Insulin Regulates Apolipoprotein B Turnover and Phosphorylation in Rat Hepatocytes

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Abstract

Our laboratory has previously shown that insulin inhibits the secretion of newly-synthesized and immunoreactive apo B from rat hepatocytes. We have also shown that apo B is secreted as a phosphoprotein and that phosphorylation is increased in hyperinsulinemic nonketotic diabetic rats. The present studies were conducted to determine whether the ability of insulin to inhibit apo B secretion is related to alterations in apo B turnover and whether insulin itself affects apo B phosphorylation. Pulse-chase studies with [35S]methionine in primary cultures of hepatocytes from normal rats in the absence and presence of insulin show that the secretion of apo B100 and apo B48 are inhibited by insulin and that this inhibition may be due in part to enhanced intracellular degradation. In addition, there is a second intracellular apo B48 pool which is not insulin regulated or degraded. In experiments in which hepatocytes were incubated with [32P]orthophosphate, insulin decreased 32P incorporation into apo B100 (42%) with only small effects on apo B48 (11%). The small insulin effect on apo B48 may relate to an insulin-insensitive apo B48 intracellular pool. These studies show that insulin can affect the intracellular turnover, secretion, degradation, and phosphorylation of apo B and emphasize the differential regulation of apo B100 and apo B48 with regard to these parameters in rat liver. (J. Clin. Invest. 1990. 86:1746-1751.) Key words: insulin • apolipoprotein B • hepatocytes

Introduction

Hepatically synthesized apo B is necessary for the normal assembly and secretion of VLDL (reviewed in reference 1). The peripheral metabolism of VLDL and apo B in part determine the level of circulating LDL cholesterol (2). Because both LDL cholesterol (3) and apo B100 (4) levels are associated with atherosclerotic disease, an understanding of the control of hepatic apo B synthesis and secretion is important. Acute studies in man (5, 6) suggest that insulin administration decreases hepatic secretion of VLDL. Using primary cultures of rat hepatocytes (7-11) and HepG2 cells (12, 13), we (7, 8) and others (9-13) have shown that insulin exposure for < 24 hours inhibits the secretion of VLDL (8-12) and apo B (7, 8, 11-13). In contrast, chronic in vivo and in vitro hyperinsulinemia stimulates VLDL secretion (reviewed in reference 14). These apparent contradictions may be best explained by the possibility that hyperinsulinemia induces resistance to the ability of insulin to alter apo B metabolism (14).

The mechanisms by which insulin acutely inhibits hepatic apo B secretion are unknown. Davis and co-workers (15, 16) recently showed that there is substantial intracellular turnover of both apo B48 and apo B100 in rat hepatocytes cultured in the presence of insulin (15) and these cells secrete apo B as a phosphoprotein with phosphorylation of serine residues (16). The specific effects of insulin are, however, unclear because in these studies no incubations in culture media lacking insulin were performed. We have shown that the secretion of both newly-synthesized and immunoreactive apo B is reduced in primary cultures of hepatocytes from hyperinsulinemic nonketotic diabetic rats (17) and is associated with an increase in phosphorylation of apo B on both tyrosine and serine residues (18). Phosphorylation of apo B may be an important modulator of the intracellular assembly and secretion of VLDL as demonstrated for vitellogenin (19). In the chick hepatocyte, phosphorylation of vitellogenin is mandatory for secretion of VLDL particles (19). Thus, insulin may be affecting the secretion of apo B from the liver by altering its intracellular turnover and phosphorylation. In addition, these effects may be time dependent.

In this report we have investigated whether alterations in intracellular apo B turnover and phosphorylation are associated with insulin’s inhibition of apo B secretion.

Methods

Materials. All culture supplies, reagents, and animals were obtained from sources previously described (8). Pansorbin cells were from Calbiochem-Behring Corp. (La Jolla, CA) and carrier-free [131I]-Na, [32P]orthophosphate, and [35S]methionine (sp act > 300 Ci/mmol) were obtained from Amersham Corp. (Chicago, IL).

Preparation of hepatocytes. Freshly-isolated hepatocytes were prepared from ad libitum fed male Sprague-Dawley rats as previously described (20). Primary cultures of hepatocytes (8) were maintained on collagen-coated plates in serum-free DME for 18 h.

Pulse-chase incubations. After overnight culture, the monolayers were switched to a methionine-free modified DME (8). Six plates each
were incubated for an additional 2 h in the absence and presence of insulin (100 nM) after which an identical amount of [35S]methionine (90-190 μCi/ml) was added to each of the 12 plates within an experiment. After 15 min of incubation, the plates were placed on ice and the media was replaced with 4.2 ml of DME containing unlabelled methionine (10 mM) (chase medium) and the incubation continued at 37°C. Insulin was readded to plates previously exposed to insulin. Incubations were terminated at 0, 10, 20, 45, 90, and 180 min after the addition of chase medium (15). The media were removed and immediately mixed with equal volumes of a quench solution consisting of PMSF (1 mM), EDTA (4 mM), sodium azide (0.1%, wt/vol), N-ethylmaleimide (8 mM), aprotinin (1,000 U/ml), soybean trypsin inhibitor (0.01%, wt/vol), and 0.001% (wt/vol) of leupeptin, peptastatin, antipain, and chymostatin. The monolayers were washed with ice-cold PBS, pH 7.4, and the cells recovered by scraping in PBS followed by centrifugation (200 g, 5 min). Lysis solution (0.5 ml) consisting of Tris-HCl (50 mM, pH 8), Triton X-100 (3%, vol/vol), sodium sacrose (1%, wt/vol), SDS (0.3%, wt/vol), aprotinin (1,000 U/ml), soybean trypsin inhibitor (0.01 U/ml), sodium orthovanadate (20 mM), PMSF (2 mM), EDTA (1 mM), N-ethylmaleimide (8 mM), and NaF (10 mM) was added to the cell pellets and the mixtures vortexed and centrifuged (10,000 g, 10 min). The supernatants were removed and mixed with equal volumes of 2 M LiCl. Media and cellular extracts were precleared using normal rabbit serum (21) and the 35S-labeled apo B immunoprecipitated overnight at 4°C. Protein A (Pansorbin cells, 100 μl/ml) was added for 2 h at 4°C. The samples were centrifuged, the supernatants removed, and subjected to two additional overnight immunoprecipitations. Preliminary experiments established that complete recovery of 35S-labeled apo B48 and apo B100 from cell extracts and media was achieved using three sequential immunoprecipitations. The pellets from each immunoprecipitation were combined, washed, and boiled in Tris-HCl buffer, (80 mM, pH 6.8), containing SDS (3.8%, wt/vol), mercaptoethanol (5%, vol/vol), and glycerol (10%, vol/vol). Insoluble material was removed by centrifugation, the supernatants made 2 M in urea, and subjected to SDS-PAGE (22) in a 4% gel containing 1 M urea. After Coomassie blue staining, gels were destained, dried, autoradiographed, and scanned with a Brumma Ultrascan XL laser densitometer (LKB Instruments, Inc., Gaithersburg, MD). Apo B band development was in a linear range.

Apo B phosphorylation. Freshly-isolated hepatocytes were preincubated in serum-free Waymouth’s medium (20). Final cell suspensions (5-10 x 10⁶ cells/ml) in Waymouth’s medium containing 10 mM orthophosphate (2 mM Ci/ml) were incubated for 60 min in the absence of insulin. Insulin (100 nM) was then added to one-half of the cells. After 90-190 min, the isometric isotonometric tension was terminated and rapid centrifugation solutions were immediately solubilized with 2 ml of lysis solution and mixed with LiCl as described above. After overnight dialysis at 4°C against PBS, pH 7.4, containing protease and phosphatase inhibitors, the dialysates were centrifuged and apo B immunoprecipitated as described above except that preclarification with normal rabbit serum was omitted. SDS-PAGE was performed as described above except that 3.5-12% linear gradient gels were employed. To establish that electrophoretically separated apo B bands were completely delipidated, samples of rat serum VLDL were identical solubilized, run on SDS-PAGE, and apo B100 and apo B48 electroeluted from the gel. Both peptides were completely devoid of free fatty acids as analyzed by gas chromatography-mass spectroscopy (data not shown).

Apo B antibody. Rabbit polyclonal antibodies to apo B100 and B48 were made from rat VLDL as described by Davis et al. (16). The antibody was tested against an affinity purified rat apo B antibody (23). Both antibodies immunoprecipitated the same bands from unlabelled VLDL, 125I-labeled VLDL, and 35S-labeled intracellular and secreted material. In addition, the same bands were identified on Western blots of rat VLDL, and immunoprecipitated material from solubilized hepatocytes and hepatocyte media. Hepatocytes contain three intracellular immunoprecipitatable phosphate proteins identified as apo B100, apo B97, and apo B48 using iodinated VLDL as a standard. This was confirmed by Western blots of VLDL and intracellular immunoprecipitated apo B using both antibodies. Apo B100 and apo B97 are designated in our study as apo B100.

Statistical analysis. Statistical analysis were performed using Student's paired t test.

Results

Intracellular apo B turnover and secretion. To evaluate the effects of insulin on intracellular apo B turnover, we performed pulse-chase experiments in primary cultures of rat hepatocytes as described in Methods. Data representing the results from five experiments are shown in Fig. 1 and Table I. The data shown in Fig. 1 represent the mean ± SEM of apo B content at each time point and the data shown in Table I represent the mean ± SEM of the respective peak values from each experiment (see table legend). By either presentation, in hepatocytes untreated with insulin, there is greater intracellular accumulation of newly-synthesized apo B48 than apo B100. In contrast to a nearly twofold higher intracellular level of apo B48 present, the absolute amounts of apo B48 and apo B100 depleted from hepatocytes after 180 min of cold methionine chase are comparable (18.3±4.3 and 15.7±2.1 U/mg protein, respectively) as is the amount of apo B secreted from these cells (13.5±1.2 and 11.0±2.4 U/mg protein representing media recoveries of 74 and 70% for apo B48 and apo B100, respectively). In the absence of insulin, the amount of intracellular apo B48 (4.9±4.7 U/mg protein) and apo B100 (4.7±3.3 U/mg protein) degraded represents ~27-30% of the apo B48 or apo B100 depleted from the cell.

Although the absolute amounts of apo B48 and apo B100 secreted and degraded in the absence of insulin are comparable, the turnover rates of intracellular apo B48 and apo B100 differ substantially. By 90 min after the cold methionine chase, 34% of the newly-synthesized apo B48 has disappeared from the cells (intracellular peak, 28.5±4.0 U/mg protein; amount depleted, 9.7±1.7 U/mg protein) compared to an 83% depletion of apo B100 (intracellular peak, 16.8±2.1 U/mg protein; amount depleted, 13.9±1.9 U/mg protein). By 180 min after the cold methionine chase, the depletions of apo B48 and apo B100 equal 64% (intracellular peak, 28.5±4.0 U/mg protein; amount depleted, 18.3±4.3 U/mg protein) and 93% (intracellular peak, 16.8±2.1 U/mg protein; amount depleted, 15.7±2.1 U/mg protein), respectively. Calculation of the half-lives of the two apo B variants from untreated hepatocytes gives values of 125 and 43 min for apo B48 and apo B100, respectively (Fig. 1, insets).

In accordance with our previous studies (7, 8), the addition of insulin causes a reduction (27% and 35%, respectively, at 180 min) in the amount of newly-synthesized 35S-labeled apo B48 (13.5±1.2 reduced to 9.9±1.4 U/mg protein, P < 0.05) and apo B100 (11.0±2.4 reduced to 7.1±1.9 U/mg protein, P < 0.05) peptides chased into the media compared to the control without insulin (Fig. 1, Table I). This effect is now shown to be posttranslational in nature because insulin has no significant effect on the rates of synthesis of labeled apo B peptides during the [35S]methionine pulse (Table I). Furthermore, the ability of insulin to inhibit both apo B48 and apo B100 secretion is not related to any significant changes in the half-lives of the two apo B variants (Fig. 1, insets, 139 and 31 min, respectively). In the presence of insulin, the amount of apo B48 and apo B100 degraded intracellularly is substantially increased but the time dependency of insulin's effects on these parame-
ters differ between apo B48 and apo B100. In the presence of insulin and after 90 min of cold methionine chase, apo B48 and apo B100 secretion are significantly inhibited (for apo B48, 8.8±1.1 reduced to 4.5±1.1 U/mg protein representing 49% inhibition, P < 0.05; for apo B100, 10.8±1.8 reduced to 5.8±1.1 U/mg protein representing 46% inhibition, P < 0.05) and intracellular degradation significantly enhanced (for apo B48, from 1.0±1.2 to 11.8±3.2 U/mg protein, P < 0.05; for apo B100, from 3.1±2.8 to 12.0±3.6 U/mg protein, P < 0.01). The comparable effect of insulin on the absolute amount of apo B48 and apo B100 secreted or degraded after 90 min of cold methionine, which is equivalent to more than two half-lives for apo B100 and less than one half-life for apo B48, is not preserved when measurements of these parameters for apo B48 are made after an additional 90 min of chase (at 180 min which is equivalent to < 1.5 half-lives for apo B48). As shown in Table I, no insulin-dependent inhibition of apo B48 secretion or enhancement of degradation are found between 90 and 180 min.

Apo B phosphorylation. In initial experiments, we attempted to evaluate the effects of insulin on apo B phosphorylation in primary cultures of hepatocytes, but due to the limitation of cell number per unit volume and low incubating specific activities, the results were inconclusive. As an alternative, we elected to employ freshly-isolated hepatocytes. Previously, we have shown freshly-isolated rat hepatocytes to be insulin-responsive with regard to aminoisobutyric acid uptake (20) and lipid synthesis (24) and have found freshly-isolated human hepatocytes to be insulin responsive with regard to apo B secretion (25). In addition, under the same incubation conditions used to evaluate apo B phosphorylation, we observed a consistent inhibition (~ 25%) of apo B secretion (as measured by RIA) from freshly-isolated rat hepatocytes incubated with insulin and found no detectable degradation of secreted VLDL apo B as measured by isopropanol (125I-VLDL; media from untreated hepatocytes, 48.1±4.6%; media from insulin-treated hepatocytes, 46.1±4.1%) and TCA precipitation (125I-VLDL precipitated; media from untreated hepatocytes, 96.7±1.4%; media from insulin-treated hepatocytes, 94.6±2.4%) of exogenously added rat 125I-VLDL. Extending our previous results with hepatocytes in culture, Fig. 2 shows that freshly-isolated hepatocytes in suspension incorporate 32P orthophosphate into intracellular apo B. Also, 32P-labeling of apo B100 relative to apo B48 is very similar to that found with [35S]methionine in the cultured hepatocytes (Fig. 1 & Table I). The addition of 100 nM insulin to hepatocytes for 90 min decreased 32P incorporation into apo B100 with only a small decrease in 32P incorporation into apo B48. This is illustrated graphically by the scan of this gel (Fig. 2 inset) demonstrating a 42% decrease in intensity of the apo B100 band with an 11% decrease in the apo B48 band.

Discussion

An understanding of the mechanisms by which insulin regulates apo B synthesis, the assembly of VLDL particles, and the secretion of VLDL are critical to our understanding of the regulation of lipoprotein production in normal physiology and in altered metabolic states such as obesity, diabetes, fasting, and glucocorticoid excess. We (7, 8) and others (11) have shown that insulin at postprandial concentrations inhibits the secretion of apo B from primary cultures of rat hepatocytes. The present study extends these observations by examining the effects of insulin on intracellular turnover and secretion of apo B48 and apo B100. The data show that in the absence of insulin most of the intracellular newly-synthesized 35S-labeled apo B48 and apo B100 depleted from the cells during a 90-min chase can be recovered in the media (91 and 78% for apo B48

Figure 1. Pulse-chase labeling of apo B in primary cultures of rat hepatocytes. Newly-synthesized apo B48 (A) and apo B100 (B) from untreated (open circles) and insulin-treated (100 nM, solid circles) hepatocytes were labeled with [35S]methionine and chased with cold methionine for various times up to 180 min as described in Methods. After immunoprecipitation and SDS-PAGE, intracellular (closed circles) and media (open circles) apo B radioactivity was determined. (Insets) First-order kinetic analysis of apo B48 (A) and apo B100 (B) depletions.
Table I. Time-dependent Changes in Intracellular, Secreted, and Degraded Apo B48 and Apo B100

A. Apo B48*

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<td>−Insulin</td>
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<td>Peak to 180 min</td>
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Time interval

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<tr>
<td>Depleted*</td>
<td>18.3±4.3</td>
<td>20.8±5.8</td>
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<tr>
<td>Secreted*</td>
<td>13.5±1.2</td>
<td>9.9±1.4</td>
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<td>Degraded†</td>
<td>4.9±4.7</td>
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B. Apo B100*

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<td>−Insulin</td>
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<td>Peak to 180 min</td>
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<td>16.8±2.1</td>
<td>20.9±3.5</td>
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Time interval

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<td>−Insulin</td>
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<tr>
<td>Depleted‡</td>
<td>15.7±2.1</td>
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<td>Secreted‡</td>
<td>11.0±2.4</td>
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<td>Degraded‡</td>
<td>4.7±3.3</td>
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* All data are expressed in scanning units per milligram protein. † Depleted apo B is the difference between the peak intracellular level and that at the designated time for each experiment. ‡ Intracellularly degraded apo B is calculated by subtracting secreted apo B from depleted apo B for each experiment. Statistical comparisons between untreated and insulin-treated hepatocytes. § P < 0.05. ‡ P < 0.01.

and apo B100, respectively). In the presence of insulin, intracellular degradation of both apo B48 and apo B100 are substantially enhanced because only 28 and 33% of intracellularly depleted apo B48 and apo B100, respectively, are recovered in the media (Table I). Because primary cultures of hepatocytes do not take up and degrade apo B-containing particles secreted into the medium (26), the results indicate that insulin-dependent enhancement of apo B48 and apo B100 intracellular degradation contributes substantially to reduced apo B secretion. However, the relative contributions of the direct inhibitory effects of insulin on apo B secretion remain to be determined.

Borchardt and Davis (15) reported significant intracellular degradation of both apo B48 (40%) and apo B100 (64%) in rat hepatocytes cultured in the presence of insulin (1 µg/ml, 167 nM). Our data after 180 min of cold methionine chase agree with their findings (Table I, apo B48 and apo B100 degradation equal 52 and 64%, respectively, in insulin-treated hepatocytes). In addition, Borchardt and Davis (15) and Swift et al. (27) have shown the secretion of apo B48 to be slower than that of apo B100 in primary cultures of hepatocytes which may relate to a larger intracellular pool of apo B48 (27). The present study confirms and extends these observations. With regard to hormonal sensitivity, we have observed that the ability of insulin to modulate apo B48 secretion and degradation is attenuated between 90 and 180 min after cold methionine chase. These observations suggest the existence of two apo B48 pools and that the acute effects of insulin as assessed with a pulse-chase protocol are primarily on a more rapidly turning over pool.

In this report we have shown that the addition of insulin decreases 32P incorporation primarily into apo B100 with less effect on apo B48. Thus, apo B100 is one of several proteins whose phosphorylation-dephosphorylation state is affected by insulin (reviewed in reference 28). Recently, Capasso et al. (29) have shown that intact rat liver Golgi vesicles translocate ATP into their cisternal space and use it to phosphorylate a set of secretory proteins. The postulated function of these phosphorylations is to render proteins (e.g., casein) resistant to proteolytic degradation (29). While our data show that in the presence of insulin and after 90 min of cold methionine chase, the absolute amount of apo B100 degraded is comparable with apo B48, this insulin effect on apo B48 is not seen from 90 to 180 min after the cold methionine chase. Relative to the peak amounts of newly-synthesized apo B100 and apo B48 present in insulin-treated cells (Table I, 20.9±3.5 and 35.9±5.6 U/mg protein, respectively), the percentages of apo B100 and apo B48 degraded (57 and 33% after 90 min and 60 and 30% after 180 min, respectively) differ and may relate to the greater inhibitory effect of insulin on apo B100 phosphorylation (42 and 11% for apo B100 and apo B48, respectively) when evaluated under conditions of continuous exposure to [32P]orthophosphate. Taken together, the 35S and 32P apo B labeling data suggest the presence of two separate hepatic pools of apo B48,
one which is phosphorylated and constitutively secreted, and one pool which is phosphorylated but whose phosphorylation is regulated by insulin allowing for channeling into a degradative intracellular pathway. The entire apo B100 pool appears to be insulin sensitive.

Another possible role for insulin-regulated changes in the phosphorylation of intracellular apo B could be to regulate VLDL particle size. Support for a role for phosphorylation in the association of proteins and lipids comes from the studies of Powell and Glenny (30) who demonstrated that dephosphorylated calpain 1 (lipocortin 1) had increased affinity for phosphatidyserine liposomes compared to the phosphorylated form. Thus, decreased phosphorylation of intracellular apo B100 in the presence of insulin could result in increased lipid association and VLDL particle size.

We have previously demonstrated (18) that both apo B100 and apo B48 are secreted as phosphoproteins and that phosphorylation of both peptides increased in hepatocytes from hypoinsulinemic nonketotic diabetic rats. This enhanced apo B phosphorylation could decrease the affinity of apo B for lipids thus leading to smaller, more dense VLDL particles characteristic of those secreted from livers of diabetic rats (31). However, whereas short-term insulin exposure and diabetes both lead to decreased apo B secretion, they have opposite effects on apo B phosphorylation.

In summary, these studies indicate that insulin acutely decreases the secretion of apo B48 and apo B100 and insulin’s effect may, in part, be related to increased intracellular degradation. The metabolism of apo B48 and apo B100 differ with regard to intracellular turnover which may relate to the presence of an insulin-insensitive intracellular apo B48 pool. Differences in apo B48 and apo B100 metabolism are further demonstrated in that the ability of insulin to inhibit apo B phosphorylation is more clearly evident in apo B100. These multiple and divergent effects of insulin on hepatic apo B variants emphasize the complexity of mechanisms by which insulin regulates the assembly, intracellular turnover, and the secretion of VLDL particles. Because human liver secretes only apo B100, our demonstration of insulin effects on apo B100 phosphorylation and turnover may be relevant to human apo B metabolism.

Acknowledgments

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References


