Regulation of Lipoprotein Lipase in Primary Cultures of Isolated Human Adipocytes

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Abstract

To study the regulation of adipose tissue lipoprotein lipase (LPL) in human adipocytes, omental adipose tissue was obtained from healthy subjects and digested in collagenase. The isolated adipocytes thus obtained were suspended in Medium 199 and cultured at 37°C. Cell viability was demonstrated in adipocytes cultured for up to 72 h by constancy of cell number, cell size, trypan-blue exclusion, and specific 125I-insulin binding. In addition, chloroquine induced an increase in cell-associated 125I-insulin at 24, 48, and 72 h after preparation. Thus, isolated adipocytes retained their ability to bind, internalize, and degrade insulin.

LPL was measured as activity secreted into the culture medium (CM), released from cells by heparin (HR), and extracted from cell digests. A broad range of heparin concentrations produced a prompt release of LPL from a rapidly replenishable pool of cellular activity. When cells were cultured in medium containing 10% fetal bovine serum, there was a marked stimulation of CM and HR. The secretory response to serum (CM) correlated strongly with HR 24 h after preparation ($r = 0.731, P < 0.001$). In addition, HR was found to correlate logarithmically and inversely with body mass index ($r = -0.731, P < 0.001$). Insulin, at 400 ng/ml only, increased HR by $36 \pm 10\%$, an effect simulated by lower concentrations of insulin-like growth factor-1 (IGF$_1$).

Thus, LPL is produced and regulated in isolated human adipocytes. The degree of adiposity and serum are important regulators of HR activity, whereas insulin is stimulatory only at a pharmacologic concentration. This effect of insulin may be mediated through the IGF$_1$ receptor. Isolated human adipocytes represent a novel and useful system for the study of LPL and lipid metabolism as well as for other aspects of adipocyte biology.

Introduction

Adipose tissue lipoprotein lipase (LPL) is the enzyme responsible for the hydrolysis of the triglyceride core of chylomicrons and very low density lipoproteins into monoaoylglycerol and free fatty acids (1). LPL deficiency may be involved in the hyperlipidemia seen in numerous disease states, such as poorly controlled diabetes mellitus (2, 3), chronic renal failure (4), and hypothyroidism (5, 6). Increases in adipose tissue LPL after weight loss suggest a role for this enzyme in weight maintenance (7). In addition, LPL-mediated triglyceride hydrolysis appears to be important in the generation of HDL$_2$ (8), the subfraction of high density lipoprotein that may protect against atherosclerosis (9).

LPL is synthesized in the adipocyte, secreted into the interstitial space, and transported to the capillary endothelium, where it is bound to glycosaminoglycans (10). Since the regulation of this enzyme may take place at numerous sites in LPL processing, studies in vivo or with whole adipose tissue in vitro have not been able to provide insight into LPL regulation at the level of the adipocyte. Thus, a system has been needed where the effects of hormones and other regulators can be directly assessed on adipose cells in vitro.

LPL has been measured in cultured 3T3-L1 cells (11) and in preadipocytes isolated from the stromal-vascular fraction of mouse (12) and rat (13) adipose tissue. These cultures, however, contain preadipocytes and other cell types in various stages of adipocyte differentiation and produce variable amounts of LPL (14, 15). Although isolated rat adipocytes produce LPL (16), studies of regulation in these buffer-suspended cells have been limited due to short-term viability. Methodology has recently been developed to study the regulation of LPL in isolated rat adipocytes (17), which remain viable in culture for several days (17, 18). However, conclusions drawn from studies using rat adipocytes may not relate to human pathophysiology. Thus, a system to study LPL in adipocytes obtained from human adipose tissue biopsies would be desirable.

In this report, we describe the preparation of human adipocytes from a collagenase digestion of omental adipose tissue, and their maintenance in a tissue culture environment for several days. These cells produce and secrete abundant LPL activity, which is regulated by factors such as serum, degree of adiposity, insulin, and insulin-like growth factor-1 (IGF$_1$).

Methods

Preparation of isolated adipocytes. After patients gave informed consent, ~10 g of omental adipose tissue was obtained from 45 healthy subjects (39 females, 6 males) under the age of 50 undergoing elective surgery for nonmalignant conditions. Patients were excluded if there was any evidence of chronic disease, and LPL data were not obtained if the patient was taking any medications known to affect lipid metabolism (such as insulin, oral contraceptives, or glucocorticoids). The mean ($\pm$SE) percentage of ideal body weight (19) of the subjects was 108$\pm$1.9% (range, 90–139%, n = 45). The nature of the surgical procedures varied but were all elective operations such as hysterectomies, ovarian cystectomies, and cholecystectomies. Patients routinely fasted for 8–14 h before surgery and were intravenously hydrated during surgery with lactated Ringer’s solution containing 5% glucose.

Omental adipose tissue was excised and immediately placed into primary cultures in Medium 199. The medium was then changed after 2 h. The cultures were changed every 2–3 days. The method used for these studies is similar to that described by Kupperman and associates (20). The cultures were examined daily for viability using cell counts and trypan-blue exclusion.

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sterile, iced phosphate-buffered saline containing 1.0 mM CaCl₂ and processed within 1 h. By the method of Rodbell (20) the tissue was divided into 1–2 g pieces, minced, and transferred to a polypropylene flask containing 7 ml of collagenase II (2 mg/ml; Worthington Biochemical Corp., Freehold, NJ) in Krebs Ringer bicarbonate buffer with 3 mM glucose and 4% bovine serum albumin. The tissue was digested at 37°C for 60–90 min in a rotary water bath at 140 rpm. The suspension was then passed through a 250-µm pore nylon mesh (Small Parts, Inc., Miami, FL) to remove undigested pieces, and Medium 199 (Gibco Laboratories, Grand Island, NY), containing 10% heat-inactivated whole fetal bovine serum (Reheis Co., Inc., New York) and penicillin/streptomycin (100 U/ml and 100 µg/ml, respectively; Gibco Laboratories) were added to the filtrate. The adipocytes were allowed to float above the heavier stromal-vascular elements, which remained in the medium below the adipocytes. This infranatant was then removed, and the adipocytes were washed twice in serum-containing medium and twice in serum-free medium. A 75-µl aliquot of concentrated cell suspension (1.17±0.10×10³ cells, mean ± SEM, n = 40) was added to 0.75 ml Medium 199, in the presence or absence of serum in sterile 12×75 mm polypropylene tubes. The cells were then incubated at 37°C in a humid 95% air-5% CO₂ environment. In some experiments, cells were cultured in Medium 199 containing 10% dialyzed fetal bovine serum. This serum was dialyzed at 4°C against 0.15 M NaCl using a 3,500-ml wt membrane with two dialysate changes over 24 h.

**Insulin binding.** Specific ¹²⁵I-insulin binding was measured by a modification of the technique of Marshall and Olefsky (21). After incubation at 37°C in Medium 199, as described above, ~1.0×10⁶ cells were washed and incubated in 1.0 ml of minimal essential medium (Gibco Laboratories) containing 25 mM 4-(2-hydroxyethyl)-1-piperazineethanesulfonic acid (Hepes, pH 7.8), 1% bovine serum albumin, and 5% fetal bovine serum (Reheis Co., Inc.). ¹²⁵I-insulin (0.3 ng) was then added in the absence or presence of a large excess (50 µg) of unlabeled porcine insulin (a gift of Dr. Ron Chance, Eli Lilly & Co., Indianapolis, IN). To assess intracellular insulin processing, experiments were performed in the presence of 50 µM chloroquine (Sigma Chemical Co., St. Louis MO), an agent that blocks intracellular insulin degradation and causes an accumulation of intact ¹²⁵I-insulin within cells (21). Termination of the binding reaction was carried out by rapidly centrifuging triplicate 200-µl aliquots of cell suspension in microtubes containing 100 µl of silicone oil. The adipocytes were thus separated from the medium, and specific cell associated ¹²⁵I-insulin was determined.

**Cell size.** An aliquot of adipocyte suspension was washed and sized microscopically by the use of methylene blue (22). In some experiments, sizing was performed with cells suspended in trypan blue. Comparisons between trypan blue and methylene blue staining revealed no difference in cell size between the two techniques.

**Cell counting.** Adipocyte number was determined according to a modification of method III of Hirsch and Gallian (23). Cells were washed and fixed in 2% osmium tetroxide in 0.05 M collidine buffer (made isotonic with saline) for 24 h at 37°C, then taken up in a known volume of 0.15 M NaCl for counting in a model ZB Coulter Counter with a 400-µm aperture (Coulter Electronics Inc., Hialeah, FL).

**Trypan blue exclusion.** By a modification of the technique used for fibroblasts (24), cells were washed with phosphate-buffered saline and incubated with 0.4% trypan blue in 0.9% NaCl for 15 min at 37°C.

**Adipose tissue LPL.** LPL activity was measured against a [¹⁴C]triolein substrate by a modification of a method previously described using cells from the stromal vascular fraction of adipose tissue (25). Three compartments of adipocyte LPL activity were measured: (a) activity secreted into the culture medium (CM), (b) activity released by heparin (HR), and (c) cellular activity extractable in deoxycholate and detergent (EXT), by a modification of the method of Iversius (26). For the measurement of CM, 150 µl of medium bathing the cells was incubated with 50 µl of substrate (described below) for 45 min at 37°C. The reaction was then terminated by the addition of 3.25 ml of the extraction mixture of Belfrage and Vaughan (27), and released fatty acids were extracted for 10 min on an Eberbach Shaker (Eberbach Corp., Ann Arbor, MI). After centrifugation at 2,000 g for 20 min, 0.5 ml of the upper phase was placed in 3.0 ml Aquasol (New England Nuclear, Boston, MA) and counted in a liquid scintillation counter. For HR, the cells were washed twice in Krebs Ringer phosphate buffer and incubated for 45 min at 37°C in 0.4 ml buffer containing 13.5 µg/ml heparin (beef lung; Upjohn Co., Kalamazoo, MI). An aliquot of buffer, containing released LPL activity, was then assayed as described above. To measure EXT, the previously heparin-released cells were transferred to a microhomogenizer, washed twice, and disrupted in the presence of 0.2 ml of solution containing 0.2 M Tris (pH 8.5), 0.73% sucrose, 1.0% albumin, 0.5% deoxycholate (Sigma Chemical Co.), 0.02% Nonidet P₄₀ (neutral detergent; Sigma Chemical Co.), and 125 µg/ml heparin. After the addition of 0.5 ml of Tris-sucrose-albumin-buffer, the suspension was centrifuged at 2,000 g for 30 min, and the middle layer, containing solubilized LPL activity, was assayed as described above.

The substance used in the LPL assay was prepared with 5 mg of unlabeled triolein (Sigma Chemical Co.), 4 µCi [¹⁴C]triolein (Amer sham Corp., Arlington Heights, IL) and 0.24 mg phosphatidyicholine (Applied Science Div., Milton Roy Co., State College, PA). After the addition of mixture of 10% fatty acid-poor albumin (Miles Laboratories Inc., Elkhart, IN), normal human serum, 2 M Tris-HCl buffer (pH 8.2), and distilled water in a ratio of 4:3:5:9:5, the substrate was emulsified by 100 s of sonication (10 s on, 10 s off, for 10 cycles) at 4°C with a cell disrupter 200 model (Heat Systems-Ultrasonics, Inc., Plainview, NY).

**IGF₁.** Two preparations of human IGF₁ were used. Three experiments were carried out with IGF₁ (batch number 165PII) that was highly purified from a Cohn fraction of plasma by Dr. R. E. Hummel (Biochemisches Institut der Universität Zürich, Switzerland) (29), and was a generous gift of Dr. Ted Ciaraldi (University of California, San Diego, La Jolla, CA). This preparation of IGF₁ migrated as a single band upon polyacrylamide gel electrophoresis (28). Further experiments were performed with an IGF₁ analogue (preparation IT-AC3) prepared by means of recombinant DNA technology by AMGen Development, Inc. (Boulder, Colorado) (29). This peptide has a threonine-for-methionine substitution at position 59 but is otherwise homologous with plasma purified IGF₁. In recent studies characterizing this product, receptor binding, biological activity, and immunoreactivity were found to be similar to those of plasma purified IGF₁ (29).

**Statistics.** All data are expressed as the mean±SE with the number of experiments given. Most data were analyzed nonparametrically by the Wilcoxon matched-paired signed-ranks test or the Spearman rank correlation coefficient test. The linear regression for the logarithmic transformation (see Fig. 9) was performed parametrically.

**Results**

The primary cultures of isolated adipocytes formed a homogeneous layer on the surface of the culture medium. To verify that only negligible cell breakage was occurring, cell number was determined on aliquots of adipocytes cultured for up to 72 h. As shown in Table I, no significant change in adipocyte number occurred. Average cell size also remained constant in cultured adipocytes, suggesting that there was no selective breakage of the larger adipocytes. Further evidence of cell membrane integrity included consistently low trypan-blue uptake (Table I).

To assess further the viability of cultured adipocytes, the ability of these cells to bind and internalize insulin was measured. After cells were cultured for up to 72 h, cell-associated ¹²⁵I-insulin was measured after a 2-h incubation at 37°C in the presence or absence of 50 µM chloroquine. Chloroquine has previously been shown to increase adipocyte-
associated $^{125}$I-insulin by inhibiting intracellular insulin degradation and thus trapping internalized insulin within the cell. As shown in Fig. 1, insulin binding in the absence of chloroquine remained constant in cells cultured for up to 72 h. Chloroquine, however, induced increases in cell-associated radioactivity of 51, 65, and 36% at 24, 48, and 72 h after preparation, respectively. This chloroquine effect was consistently present in cultured cells and was of greater magnitude than the minimal effect seen in freshly prepared cells.

LPL was measured as activity secreted into the culture medium (CM), released from cells with 13.5 μg/ml heparin (HR), and extracted from the heparin-released cells by deoxycholate and detergent (EXT). To demonstrate that HR and EXT represent two different components of cellular LPL activity, total cell activity of washed adipocytes was determined and compared with the sum of HR and EXT in parallel cultures. Total cell activity was determined in nonheparin-released cells by the extraction technique (cell disruption in neutral detergent and deoxycholate) described in Methods. As shown in Table II, the sum of HR and EXT consistently and accurately reflected the measured total cell activity, thus demonstrating the additive nature of these two components of cellular activity.

To examine the time course of LPL production by adipocytes, CM, HR, and EXT were measured after various times in culture. In cells incubated in Medium 199 alone, CM remained very low, whereas HR increased progressively, reaching a plateau between 10 and 24 h, then diminishing somewhat at 48 and 72 h (Fig. 2). Cultures that were incubated in medium containing 10% fetal bovine serum similarly showed a progressive increase in HR but also demonstrated a rapid rise in CM (Fig. 2). EXT remained relatively stable over 72 h and was not markedly influenced by the addition of serum. Because CM and HR were at a maximum at 24 h, data on LPL regulation were obtained at this time. Fig. 3 illustrates the effects of 10% serum in paired experiments of adipocytes cultured for 24 h. The mean (±SE) increases in CM and HR were 1.66±0.35 and 1.72±0.32 neq/min per 10$^6$ cells, respect-

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**Table I. Cell Number, Cell Size, and Trypan Blue Exclusion in Cultured Human Adipocytes**

<table>
<thead>
<tr>
<th>Time (h)</th>
<th>Cell number*</th>
<th>Cell size*</th>
<th>Trypan blue exclusion*</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>100 (7)</td>
<td>100 (4)</td>
<td>3.2±1.7 (7)</td>
</tr>
<tr>
<td>24</td>
<td>100±21 (7)</td>
<td>99±5 (4)</td>
<td>0.7±0.2 (3)</td>
</tr>
<tr>
<td>48</td>
<td>108±14 (7)</td>
<td>92±7 (3)</td>
<td>1.0±0.6 (3)</td>
</tr>
<tr>
<td>72</td>
<td>109±20 (7)</td>
<td>99±8 (3)</td>
<td>1.5±0.9 (3)</td>
</tr>
</tbody>
</table>

* Mean±SE.

**Table II. Division of Cellular LPL Activity into HR and EXT**

<table>
<thead>
<tr>
<th>Experiment</th>
<th>HR</th>
<th>EXT</th>
<th>Total cell activity</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Calculated*</td>
<td>Measured†</td>
<td></td>
</tr>
<tr>
<td>1</td>
<td>1.63</td>
<td>1.50</td>
<td>3.13</td>
</tr>
<tr>
<td>2</td>
<td>2.68</td>
<td>1.51</td>
<td>4.19</td>
</tr>
<tr>
<td>3</td>
<td>2.85</td>
<td>1.62</td>
<td>4.47</td>
</tr>
</tbody>
</table>

Activity is expressed as nanoequivalents of free fatty acid released per minute per 10$^6$ cells.* Sum of HR and EXT. † Nonheparin-released cells were disrupted in the presence of neutral detergent and deoxycholate. Solubilized LPL was then assayed (see Methods).

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**Figure 1.** Insulin binding in the presence or absence of chloroquine (CQ). After culturing of cells for the indicated times, 0.3 ng of $^{125}$I-insulin was added and cells were incubated at 37°C for 2 h in the presence or absence of 50 μM CQ. Nonspecific binding was determined for each point by a duplicate tube of cells to which 50 μg of unlabeled insulin was added. Cell-associated activity is expressed as percentage of total $^{125}$I-insulin added per 2 × 10$^7$ cells, after subtraction of nonspecific binding. *P = 0.05.

**Figure 2.** Time course of LPL activity. LPL was measured as (A) CM, (B) HR, and (C) EXT (see Methods). Cells were cultured in Medium 199 in the absence (×) or presence (○) of 10% fetal bovine serum (mean±SE).

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dialyzed serum was determined whether instability of serum characterized. Nevertheless, although the addition of serum to HR did not prolong the half-life further.

The release of LPL activity by heparin was characterized in cells cultured for 24 h. The addition of heparin in concentrations of 1.3 to 130 μg/ml at 37°C produced a prompt release of LPL into the medium, which reached a maximum within 15 min (Fig. 5). This maximal level of LPL was sustained in cells exposed to heparin for up to 180 min. As demonstrated in Fig. 4, the half-life of HR when separated from cells was 135 min. Thus, when cells were exposed to heparin, the released LPL maintained a steady state despite the finite half-life of the enzyme in a cell-free system. This suggests that these isolated adipocytes were able to continuously produce and release new enzyme while exposed to heparin.

To determine whether heparin could deplete cells of this pool of cellular activity, cells that had been previously exposed to heparin for 30, 60, and 120 min were washed, suspended in buffer, and then reexposed to 13 μg/ml of heparin for 45 min at 37°C. Despite the previous exposure to heparin, the second addition of heparin resulted in the release of additional LPL activity (Fig. 6). Although this second response was ~50% less than the initial heparin release, heparin was clearly unable to deplete the cells of HR activity. To determine whether this residual HR was simply due to the use of a submaximal dose of heparin in the first heparin release, the dose-response relationship between heparin concentration and LPL release was studied. As shown in Fig. 7, a broad range of heparin concentrations was found to release equivalent amounts of LPL, and the heparin concentrations used in Fig. 5 (1.3 to 130 μg/ml) released maximal amounts of LPL. Thus, heparin failed to deplete adipocytes of HR, suggesting that HR represents a rapidly replenishable pool of enzyme activity.

Although CM at 24 h (in the presence of serum) had reached a steady state, the amount of secreted activity present in CM varied considerably among primary cultures. To ascertain whether cellular LPL in part determined secreted LPL, paired experiments were conducted with adipocytes cultured for 24 h in the presence and absence of serum. As shown in Fig. 8, serum-stimulated CM correlated positively with control HR at 24 h ($r_s = 0.731$, $n = 23$, $P < 0.001$). In addition, the two components of cellular activity, HR and EXT, correlated with each other at 24 h ($r_s = 0.501$, $n = 22$, $P < 0.05$). In contrast, there was no significant relationship between serum-stimulated CM and EXT ($r_s = 0.355$, $P$, NS). Thus, HR
represented the component of cellular LPL that best predicted the ability of serum to stimulate enzyme secretion.

As illustrated in Fig. 8, HR at 24 h also varied among individual subjects. Therefore, various factors were examined to determine their effect on HR. When body mass index (weight/height$^2$) was plotted as a function of HR at 24 h, a curvilinear relationship was obtained (Fig. 9 A). A logarithmic plot of these data produced the highly significant inverse linear relationship shown in Fig. 9 B. Although fat cell volume correlated significantly with body mass index ($r = 0.775$, $n = 35, P < 0.001$), there was only a weak relationship between fat cell volume and log HR at 24 h ($r = 0.432, n = 22, P = 0.05$). CM and EXT did not correlate significantly with any index of adiposity or with fat cell size. Thus, the degree of subject adiposity played a role in determining steady-state adipocyte HR.

Because it has been suggested to be an important regulator of LPL, insulin was added to cultured adipocytes at the time of preparation over a broad range of concentrations. As shown in Fig. 10 A, 400 ng/ml of insulin produced a mean increase in HR of 0.50±0.14 neq/min per 10^6 cells (36±10%, $n = 22, P < 0.001$) above non-insulin-treated controls. In addition, this concentration of insulin increased EXT by 0.31±0.13 neq/min per 10^6 cells (28±12%, $n = 21, P < 0.01$). Although 400 ng/ml of insulin consistently increased HR, the magnitude of this insulin effect varied. As shown in Fig. 10 B, the increase in HR with 400 ng/ml of insulin correlated directly with control HR at 24 h ($r = 0.594$, $P < 0.01$). In contrast, insulin concentrations of 1, 4, and 40 ng/ml had no effect on either HR or EXT, nor did 400 ng/ml of insulin have any effect on LPL when added to cells cultured in 10% serum. Thus, only high concentrations of insulin stimulated cellular LPL, and this effect on HR was greatest in cultures with a high basal HR.

High concentrations of insulin have previously been shown to cross-react with IGF$_1$ receptors (30, 31). Because the increases in adipocyte LPL occurred only at 400 ng/ml of insulin, these effects may have been mediated through the IGF$_1$ receptor. To examine this question, experiments were conducted in which adipocytes were cultured with insulin (400 ng/ml) as well as with a spectrum of concentrations of IGF$_1$ (ranging from 0.1 to 500 ng/ml). As shown in Fig. 11, IGF$_1$ produced a dose-related increase in HR that was statistically significant at concentrations of 10, 50, and 500 ng/ml. In addition, the magnitude of the increase in HR by IGF$_1$ was similar to that produced by 400 ng/ml of insulin. Thus, concentrations of IGF$_1$ as low as 10 ng/ml produced increases in cellular LPL that paralleled the effects of pharmacologic concentrations of insulin.

**Discussion**

The cellular regulation of adipose tissue LPL has been difficult to characterize, largely because of the lack of a good in vitro model. Although LPL has previously been studied in short-term experiments with rat adipocytes, LPL has never been measured in isolated human adipocytes. Hence, an in vitro system of human cells is needed where the effects of insulin and other potential regulators of LPL can be measured in a controlled environment at the cellular level. Because attempts to use human adipocyte precursors to study LPL have been unsuccessful (25), we have focused our attention on primary cultures of mature adipocytes.

In this study, isolated human adipocytes were obtained from a collagenase digestion of omental adipose tissue. When cultured in Medium 199, these cells remained viable for up to 72 h after preparation, as evidenced by constant cell number, cell size, and low trypan blue uptake. In addition, these cells demonstrated constant $^{125}$I-insulin binding and a chloroquine-mediated increase in cell-associated radioactivity. Chloroquine

**Figure 7.** Heparin dose response. After adipocytes were cultured for 24 h, various concentrations of heparin were added to the medium and cells were incubated for the indicated times. The LPL released by each concentration of heparin is expressed as the percentage of control cells (no heparin added, mean±SE).
evidence of strong process (21), the
has previously been shown to block intracellular insulin degradation, resulting in an increase in cell-associated $^{125}$I-insulin (21). Because internalization of insulin is an energy-dependent process (21), the presence of a chloroquine effect provides strong evidence of functional viability and suggests that these
cells retain the ability to bind, internalize, and degrade insulin. Insulin binding in freshly prepared human adipocytes from subcutaneous and omental fat has been previously reported (32–35) and similar degrees of cell-associated $^{125}$I-insulin were demonstrated. In the present study, insulin binding remained constant, although the magnitude of the chloroquine effect was greater in cultured cells than in freshly prepared cells. This latter observation is consistent with a previous study (34), which determined that freshly prepared human adipocytes did not degrade $^{125}$I-insulin. In addition, Marshall (18) observed an increased chloroquine effect in isolated rat adipocytes between 12 and 48 h in culture. Thus, this increasing chloroquine effect may be due to recovery of normal cell function in culture after dissipation of residual effects of the isolation procedure. Alternatively, the culture medium environment may serve to improve coupling among insulin binding, internalization, and degradation.

Figure 8. Relationship between HR and CM. Cells from 23 subjects were cultured for 24 h in the presence and absence of 10% fetal bovine serum and then assayed for LPL. CM activity in the presence of serum is plotted as a function of HR in the absence of serum.

Figure 9. Effects of body mass index on LPL. (A) HR LPL activity in adipocytes cultured for 24 h in Medium 199 is plotted as a function of subject's body mass index (weight/height$^2$). (B) Logarithmic transformation of data in A.

Figure 10. Effect of insulin on HR LPL. (A) Adipocytes were cultured in Medium 199 for 24 h in the absence and presence of insulin in the indicated concentrations. HR LPL activity is shown for each insulin concentration and expressed as percentage of control HR. (B) Increasing insulin effect with increasing basal activity. The change in HR in response to insulin 400 ng/ml is plotted as a function of control HR at 24 h.

Figure 11. Effect of IGF$_1$ on HR LPL. Adipocytes were cultured overnight in Medium 199 containing the indicated concentrations of IGF$_1$. The effect of insulin (400 ng/ml) was assessed in parallel adipocyte cultures. Data are expressed as the increase in HR activity over the control cultures (mean±SE).
Soon after preparation, LPL was measurable as CM and in the cells as HR, and as EXT after the release by heparin. CM was markedly stimulated by the presence of 10% whole and dialyzed fetal bovine serum. This is in agreement with observations in rat adipocytes (36–38) in which LPL secretion was stimulated by a factor in serum with a molecular weight of >100,000 (38). The effect of serum in the present study may have been not to stimulate secretion but to stabilize LPL. The half-life of CM LPL in the presence of serum at 37°C was 66 min, and the addition of serum to CM from control cultures produced no increase in LPL activity. Hence, LPL may be spontaneously secreted from adipocytes but then inactivated virtually instantaneously in the absence of serum. The addition of serum to HR did not prolong its half-life significantly. However, the HR enzyme is already stabilized by heparin, and serum may not stabilize LPL further. Thus, these data cannot definitively distinguish between a stabilization of LPL by serum and a direct stimulation of secretion. The longer half-life of HR is consistent with the tendency of heparin to stabilize LPL in nonpurified preparations (39).

The release of enzyme activity by heparin occurred quickly and was dose dependent. After 15 min of heparin exposure, LPL activity that was released reached a plateau. Because the HR enzyme has a finite half-life of 135 min, this plateau of activity probably represents a steady state of continuous new enzyme release along with inactivation of enzyme already released. In addition, HR activity was not depleted by a prior maximal release of enzyme activity by heparin. Together, these data suggest that heparin causes the release of LPL from a pool of activity that can be rapidly replenished from cellular stores. This replenishment may result from new enzyme synthesis or may be due to the transfer of activity from one cellular site to another.

The ability of heparin to release LPL in vivo (39), from adipose tissue pieces (40–42), and from isolated cells (17, 37, 40, 43) has long been recognized. Previous studies (2, 17, 40, 42, 44) have also demonstrated that heparin-releasable LPL represents a source of activity that is responsive to regulation by various agents. In the present study, HR at 24 h, but not EXT, correlated strongly with secreted (CM) activity in response to serum. Since LPL must be secreted from adipocytes in order to become localized to the capillary endothelium, HR represents an important functional and regulatable component of cellular LPL activity and may be a key determinant of the LPL that is available in vivo for triglyceride hydrolysis.

HR activity at 24 h was inversely correlated with the subjects’ body mass index. LPL in whole adipose tissue, when expressed per 10⁶ cells, has previously been shown to be directly related to degree of obesity (3, 7). In addition, the study by Schwartz and Brunzell (7) demonstrated that LPL activity in subcutaneous adipose tissue from obese subjects increased in response to weight loss. The subjects in the present study were in general of near-normal body weight, and LPL was measured in isolated cells from omental adipose tissue in culture, making comparisons with other studies difficult. Together, however, the data from whole adipose tissue and from isolated cells may be reconciled by the speculation that with increasing adiposity, a mobilization of LPL from cellular to extracellular stores may occur. Thus, isolated adipocytes would be expected to have less cellular LPL with increasing body weight as demonstrated in the present study, whereas whole adipose tissue would display more total LPL activity, as shown by others (3, 7).

Insulin has long been considered to be a probable hormonal regulator of LPL, although the effects of this hormone have varied and depended upon the system examined. Studies in rats have demonstrated an increase in whole adipose tissue LPL in the fed versus the fasted state (36, 40, 45). This increase in LPL, however, was found in whole fat pads, but only inconsistently in simultaneously isolated rat adipocytes. Whether this increase in whole tissue LPL was due to glucose, insulin, or some other component of the fed state could not be determined. Studies in 3T3-L1 cells have suggested that there is an effect of insulin on both LPL synthesis and secretion (44, 46), whereas other studies in rat adipose tissue or cells have not been able to distinguish between the two effects or have suggested that the predominant effect is on synthesis (47–49). An effect of insulin added in vitro has only recently been demonstrated in isolated rat adipocytes (17). This increase is predominantly on cellular LPL and is inhibited by cycloheximide, suggesting a dependence on protein synthesis.

Although insulin appears to stimulate LPL in animals, the role of insulin in the regulation of human LPL is less clear (50). Decreases in LPL in fat biopsies have been observed when insulin levels are low, as in starvation (51), during hypocaloric feeding (41, 52), and in poorly controlled insulin-dependent diabetes (2). However, wide variations in basal activity create substantial overlap between controls and uncontrolled diabetics (3). Studies in normal humans have also been difficult to interpret. Feeding studies have shown inconsistent increases in LPL activity between 1 and 6 h after feeding (3, 53). Using the euglycemic clamp technique, Sadur and Eckel (42) have shown a stimulatory effect on insulin on whole adipose tissue LPL in normal subjects. In contrast to the results of feeding studies, however, this effect required 6 h of relative hyperinsulinemia (~70 μU/ml) and a simultaneous glucose infusion. Furthermore, attempts to demonstrate an effect of insulin added to whole human adipose tissue in vitro were unsuccessful (54).

In the present study, insulin was added to primary cultures of isolated human adipocytes over a broad range of concentrations. Only at 400 ng/ml, a pharmacologic concentration, was there an increase in cellular (predominantly HR) LPL activity. This effect was consistently seen, although the magnitude of this high-dose insulin effect on HR varied directly with control HR. Physiologic concentrations of insulin had no consistent effect on LPL, regardless of the control activity, and 400 ng/ml of insulin had no effect in the presence of 10% fetal bovine serum. Because high insulin concentrations have been shown to cross-react with IGF₁ receptors in many systems (30, 31), the effect of a spectrum of concentrations of IGF₁ was compared with the effect of 400 ng/ml of insulin. IGF₁ elicited a dose-dependent increase in HR that was statistically significant at a concentration as low as 10 ng/ml. In addition, the magnitude of the increase in HR with IGF₁ was similar to that produced by 400 ng/ml of insulin. This suggests that the effect of high insulin concentration on LPL may actually be mediated through low affinity IGF₁ receptor binding. Consistent with this hypothesis is the lack of an additive effect when insulin is added to cells cultured in 10% serum. Although the increase in HR by serum is probably mediated by a number of factors, the IGF₁ in serum would be expected to contribute to this
effect and thus prevent any further action of insulin on IGF₁ receptors.

The response of human adipocytes to insulin differs from that observed by us in isolated rat adipocytes (17) and in vivo during the euglycemic clamp (42). During a euglycemic clamp, however, the insulin infusion is accompanied by an elevated glucose disposal rate, which may have independent effects on LPL. Furthermore, in rat adipocytes cellular LPL continues to increase with insulin concentrations well above the physiologic range, suggesting that there may be a role for IGF₁, as well as for insulin, in these cells.

Although the concentration of insulin required to stimulate LPL was clearly pharmacologic, it is difficult to estimate what concentration of IGF₁ in vitro truly reflects in vivo physiology. Various studies have measured plasma IGF₁ concentrations in humans and reported concentrations as low at 24 ng/ml in growth hormone-deficient subjects (55) and as high as 722 ng/ml in diabetics with proliferative retinopathy (56). Further complicating the matter, however, is the degree to which IGF₁ is protein bound. Several independent studies failed to detect any free IGF₁ in whole serum (57, 58). In addition, it is not known whether biologically active IGF₁ is the free or the protein-bound moiety (59). Thus, the concentrations of IGF₁ in cell culture that properly reflect physiological concentrations cannot be stated with certainty. However, the concentrations that stimulate LPL in this study closely parallel the concentrations that effectively compete for IGF₁ binding in other systems (31, 60).

As the structure and function of insulin, IGF₁, and insulin-like growth factor 2 have become better characterized, two types of receptor-mediated, anabolic processes have been defined. Rapid cellular responses, such as glucose transport, occur via the insulin receptor, whereas slower, growth-promoting responses are mediated through the insulin-like growth factor receptors (30, 61). Previous studies have demonstrated specific IGF₁ binding to rat adipocytes as well as a stimulation of glucose transport and an inhibition of lipolysis (60). These biological effects, however, are acute responses and indeed were abolished by the removal of the insulin receptors by a brief exposure to trypsin (62). This led to the conclusion that the acute biological effects of IGF₁ and adipocytes are mediated through cross-reactivity with the insulin receptor (61, 62). However, because adipocytes could not previously be sustained for more than a few hours in vitro, it was not possible to measure any long-term biological effects of IGF₁. Because of the demonstrated viability of the cultured adipocytes described herein, this system is better able to study IGF₁ mediated processes. Our data, which demonstrate an effect on LPL by insulin at 400 ng/ml and by IGF₁ at 10 ng/ml, represent a response characteristic of one mediated through an IGF receptor. In addition, preliminary data from our lab, using the IGF₁ analogue described in Methods, confirm that both cultured rat and human adipocytes specifically bind IGF₁ (Kern, P. A., D. Graves, J. J. Van Wyk, and R. H. Eckel, unpublished observations). Whether or not IGF₁ plays an important role in LPL regulation in vivo remains to be determined.

Together, these studies demonstrate that isolated human adipocytes can survive for several days in a tissue culture environment and produce and secrete LPL. Heparin causes the release of LPL activity from a rapidly replenishable pool of cellular activity (HR) which correlates strongly with the secretory response to serum. In addition, HR is regulated by such factors as body mass index and serum. Insulin in a high concentration and IGF₁ at lower concentrations both stimulate HR to a similar degree, indicating that the effect of insulin on LPL in isolated human adipocytes could occur via cross-reactivity with IGF₁ receptors. This method for culturing adipocytes should prove useful for further studies on the regulation of LPL and for investigations into other aspects of human adipocyte cell biology.

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