expression levels were determined by immunoblotting. In control experiments, targeting siRNAs were replaced with lamin A/C siRNA (Qiangen). The experiments were repeated thrice for HIF-1α, four times for Sp1, and thrice for both siRNAs.

Leptin detection by immunofluorescence/deconvoluted microscopy. MDA-MB-231 cells were plated in 2-well Permanox chamber slides (Nunc). Subconfluent cultures were transfected with HIF-1α siRNA and/or Sp1 siRNA and treated with 340 nmol/L insulin for 16 h for leptin detection. Then, the cells were fixed for 10 min at −20 °C in methanol. Next, the cells were permeabilized with 0.2% Triton X-100 for 5 min, and unspecific binding was blocked with 7.5% bovine serum albumin fraction V for 1 h at room temperature. Leptin expression was detected using 200 µg/mL Ob pAb A-20 (Santa Cruz) and donkey anti-rabbit IgG-FITC (Santa Cruz). The
slides were covered with Vectashield mounting medium containing 4',6-
diamidino-2-phenylindole (DAPI; Vector laboratories, Inc.) to allow visualization of cell nuclei. The abundance of leptin under different conditions was assessed using Olympus IX81 deconvoluted microscope and Slidebook software. Leptin expression was quantified by determining the percentage of positive cells in at least 10 viewing fields.

**Leptin detection by ELISA.** Subconfluent MDA-MB-231 cells were left untreated or were treated for 4 h with 340 nmol/L insulin, and/or 50 μmol/L LY, and/or 50 μmol/L UO. After that, the cells were counted and conditioned medium was collected and concentrated with Amicon centrifugal filter 10K (Millipore) to final volume of 500 μL. Leptin concentration in conditioned medium was measured using the Human ELISA kit (R&D), which specifically detects leptin without significant cross-reactivity or interference with factors related to or associated with leptin. The lowest detectable concentration was 7.8 pg/mL, intra-assay variation was 4.8%, and inter-assay variation was < 7.2%. The standard curve was done using 0, 7.8, 15.6, 31.2, 62.5, 125, 250, 500, and 1,000 pg/mL leptin concentrations. Leptin concentrations detected in conditioned medium were within the range of the standard curve. The concentrations are expressed as pg/mL per 1.5 × 10^7 cells. All points were done in triplicate and the experiments were repeated thrice.

**Statistical analysis.** The correlations were studied by Student’s t test. P values of < 0.05 were considered statistically significant.

**Results**

Insulin treatment activates the leptin promoter and increases leptin mRNA and protein expression in MDA-MB-231 cells. We have previously shown that leptin is overexpressed in human breast cancer biopsies and that this overexpression might be caused by obesity-related stimuli (10). Here, we focused on the molecular mechanism of leptin activation by high doses of insulin in MDA-MB-231 cells.

To assess whether insulin can affect leptin transcriptional activity, we used the reporter plasmid, pGL3-OB1, containing the luciferase gene under the control of leptin regulatory sequences

\[ \text{Leptin concentration} = \frac{\text{leptin protein}}{\text{leptin mRNA}} \]

We found that insulin significantly (by ~5-fold) increased leptin mRNA concentration in cells transfected with pGL3-OB1, relative to cells transfected with an empty vector pGL3 (Fig. 1A). Using QRT-PCR, we showed that insulin stimulates leptin mRNA synthesis by ~1.8-fold, relative to untreated cells. Using immunofluorescence deconvoluted microscopy, we observed that cellular leptin protein was highly expressed in 65% ± 2% of insulin-treated cells but was detectable only in 10% ± 1% of control cells (Fig. 1B and C). Similarly, insulin significantly (~4.0-fold) increased the levels of secreted leptin, as measured by ELISA (Fig. 1D).

**Insulin stimulates nuclear expression of HIF-1α and Sp1.** Insulin can activate gene transcription through different transcription factors, including both Sp1 and HIF-dependent mechanisms (32, 37–40). Here, we assessed the abundance of nuclear Sp1 and HIF-1α in MDA-MB-231 cells in response to insulin treatment (Fig. 2A). In untreated cells, similar levels of Sp1 were found in the cytoplasm and nucleus, whereas HIF-1α was present only in nuclear protein fractions. Insulin stimulation significantly increased nuclear abundance of both factors (Fig. 2A). The cytoplasmatic protein β-tubulin and nuclear protein nucleolin were used as control of protein loading.

**Insulin stimulates loading of transcription factors on the leptin promoter in MDA-MB-231 cells.** To test if insulin stimulates the loading of Sp1 and HIF-1α as well as other transcription regulators on the leptin promoter, we first used ChIP assays (Fig. 2B). Because the leptin promoter contains multiple potential HIF and Sp1 binding sites, we hypothesized that insulin-induced leptin transcription (Fig. 1A) might be related to the increased association of HIF-1α and Sp1 with leptin regulatory sequences. Indeed, insulin significantly increased HIF-1α and Sp1 binding to two leptin promoter regions: Lep1 region (~992 to +377) containing four HRE

\[ \text{Lep1 region} = \frac{\text{leptin expression}}{\text{leptin promoter}} \]

Thus, we concluded that insulin stimulates leptin expression in breast cancer cells through a mechanism involving the activation of the leptin promoter by HIF-1α and Sp1.
Similarly, insulin stimulated the association of Sp1 by Lep1 and by insulin-dependent leptin mRNA and protein expression. However, inhibition of ERK-1/2 and PI-3K pathways (41, 42) using UO and LY 274352 (UO) inhibited insulin-induced leptin mRNA expression by ~38%. Similar effects were observed with LY (Fig. 3A). LY, but not UO, also blocked basal leptin mRNA by ~50% (Fig. 3A). Similarly, insulin-dependent leptin levels were reduced by UO and LY by ~38% and 41%, respectively, whereas basal levels were not significantly affected (Fig. 3B).

Inhibition of PI-3K reduces insulin-dependent binding of HIF-1α, but not of Sp1, to the proximal leptin promoter. PI-3K and ERK-1/2 pathways are known to regulate HIF-1α expression and activity, whereas Sp1 is influenced mostly by ERK-1/2 (43–46). Here, we studied how inhibition of these pathways can affect insulin-induced recruitment of HIF-1α and Sp1 to the leptin promoter. Using Chip technology, we found that LY significantly (by ~25%) reduced insulin-dependent binding of HIF-1α to Lep1; however, it did not modulate binding of Sp1 to Lep1 or Lep2 regions. Samples precipitated with control IgGs did not reveal any PCR products for Lep1 or Lep2 (data not shown).

Inhibition of ERK1/2 and PI-3K pathways down-regulates insulin-dependent leptin mRNA and protein expression. Leptin expression and activity are regulated by insulin-sensitive ERK1/2 and PI-3K/Akt signaling pathways (41, 42). Using UO and LY inhibitors, we studied whether these pathways are required for leptin expression in MDA-MB-231 cells. The treatment of cells with UO inhibited insulin-induced leptin mRNA expression by ~38%. Similar effects were observed with LY (Fig. 3A). LY, but not UO, also blocked basal leptin mRNA by ~50% (Fig. 3A). Similarly, insulin-dependent leptin levels were reduced by UO and LY by ~38% and 41%, respectively, whereas basal levels were not significantly affected (Fig. 3B).

Figure 2. Insulin increases nuclear levels of HIF-1α and Sp1 and stimulates their loading on the leptin promoter. A, MDA-MB-231 cells were treated for 4 h with 340 nmol/L insulin or left untreated. The expression of HIF-1α (~120 kDa) and Sp1 (~106 kDa) was assessed by WB in 70 μg of cytoplasmic or nuclear proteins using specific Abs, as described in Materials and Methods. Protein loading and purity of fractions were controlled by reprobing WB filters for the expression of a cytoplasmic protein β-tubulin (Tub) and a nuclear marker nucleolin (Nuc). Graphs represent relative HIF-1α and Sp1 expression levels normalized to nuclear marker nucleolin or cytoplasmic protein β-tubulin; columns, mean; bars, SD; *, P < 0.05, control versus insulin. B, soluble chromatin was isolated from MDA-MB-231 cells stimulated with insulin for 4 h, and from untreated cells. The binding of HIF-1α, Sp1, p300, and acetylated histones H3 (H3) to two different leptin promoter regions, Lep1 and Lep2, was assessed by Chip, as described in Materials and Methods. Nonimmune IgG was used as control for precipitating Abs. DNA input in PCR reactions was assessed using Lep1 and Lep2 primers on samples before immunoprecipitations. The graph represents the abundance of HIF-1α, Sp1, p300, and p300 on Lep1 and Lep2 under different conditions ± SD; *, P < 0.05, control versus insulin.

Consensus sequences (~765/760, −632/−627, −571/−566, and −120/−115) and five Sp1 binding motifs (~655/−649, −353/−343, −128/−122, −103/−90, and −22/−16) and Lep2 (regions from −2742 to −1403) containing two Sp1 motifs (~2539/−2534 and −1913/−1918) and four HRE (~2593/−2588, −2467/−2462/−1830/−1825, −1506/−1501; Fig. 2B). Specifically, under insulin treatment, the recruitment of HIF-1α increased by ~6-fold on Lep1 and by ~3.5-fold on Lep2, relative to untreated cells. Similarly, insulin stimulated the association of Sp1 by ~3- and ~2.5-fold on Lep1 and Lep2, respectively. In parallel, we noted an increased association of p300, a histone acetyl transferase and a common coregulator of HIF and Sp1 (28–30) with Lep1 and Lep2, paralleled by higher abundance of acetylated H3 at these regions (Fig. 2B). Samples precipitated with control IgGs did not reveal any PCR products for Lep1 or Lep2 (data not shown).

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unlabeled probes, used as controls, did not precipitate HIF-1α or Sp1 proteins (data not shown).

**Inhibition of ERK-1/2 significantly reduces insulin-dependent binding of HIF-1α to the proximal and distal leptin promoter and of Sp1 to the proximal leptin promoter.** In the presence of UO, insulin-stimulated HIF-1α loading on Lep1 was reduced by ~50% and on Lep2, by ~46%, according to our ChIP analysis. UO also nearly completely blocked the basal level of HIF-1α binding to these regions (Fig. 4A). Furthermore, UO inhibited insulin-dependent association of Sp1 with Lep1 by ~62%. However, no effects of UO were noted for Sp1 binding to Lep2 (Fig. 4A). Samples precipitated with control IgGs did not reveal any Lep 1 or Lep2 PCR products. Similar effects of inhibitors were obtained in DAPA assays, where UO down-regulated HIF-1α binding to ~602/–631 and ~2592/–2562 sequences in Lep 1 and Lep 2, respectively, and reduced the association of Sp1 to ~354/–323 Lep1 region (Fig. 4B).

**Insulin-induced expression of leptin in breast cancer cells requires HIF-1α and Sp1.** To determine whether HIF-1α and Sp1 are required for insulin-dependent leptin expression in breast cancer cells, we inhibited basal and insulin-induced HIF-1α and Sp1 expression with RNA interference. Targeting siRNAs down-regulated basal HIF-1α and Sp1 protein levels by ~98% and 85%, respectively, whereas unrelated lamin A/C siRNAs had no effects on these proteins (Fig. 5A; data not shown). HIF-1α and Sp1 knockdowns were paralleled by ~47% and 50% inhibition of insulin-induced leptin mRNA levels, respectively. Using a mixture of both siRNAs, we observed further (~70%) reduction of insulin-dependent leptin mRNA expression (Fig. 5B). No significant effects of single siRNAs were seen on basal leptin mRNA expression, but the combination of both siRNAs reduced basal leptin levels by ~50% (Fig. 5B). Similarly, siRNAs targeting either HIF-1α, Sp1, or both transcription factors reduced cellular leptin expression by 58% ± 4%, 46% ± 2%, and 100% ± 0%, respectively (Fig. 5C).

The hypothetical model of insulin-dependent leptin expression through Sp1- and HIF-1α–dependent transcription in MDA-MB-231 cells is presented in Fig. 6.

**Discussion**

Pathologic conditions associated with hyperinsulinemia, such as obesity, metabolic syndrome, and diabetes, have been shown to increase the risk of breast cancer (1, 2, 5). Insulin is known to stimulate breast cancer cell growth and transformation; however, molecular mechanisms of these effects are only partially known (47). Here, we studied how insulin activates the expression of obesity hormone leptin in breast cancer cells.

Studies in cellular and animal models showed that leptin can increase breast cancer cell growth, survival, tumorigenicity, and drug resistance (2, 7). Current evidence suggests that leptin is significantly overexpressed in breast cancer relative to noncancer mammary epithelium (10–12). As suggested by our previous work, leptin mRNA overexpression can be induced by obesity-related stimuli, such as insulin, insulin-like growth factor 1 (IGF-I), estrogens, or hypoxia (10). Interestingly, compared with other mitogens or hypoxia-mimetic agents, insulin was the most effective leptin inducer in ER-negative, metastatic MDA-MB-231 cells (10). Our studies on the molecular mechanism of this phenomenon revealed that: (a) in MDA-MB-231 cells, insulin stimulates leptin mRNA and protein expression, which is associated with transcriptional activation of the leptin gene promoter; (b) insulin increases nuclear accumulation of transcription factors HIF-1α and Sp1 and their loading on the leptin promoter, especially association of HIF-1α with ~602/–631 and ~2592/–2562 sites, and Sp1 with ~354/–323 and ~2540/–2510 sites; (c) siRNA-mediated knockdown of either HIF-1α or Sp1 significantly down-regulates insulin-induced leptin mRNA and protein expression; this effect is further amplified by double knockdown of both factors; and (d) inhibition of ERK1/2 and PI-3K pathways partially decreases insulin-dependent leptin expression, which is associated with reduced association of HIF-1α and/or Sp1 with specific leptin promoter regions.

In MDA-MB-231 cells, 340 nmol/L insulin increased leptin expression through transcriptional mechanism (Figs. 1 and 2). Because the human leptin promoter contains multiple Sp1 binding sites and HRE motifs, we tested whether these transcription factors are involved in insulin-induced leptin expression. ChIP assays showed that insulin increased nuclear abundance of both Sp1 and HIF-1α, and elevated binding of these transcription factors to Lep1 and Lep2 promoter regions. These effects were paralleled by increased binding of p300 and acetylation of histone H3 (Fig. 2). Further DAPA analysis proved that within the studied promoter regions, ~602/–631 and ~2592/–2562 were the preferential HIF-1α binding sites, and

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**Figure 3.** Insulin-induced leptin mRNA and protein expression is mediated by PI-3K and ERK1/2. A. MDA-MB-231 cells were treated for 4 h with 340 nmol/L insulin, and/or 50 μmol/L LY, and/or 5 μmol/L UO, as described in Materials and Methods. The abundance of leptin mRNA was studied with QRT-PCR in MDA-MB-231 cells untreated or treated with insulin, and/or LY, and/or UO as described in Materials and Methods. The graph represents the increase of leptin mRNA relative to the increase of α-actin mRNA in the same sample ± SD. *, *P < 0.05 basal versus inhibitor-treated or insulin versus inhibitor-treated. B. The cells were treated with insulin in the presence or absence of LY or UO. Secreted leptin concentrations were measured as described in Materials and Methods and represent pg leptin per milliliters of conditioned medium from 1.5 × 10⁶ cells. Columns, mean; bars, SD. *, *P < 0.05 basal versus inhibitor-treated or insulin versus insulin + inhibitor–treated.
-354/-323 and -2540/-2510 were the sites for Sp1 binding (Figs. 4 and 6). Interestingly, Sp1 and HIF-1α did not associate with other individual consensus sites in response to insulin (data not shown). This could reflect true transcriptional preferences under insulin treatment, or be simply related to high CG content and tertiary structures of our DNA-DAPA probes and their poor hybridization with transcription factors in vitro. Interestingly, in ChIP assays, loading of HIF-1α and Sp1 tended to be more abundant on Lep1 than on Lep2, possibly due to preferential chromatin structure within the region containing -602/-631 HRE and -354/-323 Sp1 motifs (Fig. 6).

The importance of HIF-1α and Sp1 for leptin mRNA and protein expression was confirmed using specific siRNAs. Indeed, down-regulation of both transcription factors produced nearly complete inhibition of insulin-dependent leptin mRNA expression and resulted in total blockade of insulin-induced leptin protein expression (Fig. 5). This is in agreement with the notion that HIF-1α and Sp1 cooperate in the regulation of insulin-induced

Figure 4. Insulin controls HIF-1α and Sp1 loading on the leptin promoter. A, MDA-MB-231 cells were treated for 4 h with insulin, and/or LY, and/or UO or left untreated. The binding of HIF-1α and Sp1 to Lep1 and Lep2 regions was assessed by ChIP with specific Ab, or control IgG, as described in Materials and Methods. DNA input in PCR reactions was assessed using Lep1 and Lep2 primers on samples before immunoprecipitations. The graph represents relative abundance of HIF-1α and Sp1 on Lep1 and Lep2 under different conditions ± SD; *, P < 0.05 basal versus inhibitor-treated or insulin versus insulin + inhibitor–treated. B, the cells were treated as described above. The binding of nuclear HIF-1α or Sp1 to specific leptin promoter regions was analyzed by DAPA, as described in Materials and Methods. Columns, mean; bars, SD; *, P < 0.05 basal versus treated samples.
leptin transcription in adipocytes (17). However, effective down-regulation of HIF-1α (98%) and Sp1 (85%) did not produce similarly profound effects on leptin mRNA, suggesting that other transcription factors might be involved.

The transcriptional activity of the leptin promoter seemed to be differentially regulated by PI-3K and ERK1/2, two major pathways activated by insulin. The ERK1/2 pathway controlled the association of HIF-1α with Lep1 and Lep2 as well as Sp1 binding to Lep1, which might be related to significant regulation of both factors by ERK-1/2–dependent phosphorylation (27, 43, 44). On the other hand, PI-3K regulated HIF-1α binding only on the proximal Lep1 promoter region (Figs. 4 and 6). Interestingly, loading of Sp1 on Lep2 was not blocked by the PI-3K or ERK1/2 inhibitors (Figs. 4 and 6), suggesting that insulin used other pathways, for instance PKCα to phosphorylate Sp1 and activate Sp1/DNA interaction (48). Insulin could also affect leptin transcription through crosstalk with other receptors or intracellular kinases, such as src. The substantial, albeit partial, involvement of the PI-3K and ERK1/2 pathways in insulin-dependent leptin mRNA and protein expression was confirmed by QRT-PCR and ELISA data (Fig. 3). The basal leptin mRNA and protein expression seemed to be regulated by HIF and Sp1, and PI-3K could be involved on the transcriptional level (Figs. 3 and 5). Although PI-3K might influence basal HIF activity, it is not clear which pathways affected basal Sp1 activity.

Our results demonstrating insulin-dependent leptin transcription in MDA-MB-231 breast cancer cells are in agreement with data obtained in other cells models (17, 36). We are the first to show that in MDA-MB-231 cells, insulin-induced leptin expression is regulated by PI-3K and ERK1/2 pathways and requires Sp1- and HIF-1α–mediated transcription. Previously, we showed that MCF-7 breast cancer cells can produce leptin in response to estradiol, insulin, IGF-1 treatment, as well as in response to physiologic or pharmacologic hypoxia (10). However, in MCF-7 cells, the action of insulin was mediated by HIF-1α, although Sp1 was not involved (data not shown; ref. 32). This implicates that the mechanisms of leptin synthesis in breast tumors might be cell context–dependent;
for instance, the process is somehow dependent on the presence or absence of ER, as this receptor is known to influence insulin and IGF-1 signaling pathways and their outcome (33, 49).

In summary, our data suggest that hyperinsulinemia could induce breast cancer progression through leptin-dependent mechanisms.

Disclosure of Potential Conflicts of Interest

No potential conflicts of interest were disclosed.

References


Figure 6. Model of insulin-induced leptin transcription in MDA-MB-231 cells. Insulin induces HIF-1α and Sp1 binding with the leptin promoter regions Lep1 and Lep2. Potential HIF-1α and Sp1 binding sites are indicated by open ovals or rectangles, respectively, whereas identified by DAPA sites are indicated by black shapes. Association of HIF-1α with Lep1 is controlled in part by PI-3K and ERK1/2. Binding of Sp1 to Lep1 is enhanced by ERK1/2. On Lep2, binding of HIF-1α is partially controlled by ERK1/2, whereas neither PI-3K nor ERK1/2 modulates Sp1 association.

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