Role of Insulin in the Regulation of Leucine Kinetics in the Conscious Dog

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ABSTRACT To study the effect of insulin on leucine kinetics, three groups of conscious dogs were studied after an overnight fast (16–18 h). One, saline-infused group (n = 5), served as control. The other two groups were infused with somatostatin and constant replacement amount of glucagon; one group (n = 6) received no insulin replacement, to produce acute insulin deficiency, and the other (n = 6) was constantly replaced with 600 μU/kg per min insulin, to produce twice basal hyperinsulinemia. Hepatic and extrahepatic splanchnic (gut) balance of leucine and α-ketoisocaprate (KIC) were calculated using the arteriovenous difference technique. [3-4,5-3H]Leucine was used to measure the rates (micromoles per kilogram per minute) of appearance (Ra) and disappearance (Rd), and clearance (Cl) of plasma leucine (milliliters per kilogram per minute).

Saline infusion for 7 h resulted in isotopic steady state, where Ra and Rd were equal (3.2±0.2 μmol/kg per min). Acute insulin withdrawal of 4-h duration caused the plasma leucine to increase by 40% (P < 0.005). This change was caused by a decrease in the outflow of leucine (Cl) from the plasma, since Ra did not change. The net hepatic release of the amino acid (0.24±0.03 μmol/kg per min) did not change significantly; the arterio-deep femoral venous differences of leucine (−10±1 μmol/liter) and KIC (−12±2 μmol/liter) did not change significantly indicating net release of the amino and ketoacids across the hindlimb. Selective twice basal hyperinsulinemia resulted in a 36% drop in plasma leucine (from control levels of 128±8 to 82±7 μmol/liter, P < 0.005) within 4 h. This was accompanied by a 15% reduction in Ra and a 56% rise in clearance (P < 0.001, both). Net hepatic leucine production and net release of leucine and KIC across the hindlimb fell markedly. These studies indicate that physiologic changes in circulating insulin levels result in a differential dose-dependent effect on total body leucine metabolism in the intact animal. Acute insulin withdrawal exerts no effect on leucine rate of appearance, while at twice basal levels, insulin inhibited leucine rate of appearance and stimulated its rate of disappearance.

INTRODUCTION

The effects of insulin on protein turnover in the intact animal (1, 2) or man (3–6) have been primarily obtained from nitrogen balance studies. In the absence of insulin, the untreated insulin-dependent diabetic (7) or depancreatized dog (2) has enhanced nitrogen excretion, which is reversed with insulin replacement. These studies, however, fail to distinguish between changes in the rates of protein synthesis and degradation. In vitro studies in rat skeletal or cardiac muscles have established that insulin stimulates protein synthesis by enhancing peptide chain initiation and inhibits protein breakdown by a yet unknown mechanism (8). In the liver, insulin exerts an antikatabolic effect, primarily due to stabilization of liver lysosomes (9, 10). It is not clear from these studies, however, whether it is the effect of insulin on protein synthesis or degradation that predominates in vivo.

The effect of insulin on branched-chain amino acid homoeostasis in the intact animal or man is not very well delineated. In high amounts, insulin suppresses the plasma concentration of most of the amino acids,
and particularly leucine, isoleucine, and valine (3). On the other hand, in situations of relative or absolute insulin deficiency, as in obesity (7), starvation (4, 11), and diabetes mellitus (7) the concentrations of the branched-chain amino acids are elevated. It is not clear however, whether the changes observed in the plasma concentrations of the branched-chain amino acids are due to changes in their rates of appearance into or disappearance from the plasma compartment. Leucine, being an essential amino acid, its only source after an overnight fast must be from protein, and therefore changes in its rate of turnover in plasma will reflect changes in protein turnover. The studies presented herein were undertaken to examine the effect of physiologic changes in insulin levels on total leucine flux and on leucine balance across the hindlimb, the splanchnic and hepatic beds. To define the effects of acute insulin deficiency, independent of changes in glucagon secretion, somatostatin (SIF)1 was used to inhibit the endocrine pancreas and glucagon was replaced to basal levels via the intraportal route. The effects of twice basal insulin, i.e., hyperinsulinemia, were also examined in a similar fashion. SIF was used to inhibit the endocrine pancreas; glucagon was replaced to basal levels, while insulin was replaced to twice basal levels via the intraportal route.

METHODS

Experiments were carried out on 17 overnight fasted mongrel dogs (20–26 kg) of either sex, which had been fed a regular dog chow diet (Wayne Dog Food, Allied Mill, Inc., Chicago, IL) for 3 wk before their use. Silastic catheters were implanted under general anesthesia in the portal vein, proximal splenic vein, hepatic vein, and femoral artery, as described (12), 17 d before the study. After the catheters were implanted they were filled with saline containing heparin (200 U/ml, Abbott Diagnostics, Diagnostic Products, North Chicago, IL) and their free ends were knotted and placed in a subcutaneous pouch so that complete closure of skin incisions was possible. 1.6 d after surgery, blood was drawn from each animal to determine leukocyte count and hematocrit. Only animals that had a hematocrit > 38%, a leukocyte count < 15,000/mm³, a good appetite (consuming greater than two-thirds of their daily rations), and normal stools were used in the studies to be described. On the day of the study the catheters were removed from the subcutaneous pockets through skin incisions made under local anesthesia (1% lidocaine, Astra Pharmaceutical Products, Inc., Worcester, MA). The contents of each catheter were aspirated and saline was infused through them at a slow rate (0.1 ml/min) until the experiment began. The portal vein, hepatic vein, and arterial catheters were used for blood sampling, while the splenic vein catheter was used for hormone infusion. Angiocatheters (No. 18 gauge, Abbott Diagnostics) were inserted percutaneously into the two cephalic veins as well as the right saphenous vein. An angiocatheter (No. 18 gauge) was introduced into the left saphenous vein and threaded into the deep femoral vein and used for blood sampling across the hindlimb. The right and left cephalic veins were used for the [3H]leucine and cardiogreen infusion (0.075 mg/m² per min), respectively, and the saphenous vein was used for SIF infusion. After completion of preexperimental preparation, the conscious dogs were placed in a Pavlov harness and were allowed to rest for ~1 h before the beginning of the intravenous infusion of the tracer.

Experimental design. All experiments consisted of a 180-min tracer equilibration period (from −210 to −30 min) followed by a 30 min basal period (from −30 to 0 min) and a 240-min period of hormone perturbation (0–240 min). A constant infusion of L-4,5-[3H]leucine was started at −210 min and continued throughout the study.

SIRF (0.8 µg/kg per min) was infused from 0 to 240 min in combination with intraportal glucagon and/or insulin. Glucagon was infused at a constant rate of 0.65 ng/kg per min and insulin, when used, was infused at a constant rate of 600 µU/kg per min.

Three types of experiments were performed: (a) Group I (n = 5)—saline was infused to assess the stability of the animals during the experimental period. (b) Group II (n = 6)—SIF plus intraportal glucagon, to determine the effect of acute (4 h) selective insulin lack. (c) Group III (n = 6)—SIF plus intraportal glucagon and insulin, to examine the influence of a twofold rise in circulating insulin on leucine kinetics; during this study glucose (50% dextrose in water) was infused at a variable rate to maintain euglycemia. The L-4,5-[3H]leucine infusion differed amongst the three groups and amounted to (18.9±0.5)×10⁴, (12.8±0.6)×10⁴, and (3.34±0.4)×10⁴ dpm/min for groups I, II, and III, respectively.

Processing of blood samples. Blood samples were drawn every 10 min throughout the basal period and every 30 min thereafter. The collection and immediate processing of blood samples have been described (12). In group III, plasma glucose was monitored every 15 min and the exogenous infusion of dextrose solution was adjusted accordingly to maintain euglycemia. Plasma immunoreactive glucagon was assayed using Unger’s (13) 30K antibody obtained from the University of Texas, Southwestern Medical School. Immunoreactive insulin was measured by the Sephadex-bound antibody procedure (Pharmacia Fine Chemicals, Piscataway, NJ) (14). Indocyanine green (cardiogreen) was measured spectrophotometrically at 810 nm for the determination of hematric blood flow (15).

The radioactivity and the concentration of [3H]leucine in plasma samples were determined by rapid column chromatography procedure. The system consists of two identical jacketed chromatographic columns (0.9 cm i.d.) packed with resin BP-AN-6 (Benson Co., Reno, NV). Separate buffer pumps are used on each column and are fed from the common buffer through three-way solinoid operated valves. Flow of the two pumps is equal and the effluents from the two columns lead to a four-way valve arranged so that when one column is directed to a fraction collector through a “T” fitting, the other column effluent is directed to waste. The effluent stream directed to the fraction collector is continuously sampled from the T fitting by an autoanalyzer (Technicon Instruments Corp., Tarrytown, NY) that reacts the stream with ninhydrin for quantitation of the leucine peak. The current method uses the following conditions: column height 12.2 cm, sodium citrate buffer (0.2 M) with pH of 3.24 and buffer flow rate of 2.2 ml/min. Under these con-

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1 Abbreviations used in this paper: KIC, α-ketoisocaproic acid; Ra, rate of appearance of plasma leucine; Rd, rate of disappearance of plasma leucine; SIF, somatostatin.
concentrations, methionine and the branched-chain amino acids are well separated with peak time for leucine at 27 min, depending on the pH of the buffer. Quantitation of leucine concentration using this method was carried out by measuring peak heights, having first established that peak height was linear with concentration. A factor was established, using leucine standards of various concentrations, to convert peak heights to micromole leucine per milliliter. Radioactivity in the fraction collected during elutions of the leucine peak was determined using established liquid scintillation techniques. The disintegrations per minute per milliliter of the effluent were multiplied by the total volume of the fraction to obtain the total radioactivity in the sample applied. This was further corrected for the fractional recovery of [3H]leucine determined for each run. In addition, the specific radioactivity of [3H]leucine in the infusate solution was checked in each experiment in a similar fashion to the plasma samples. The coefficients of variation, in duplicate pairs (n = 12), were 2.3% for arterial plasma leucine concentration and 3.1% for leucine specific activity.

The concentration of α-ketoisocaproic acid (KIC) in plasma was measured by high performance liquid chromatography (HPLC) (16). In brief, plasma is precipitated with 0.6 M perchloric acid and the filtrate is reacted with 2,4-dinitrophenylhydrazine (1:1, vol/vol); the filtrate is then extracted with an equal volume of cyclohexane/heptane/ether solution (10:1:1) (17). The organic layer is removed, dried, and kept at 4°C overnight. The sample is then reconstituted with methanol and run on the HPLC using reverse-phase C-18, μ-Bondapak column (Waters Instruments, Inc., Rochester, MN) and eluted with 50 mM Na₂HPO₄ buffer, pH 7.0 at a rate of 2 ml/min. The effluent is monitored spectrophotometrically at 365 nm. This method allows the separation of ketoacids of leucine and valine, KIC, and α-keto-β-methyl-valeric acid, at 3.5 and 4.5 min, respectively.

Materials. 1-4,5[3H]Leucine (New England Nuclear, Boston, MA) was used as the radioactive tracer (55 μCi/μmol). The purity of the tracer was determined for each batch of [3H]leucine used; 98–99% of the radioactivity infused was in leucine. Insulin and glucagon were purchased from Eli Lilly & Company (Indianapolis, IN); trasylool from FBA Pharmaceuticals Inc. (New York) and [125I]-glucagon was obtained from Novo Research Institute (Copenhagen, Denmark). Phadebas insulin radioimmunoassay kit was purchased from Pharmacia Fine Chemicals, Inc. All hormones solutions were prepared with normal saline and contained 0.3% bovine albumin.

Tracer methods and calculations. The rates of leucine flux in and out of the plasma compartment were determined by the constant tracer infusion method. At maximal steady-state ("plateau") arterial leucine specific activity, the amount of [3H]leucine entering the pool is equivalent to the amount of leaving the pool, assuming that the pool size remains constant. Thus, under steady-state conditions, the rate of appearance (Ra) of leucine in the plasma compartment is equal to its rate of disappearance (Rd) and is equivalent to the rate of isotope infusion in disintegrations per minute per kilogram divided by the plateau arterial specific activity in disintegrations per minute per micromole per 10⁶. This method assumes that the reentry of isotope from the protein pool is negligible during the infusion period. Clearance (Cl) of leucine from the plasma compartment is calculated by dividing Rd (micromoles per kilogram per min) by the plasma leucine concentration (micromoles per milliliter).

During nonsteady-state conditions, calculations of the rates were carried out according to the method of Wall et al. (18) as simplified by DeBodo et al. (19). This method is based on a single compartment analysis in which it is assumed that rapid changes in the specific activity and concentration of leucine did not occur uniformly within the entire leucine pool.

Net hepatic, splanchnic and "gut" (extrahepatic splanchnic tissue) balance of leucine and other substrates were calculated as follows:

Net hepatic balance

\[ = 0.72(HV-PV)_{conc} + 0.28(HV-A)_{conc} \times EHPF \]  (1)

Net splanchnic balance = (HV-A)_{conc} × EHPF

Net "gut" balance = (PV-A)_{conc} × 0.72EHPF,  (3)

where A, PV, and HV represent the plasma concentration of leucine in the artery, portal, and hepatic veins, respectively; EHPF represents the estimated hepatic plasma flow; it is estimated that portal vein supplies 72% and the artery 28% of the hepatic blood flow (12).

Statistical analyses were performed using the Student's t test, the paired t test, and, where applicable, analysis of variance. The data in the figures and tables are expressed as mean±SEM.

RESULTS

Steady-state leucine kinetics. Fig. 1 shows the plasma leucine concentration and arterial leucine specific activity in the saline-infused control group. There was no significant change in plasma leucine concentration during 7 h of saline infusion. Leucine specific activity did not reach steady-state levels until 180 min after the start of the infusion. Because of this fact all studies were performed allowing a 3-h isotopic equilibration period, a 0.5-h basal period, followed by a perturbation period as outlined in the experimental design.

To establish whether SRIF has any effect on leucine metabolism and kinetics in the conscious dog, we measured plasma leucine concentration and flux in two overnight fasted dogs for 6 h. When SRIF was infused with intraportal replacement of basal insulin (225 μU/kg per min) and glucagon (0.65 ng/kg per min), plasma leucine (113±3 μmol/ml) and its specific activity (254±21 dpm/μmol per 10⁶) showed no significant change from saline controls (plasma leucine = 118±6 μmol/liter and specific activity = 261±19 dpm/μmol per 10⁶).

Table I summarizes the mean values of plasma leucine and KIC obtained during the basal period from the three study groups. There is a stepwise dilution of the arterial specific activity across the splanchnic bed. The portal and the hepatic vein specific activities were 22 and 36%, respectively, lower than that of the artery. The dilution of the specific activity across the gut is mainly due to the exchange of labeled leucine for unlabeled molecules. This is evident by the fact that the ratio of portal venous specific activity to that of the
artery was 0.79±0.01, while the ratio of plasma leucine concentrations in the two vessels was 0.95±0.01. To determine whether exchange of label occurred across the liver during the basal period, the specific activity of leucine entering the liver was compared with that leaving the liver and the ratio was 0.78±0.02. Because the ratio of unlabeled leucine in the hepatic vein to that entering the liver was 1.07±0.01, then about one-third of the dilution occurring across the liver must have resulted from a net addition of unlabeled leucine to the plasma (0.24±0.03 μmol/kg per min, P < 0.05; Table II) and the remainder from the exchange of labeled for unlabeled molecules as plasma traversed the liver. The mean rate of [3H]leucine infusion was $591 \times 10^3$ dpm/kg per min, whereas the mean rates of hepatic and gut [3H]leucine uptake were $76.8 \times 10^3$ and $81.6 \times 10^3$ dpm/kg per min, respectively. Thus, 27±3% of the infused label was removed by the splanchnic bed (14±2% by the gut and 13±2% by the liver) indicating that ~70% of leucine flux out of the plasma compartment occurred in the extrasplanchnic tissues.

Plasma leucine concentrations in the three vessels were four- to sevenfold higher than those of KIC. The concentrations were such that there was a consistent net hepatic uptake of KIC (0.20±0.08 μmol/kg per min) and an equivalent net hepatic output of leucine (0.24±0.03 μmol/kg per min); the extrahepatic splanchnic tissues (the gut) extracted both KIC and leucine (Table II). Although these values were very small, they were all statistically significant (P < 0.05). Arteriovenous differences across the hindlimb were $-10\pm1$ and $-12\pm2$ μmol/liter for both leucine and KIC, respectively. Despite this net release of the amino acid.
and ketoacids, there was net extraction of [\textsuperscript{3}H]leucine 16±1.2% as it traversed the hindlimb (data not shown).

**Effects of insulin on leucine kinetics.** Fig. 2 shows the plasma insulin and glucagon concentrations for the three experimental groups. The intraportal replacement of glucagon resulted in comparable glucagon levels to those obtained in the saline-infused control group. When insulin was not infused, there was an 80% drop in plasma insulin from basal values of 12±1 to 2±1 μU/ml (P < 0.001) during the experimental period; the radioimmunoassay used for insulin measurement does not distinguish these latter values from zero. Infusion of intraportal insulin (600 μU/kg per min) resulted in just over a twofold (28±2 μU/ml, P < 0.001) increase in the peripheral insulin levels.

The effect of selective changes in basal insulin on leucine metabolism can be seen in Fig. 3. The acute withdrawal of insulin resulted in a progressive increase in plasma leucine, from the basal values of 131±9 to 187±12 μmol/liter (P < 0.005) by the end of 4 h. No significant change in arterial leucine specific activity was observed during this period as shown in panel B. On the other hand, selective hyperinsulinemia resulted in rapid changes in leucine metabolism, such that plasma leucine fell by 36% (from basal levels of 128±8 to 82±7 μmol/liter by 240 min, P < 0.005) and leucine specific activity rose by 21% (from basal values of 44±5 dpm/μmol per 10\(^3\) to 53±9 dpm/μmol per 10\(^3\), P < 0.005).

The effect of physiologic changes in insulin levels on the rates of appearance and disappearance of leucine into and out of the plasma compartment are shown in Table III. In animals that received only saline, there was no significant change in Ra during the course of the experiment. Similarly there was no significant change in Ra with acute induction of insulin deficiency. In contrast, the induction of hyperinsulinemia resulted in a decline in Ra, such that after 60 min it had fallen by 15% (from 3.2±0.1 μmol/kg per min during the basal period to 2.73±0.15 μmol/kg per min, P < 0.001) and remained depressed for the rest of the experimental period. The clearance of leucine from the plasma compartment (Table III) dropped significantly when insulin was withdrawn (from 25±3 ml/kg per min during the basal period to 15±0.3 ml/kg per min by the end of 4 h, P < 0.001), while with hyperinsulinemia, there was a 65% rise in clearance (from 26±0 to 42±5 ml/kg per min by 4 h, P < 0.001). The changes in Rd were such that Ra minus Rd was always positive with insulin deficiency and always negative with hyperinsulinemia (Fig. 4). These changes were not significant at any particular interval, however, the cumulative changes over the entire experimental period (i.e., 4 h) were significant in both groups (cumulative Ra-Rd was +0.58±0.08 μmol/kg per min with insulin deficiency and −0.69±0.12 μmol/kg per min with twice basal hyperinsulinemia, P < 0.001 for both). No significant changes were observed for the saline-infused control group.

Saline infusion or acute insulin withdrawal for 4 h,
TABLE II
Leucine and KIC Balance Across the Hindlimb, Gut, Splanchnic, and Hepatic Beds of Overnight Fasted (16–18 h) Conscious Dogs*

<table>
<thead>
<tr>
<th></th>
<th>Basal period</th>
<th>Experimental period</th>
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</thead>
<tbody>
<tr>
<td></td>
<td>Saline</td>
<td>Saline</td>
</tr>
<tr>
<td>Leucine</td>
<td></td>
<td></td>
</tr>
<tr>
<td>Net hepatic balance</td>
<td>-0.24±0.03(\mu)mol/kg/min)</td>
<td>-0.25±0.04(\mu)mol/kg/min)</td>
</tr>
<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
</tr>
<tr>
<td>Net splanchnic balance</td>
<td>-0.16±0.03(\mu)mol/kg/min)</td>
<td>-0.19±0.03(\mu)mol/kg/min)</td>
</tr>
<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
</tr>
<tr>
<td>Net gut balance</td>
<td>+0.09±0.01(\mu)mol/kg/min)</td>
<td>+0.07±0.03(\mu)mol/kg/min)</td>
</tr>
<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
</tr>
<tr>
<td>Hindlimb A-DV differences</td>
<td>-10±1(\mu)mol/liter)</td>
<td>-13±2(\mu)mol/liter)</td>
</tr>
<tr>
<td></td>
<td>(9)</td>
<td>(3)</td>
</tr>
<tr>
<td>KIC</td>
<td></td>
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</tr>
<tr>
<td>Net hepatic balance</td>
<td>+0.20±0.08(\mu)mol/kg/min)</td>
<td>+0.21±0.06(\mu)mol/kg/min)</td>
</tr>
<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
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<tr>
<td>Net splanchnic balance</td>
<td>+0.37±0.07(\mu)mol/kg/min)</td>
<td>+0.39±0.06(\mu)mol/kg/min)</td>
</tr>
<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
</tr>
<tr>
<td>Net gut balance</td>
<td>+0.17±0.04(\mu)mol/kg/min)</td>
<td>+0.18±0.05(\mu)mol/kg/min)</td>
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<tr>
<td></td>
<td>(17)</td>
<td>(5)</td>
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<tr>
<td>Hindlimb A-DV differences</td>
<td>-12±2(\mu)mol/liter)</td>
<td>-13±2(\mu)mol/liter)</td>
</tr>
<tr>
<td></td>
<td>(9)</td>
<td>(3)</td>
</tr>
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</table>

* For each dog, the four values obtained during the basal period and the five values during the last 2 h of the experimental period were averaged. The values during the basal period did not differ among the three study groups and the values were pooled. The numbers in parentheses denote the number of dogs studied. (−) denotes release and (+) denotes uptake. All values are expressed as mean±SEM.

1 \(P < 0.05\)

§ \(P < 0.05\)

A-DV differences, arterio-deep femoral venous differences of leucine.

had no effect on the net balance of KIC and leucine across the liver, gut, or hindlimb as shown in Table II. Twice basal hyperinsulinemia resulted in a significant reduction in net hepatic output of leucine (to 0.04±0.04 \(\mu\)mol/kg per min during the last 2 h of the infusion), while net hepatic uptake of KIC persisted of 0.20±0.09 \(\mu\)mol/kg per min \((P < 0.05)\); the specific activity of \(^3\)Hleucine in the hepatic vein increased from 34±5 dpm/\(\mu\)mol per \(10^3\) in the basal period to 42±6 dpm/\(\mu\)mol per \(10^3\) by 4 h of insulin infusion. Concomitantly, arterio-deep femoral venous differences of leucine and KIC were −2±1 and 0±2 \(\mu\)mol/liter, respectively, indicating a marked reduction in the net release of the amino and ketoacids across the hindlimb.

The effect of saline and hormone infusion on plasma glucose were consistent with those previously reported from this laboratory (22). During the saline control, and during the infusion of SRIF plus intraportal re-placement of basal insulin and glucagon, plasma glucose levels remained stable. With twice basal hyperinsulinemia, euglycemia was maintained by an infusion of exogenous glucose. Selective insulin deficiency resulted in plasma glucose levels of 150–170 mg%, which over a period of 4 h progressively dropped to 120±4 mg% (or 23±2 mg% above basal values).

**DISCUSSION**

This study addresses the effects of physiologic changes in insulin concentrations on leucine metabolism in the intact animal, with special reference to the role of basal insulin in maintaining normal whole body protein homeostasis. Furthermore, the combination of the arteriovenous difference and isotopic techniques used in the present study provided valuable information regarding leucine flux that could not be obtained by either technique alone.
The flux of an amino acid, defined as the rate of entry or exit from a pool can be estimated by the continuous infusion of the isotopically labeled amino acid. At steady state, the rate of appearance (Ra), or its equivalent the rate of irreversible disposal (Rd) can be calculated using the plateau-specific activity of the labeled moiety in the infused pool regardless of the volume of the pool or the configuration of the system.

Ra determined in this fashion measures the rate of input of unlabeled substrate into the infused pool and does not necessarily represent the total rate of production (20). In the current study, we have measured the influx of unlabeled leucine into the plasma pool from all sources. The leucine turnover rates measured during the basal period in all three groups of dogs were in agreement with those previously reported by us (11) and by others (21).

Additionally, the application of the arteriovenous difference technique across the splanchnic bed and the hindlimb allows us to calculate the contribution of these tissues to total leucine turnover. During the steady-state control period, ~30% of the infused tracer was extracted across the splanchnic bed; half by the liver and the other half by the gut. This is slightly lower than alanine flux, of which >50% is accounted for by the splanchnic bed and particularly by the gut (22). During the transit of blood from the artery to the deep femoral vein, there was a net release of leucine; this was accompanied by a net extraction of leucine radioactivity showing a process of exchange occurring across the hindlimb. If we assume that plasma flow was 0.101 liter/min (21), then from the change in the specific activities of the artery (122±25 dpm/μmol per 10^9) and the deep veins (95±20 dpm/μmol per 10^9), at least 36% of the dilution occurring across the hindlimb must have resulted from the net addition of non-radioactive leucine to plasma.

The small, but consistent net release of leucine from the splanchnic bed and the hindlimb, raises an interesting question concerning the site of net consumption

![Figure 2](image_url)
of leucine. Since after an overnight fast, both muscle and splanchnic beds contribute leucine to the plasma compartment, then other sites must utilize leucine during that period. Recent data from this laboratory, obtained from a similarly fasted group of dogs, have demonstrated the capacity of the kidneys to utilize leucine at a basal rate of 0.25±0.03 μmol/kg per min (15). Furthermore, in vitro evidence exists for the metabolism of these amino acids by the heart (23), adipose (24), and nervous tissues (25).

With the use of isotope tracers, a precise quantification of protein synthesis and degradation cannot be made from plasma specific activities alone. Assumptions about the relation of the plasma to the intracellular specific activities in the different tissues must be made, or an independent estimate of the precursor pool specific activity is required. Despite these limitations, changes in the rate of appearance of leucine into the plasma compartment should reflect changes in protein degradation; this is based on the fact that

**FIGURE 3** The arterial plasma leucine concentration and the percent change from basal of leucine specific activity in 16-18-h fasted conscious dogs maintained on infusions of SRIF (0.8 μg/kg per min) and concurrent intraportal infusions of glucagon alone (0.65 ng/kg per min—no insulin, broken line) or in combination with insulin (600 μU/kg per min—elevated insulin, solid line). The changes in the concentration of plasma leucine from control levels were significant (P < 0.05) from 90 min on, when using paired or unpaired t tests. The changes in leucine specific activity were significant (P < 0.05) only in the elevated insulin group from 60 min on (paired or unpaired t tests), when compared to both saline control or no insulin groups.
leucine is an essential amino acid and its only source after an overnight fast must be from protein.

During the period of hormonal perturbations, with changing concentration and/or specific activity, another caveat needs to be kept in mind; namely, that calculations of Ra, Rd, and clearance are dependent on the assumed volume of distribution. However, although assumption of an incorrect volume of distribution would introduce errors in the absolute values for Ra, Rd, and clearance it would not change either the direction of changes of these parameters or the relative contribution of each to the observed changes in leucine flux into and out of the plasma compartment. Keeping these considerations in mind, one can draw valid conclusions about the effect of changes in insulin concentration on the relative contribution of changes in Ra, Rd, and clearance on the flux of plasma leucine and by inference on protein degradation and synthesis.

The data obtained using the arteriovenous difference technique represent a balance between the factors influencing protein synthesis and protein breakdown across the appropriate beds. The net negative nitrogen balance observed across both skeletal muscle and splanchnic beds of 18-h fasted dogs, similar to those previously observed across the human forearm (28–30), suggest that at basal levels, insulin has a function to maintain protein synthesis slightly less than that of protein breakdown.

During acute insulin withdrawal, the isotopic data indicated that the rise in plasma leucine was not due to an increase in leucine rate of appearance in the plasma compartment but to a relative decrease in its rate of disappearance out of plasma (Fig. 4). However, we cannot rule out the possibility that a minimum increase in Ra did occur, which in the present study would have been masked by a slight increase in recycling of 3H-label from the protein compartment. Nevertheless, most of the observed change was mainly due to the 40% decrement in leucine clearance (Table III). Since the observed changes in plasma leucine concentrations over the 4-h experimental period would only require the net addition of 0.06 μmol/kg per min of leucine, the arteriovenous technique might not be sensitive enough to pick these changes.

It is apparent that the initial event in twice basal hyperinsulinemia is a rapid fall in Ra. Rd also dropped in conjunction with Ra, however, the difference between the rates of entry and outflow of leucine into and out of the circulation (denoted by Ra minus Rd in Fig. 4) had decreased significantly by the end of the 4-h infusion. Under these conditions, it could be calculated that the change of Ra accounted for ~50% of the change in the net balance, while Rd accounted for the rest. In this regard the net output of cold leucine by the liver and of both KIC and leucine by the hindlimb became negligible; this indicates that either the uptake of leucine by these tissues had increased and/or its release had decreased. Thus, when the findings obtained by the arteriovenous difference technique are combined with those obtained by the tracer method, they would suggest that with hyperinsulinemia, both the liver and the hindlimb tissues (most likely skeletal muscle) participate in the observed decrement in leu-

### Table III

**Effect of Physiologic Changes in Insulin Levels on Plasma Leucine Turnover in the Conscious, Overnight (16–18 h)-Fasted Dogs**

<table>
<thead>
<tr>
<th>Time interval</th>
<th>Saline control (n = 5)</th>
<th>No insulin (n = 6)</th>
<th>Twice insulin (n = 6)</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Ra</td>
<td>Rd</td>
<td>Clearance</td>
</tr>
<tr>
<td>Basal period</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>−30–0</td>
<td>3.11±0.19</td>
<td>3.18±0.15</td>
<td>25±2</td>
</tr>
<tr>
<td>Experimental period</td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>0–30</td>
<td>3.18±0.16</td>
<td>3.17±0.16</td>
<td>24±2</td>
</tr>
<tr>
<td>30–60</td>
<td>3.09±0.21</td>
<td>3.18±0.20</td>
<td>25±1</td>
</tr>
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<td>60–90</td>
<td>3.13±0.17</td>
<td>3.10±0.18</td>
<td>24±3</td>
</tr>
<tr>
<td>90–120</td>
<td>3.15±0.16</td>
<td>3.14±0.16</td>
<td>26±3</td>
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<tr>
<td>120–150</td>
<td>3.16±0.14</td>
<td>3.13±0.16</td>
<td>23±4</td>
</tr>
<tr>
<td>150–180</td>
<td>3.19±0.17</td>
<td>3.24±0.21</td>
<td>24±2</td>
</tr>
<tr>
<td>180–210</td>
<td>3.20±0.16</td>
<td>3.22±0.20</td>
<td>25±3</td>
</tr>
<tr>
<td>210–240</td>
<td>3.18±0.18</td>
<td>3.14±0.15</td>
<td>25±2</td>
</tr>
</tbody>
</table>

*Saline, SRIF plus basal intraportal glucagon, or SRIF, basal intraportal glucagon, and twice basal insulin were infused from 0 to 240 min. All values are expressed as mean±SEM.

1 Significance from corresponding values in the basal period, P < 0.05.
FIGURE 4 The cumulative differences between the rates of appearance (Ra) and disappearance (Rd) in 16-18-h fasted conscious dogs during saline infusion or during the infusion of SRIF (0.8 μg/kg per min) and concurrent intraportal infusion of glucagon alone (0.65 ng/kg per min—insulin deficiency), or with insulin (600 μU/kg per min—elevated insulin). The values represent an average of the cumulative differences for each time period per dog and are expressed as mean±SEM. The number of dogs studied in each group are the same as those shown in Table III. Changes from control levels were significant (P < 0.05) in the insulin deficiency and elevated insulin groups, from 60 min on when using paired or unpaired t tests.

cine rate of appearance and in the increase in its rate of clearance from the plasma compartment.

Though the rates of oxidation of leucine were not measured in the present study, it would be reasonable to assume that, if anything, leucine oxidation had increased with acute insulin withdrawal, in accordance with studies on isolated skeletal muscle preparations (8, 26) and declined in response to twice physiologic levels of insulin. Recent studies in man have shown that a comparable twofold rise in peripheral insulin levels have resulted in a 60% drop in leucine oxidation. 2 Since Rd reflects changes occurring in protein synthesis and/or in the rate of leucine oxidation, it could be speculated that the initial event in selective insulin deficiency is not an increase in protein breakdown, while a slight physiologic rise in insulin may exert a dual effect on total body protein turnover, namely, an enhancement of protein synthesis and an inhibition of protein breakdown.

2 Abumrad, N. N. Unpublished observations.
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