Inverse Relationship of Leucine Flux and Oxidation to Free Fatty Acid Availability In Vivo

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Abstract

To determine the effect of fatty acid availability on leucine metabolism, 14-h fasted dogs were infused with either glycerol or triglyceride plus heparin, and 46-h fasted dogs were infused with either nicotinic acid or nicotinic acid plus triglyceride and heparin. Leucine metabolism was assessed using a simultaneous infusion of L-[4,5-3H]leucine and α-[1-14C]ketoisocaproate. Leucine, α-ketoisocaproate (KIC), and totalleucine carbon (leucine plus KIC) flux and oxidation rates were calculated at steady state. In 14-h fasted animals, infusion of triglyceride and heparin increased plasma free fatty acids (FFA) by 0.7 mM (P < 0.01) and decreased leucine (P < 0.01), total leucine carbon flux (P < 0.02), and oxidation (P < 0.05). The estimated rate of leucine utilization not accounted for by oxidation and KIC flux decreased, but the changes were not significant. During glycerol infusion, leucine and KIC flux and oxidation did not change.

In 46-h fasted dogs, nicotinic acid decreased FFA by 1.0 mM (P < 0.01) and increased (P < 0.05) the rate of leucine and total leucine carbon flux, but did not affect KIC flux. Leucine oxidation increased (P < 0.01) by nearly threefold, whereas nonoxidized leucine utilization decreased. Infusion of triglyceride plus heparin together with nicotinic acid blunted some of the responses observed with nicotinic acid alone.

In that changes inoxidation under steady state condition reflect changes in net leucine balance, these data suggest that FFA availability may positively affect the sparing of at least one essential amino acid and may influence whole body protein metabolism.

Introduction

Prolonged fasting results not only in depletion of carbohydrate and fat stores, but also in the progressive loss of body protein ultimately resulting in inanition and death. Therefore, any mechanism which improves protein balance by decreasing proteolysis and/or oxidation of essential amino acids during inadequate caloric intake may improve the individual's duration of survival. Such relative sparing of body protein in fasted man has been proposed on the basis of decreased excretion of urinary nitrogen and 3-methylhistidine in obese subjects fasted 21 d (1–3). In most mammalian species, body protein cannot be totally conserved during prolonged periods of fasting, in contrast to the hibernating bear (4).

A variety of in vitro (5–7) and in vivo (8–13) experiments using isotopes of leucine have been carried out to determine the effects of fasting on amino acid and protein metabolism; however, the results of these studies are not entirely consistent. The oxidation of leucine by skeletal muscle is increased threefold in fasted when compared to fed rats, whereas protein synthesis, but not proteolysis, is decreased (7), suggesting that essential amino acids are not spared under these fasting conditions in this species. In 7-d fasted obese subjects, rates of leucine oxidation have also been reported to be increased when compared to the immediate postabsorptive period (13). In contrast, the results of leucine flux and oxidation studies in fasted dogs (12) and the results of leucine and lysine kinetic data and [15]glycine studies in fasted or partially fasted obese subjects, suggest a decreased whole body protein turnover (13–15).

During fasting, plasma glucose and insulin concentrations decrease, whereas those of free fatty acids (FFA),1 ketone bodies, and branched chain amino acids increase. Acute insulin withdrawal in somatostatin- and glucagon-infused dogs (16) results in an increase in leucine concentrations, and either in an increase or in no change in rate of appearance. Infusion of Na D,L-β-hydroxybutyrate in man increased leucine nitrogen flux but did not affect the rate of leucine carbon flux (11). These data collectively suggest that neither a decrease in plasma insulin nor an increase in plasma ketone body concentrations per se is responsible for the decrease in leucine flux observed with fasting. It is possible that increased availability of FFA alone could account for the decrease in leucine flux and possibly oxidation observed with fasting (12, 13). Therefore, the present studies were designed to determine the effects of an acute increase or decrease of plasma FFA on leucine flux and oxidation in dogs.

Methods

Isotopes. L-[4,5-3H]leucine (100 mCi/mmol) and L-[1-14C]leucine (55 mCi/mmol) were purchased from Amersham Corp., Amersham, United Kingdom. [1-14C]α-ketoisocaproate (KIC) was prepared by the method of Rudiger et al. (17) with final purification by high performance liquid chromatography (HPLC) (18). Na[14C]HCO3 (66 mCi/mmol) was obtained from New England Nuclear, Boston, MA.

Analytical methods. Plasma KIC concentrations and specific activities were determined by HPLC (18). Plasma leucine concentration and specific activity (SA) were determined using either a combination of HPLC and ion exchange chromatography (12, 19), or our recently developed HPLC method (18). Leucine containing aliquots derived from plasma

1. Abbreviations used in this paper: FFA, free fatty acids; HPLC, high performance liquid chromatography; KIC, α-ketoisocaproate; SA, specific radioactivity; TG+H, triglyceride plus heparin.
samples were collected in scintillation vials, scintillation cocktail added, and radioactivity determined by scintillation spectrometry. Radioactivity was corrected for quench and 14C spillover into the tritium channel using an external standard method. No differences were observed in the leucine SA among the dogs analyzed by the two methods.

To determine the rates of 14CO2 expired, a funnel made from a plastic bottle and fitted with a rubber diaphragm was placed over the dog's muzzle and the expired air was directed either to the atmosphere or to a collection bag using a three-way valve. Expired air was collected in Douglas bags (Warren E. Collins, Inc., Braintree, MA) over a 2-min period, and the collected gas subsequently aspirated slowly through a 1-M ethanolamine solution to trap CO2. A 1.0-M aliquot of this solution was pipetted in triplicate into scintillation vials, scintillation cocktail added, and the 14C radioactivity determined.

Plasma concentration of FFA (20), β-hydroxybutyrate, acetoacetate (2), lactate, and pyruvate (21) were determined as previously described. Plasma glucose concentrations were determined using a glucose oxidase method (Beckman II glucose analyzer; Beckman Instruments, Inc., Palo Alto, CA). The plasma concentration of insulin and glucagon was determined by radioimmunoassay (22, 23).

Experimental design. Healthy mongrel dogs weighing between 15 and 20 kg were used. Animals were housed in the Mayo Foundation animal care facility and fed a standard dry dog food (Dog Chow; Ralston Purina Co., Chow Div., St. Louis, MO). Dogs were fasted for either 14 or 46 h before the study as dictated by the protocols below, and allowed free access to water. The last meal before the study was offered under direct observation and entirely consumed within 5 min.

On the morning of the study the dog was placed in a standing sling and two cephalic veins were cannulated with 18-gauge catheters. One vein was used for isotopic infusions and the other for either triglyceride plus heparin (TG+H), nicotinic acid, or glyceral infusions. A catheter was also advanced through a saphenous vein to the proximity of the right atrium to obtain mixed central blood samples.

At 150 min, a primed-constant infusion of 4.5H leucine (1.0–1.8 μCi/kg, 0.02–0.03 μCi/kg·min) and [1-14C]KIC (0.05 μCi/kg, 0.01 μCi/kg·min) was initiated. The bicarbonate pool was primed with a 4–5 μCi bolus of [1-14C]HCO3 to facilitate equilibration of 14CO2 in expired air (24). Baseline samples for plasma substrate and SA determination and for expired 14CO2 determination were collected at −30, −20, −10, and 0 min. Thereafter, animals were studied according to four different protocols. Four groups of dogs were studied.

Group I: TG+H. To acutely increase plasma FFA concentrations, 14-h fasted dogs were infused from 0 to 180 min with a 10% triglyceride-glycerol emulsion (Intralipid; Cutter Laboratories, Inc., Berkeley, CA) at 0.4 ml/min. An intravenous bolus (20 U/kg) of Na heparin was administered at 0 min and subsequently at 30-min intervals throughout the study.

Group II: glycerol. In order to distinguish between the effects of increased FFA and glycerol release upon hydrolysis of the triglyceride emulsion from those induced by the free glycerol present in the commercial lipid emulsion, four additional dogs were studied under identical conditions, except that they were infused with glycerol alone at a mean rate of 11 mg·kg·min, which is comparable to the amount of free glycerol infused in group I.

Group III: nicotinic acid. To determine the effects of an acute decrease in the plasma FFA concentration, six 46-h fasted dogs were infused for 150 min with a primed dose (1 mg/kg), continuous (0.2 mg·kg·min) infusion of nicotinic acid, a rate approximately twice that employed in man (25).

Group IV: nicotinic acid plus TG+H. In an attempt to control for potential direct effects of nicotinic acid, four additional animals were studied as described in group III above, except that triglyceride and heparin were infused as described in group I.

Calculations. Since plasma concentration of leucine and KIC and SA of [1-14C] and [4-14C] leucine and [1-3H] and [4-3H] KIC were nearly constant during the periods of study, estimates of flux and oxidation were calculated assuming steady state conditions. The apparent rates of leucine flux were calculated using the following standard formula: 

\[ \text{flux rate} = \frac{\text{radioactivity injected}}{\text{SA} \times \text{time}} \]

where \( \text{flux rate} \) is the rate of isotopic infusion (dpm/kilograms·minutes) and \( \text{SA} \) is the SA of [3H]leucine in plasma. Similar calculations were carried out using the rate of [1-14C]KIC and the plasma [1-14C]KIC SA to determine KIC flux. Total leucine carbon flux was calculated by adding the leucine and KIC flux. The apparent rates of leucine carbon oxidation were calculated using standard product-precursor relationships: apparent leucine carbon oxidation = (expired 14CO2)/[SA(14C) × 0.8] where expired 14CO2 is the rate (dpm/kilograms·minutes) of 14CO2 expired and SA(14C)KIC is the plasma SA of [1-14C]KIC. The constant 0.8 represents the fractional recovery of 14CO2 in expired breath during 14CO2-bicarbonate infusion and corrects for CO2 fixation and other losses. In addition, leucine carbon oxidation was calculated using the [1-3H]leucine SA (25), analogous to the use of [1-14C]KIC enrichment during infusion of [1-14C]leucine (27).

Leucine flux not accounted for by oxidation, an estimate of the rate of leucine entering protein, was calculated by subtracting the rate of leucine oxidation using the [1-14C]KIC SA or the rate of leucine oxidation using the [1-3H]leucine SA from the total (leucine plus KIC) leucine carbon flux.

Results

Effects of TG+H or glycerol infusions in 16-h fasted dogs. During TG+H infusion, plasma FFA increased (P < 0.01) from 0.5 ± 0.1 to 1.2 ± 0.2 mM, whereas during glycerol infusion, FFA concentrations did not change (0.6 ± 0.1 to 0.7 ± 0.1 mM, Table I). Total ketone body concentrations, (β-hydroxybutyrate plus acetoacetate) were similar in both groups of dogs studied and did not change during either TG+H or glycerol infusions (Table I). The TG+H infusion increased significantly with glycerol infusion, whereas TG+H infusion, but neither change was significant. No changes were observed in plasma glucose, lactate, pyruvate, insulin, or glucagon values with the infusion of either TG+H or glycerol (Table I), although some small differences were observed in baseline values between the two groups of animals studied.

Plasma leucine concentrations increased slightly with the glycerol infusion and were decreased slightly with TG+H infusion, but these changes were not significant (Table II). The plasma concentration of KIC decreased (P < 0.05) with glycerol infusion (22 ± 3 to 16 ± 1 μM) but was not affected by TG+H infusion (Table II). The [3H]- and [14C]-SA of leucine and KIC remained constant during the glycerol infusion. In contrast, the [1-14C]- and [3H]leucine SA increased during TG+H infusion, but those of KIC remained constant. During glycerol infusion the rate of 14CO2 expired increased (9.1 ± 0.9 to 11.6 ± 3.0 × 10−3 dpm/kg·min), primarily due to an increase in 14CO2 expired in one dog; overall, no significant change in the rate of 14CO2 expired was observed. During TG+H infusion, the rate of 14CO2 expired increased (P < 0.02) from 7.3 ± 0.8 to 5.6 ± 0.5 × 10−3 dpm/kg·min (Table II).

During glycerol infusion, total leucine carbon entry (6.5 ± 0.41 to 6.49 ± 0.37 μmol/kg·min) did not change. In contrast, the rate of total leucine carbon decreased from 7.04 ± 0.83 to 6.18 ± 0.64 μmol/kg·min (P < 0.02) during the TG+H infusion as the result of a decrease in both leucine and KIC flux (Table III).

The rate of leucine oxidation increased during glycerol infusion, but this increase was not significant, regardless of the precursor pool SA used. In contrast, TG+H infusion resulted in a decrease in leucine oxidation, using either the [3H]leucine SA (2.08 ± 0.35 to 1.25 ± 0.16 μmol/kg·min, P < 0.05), or the [1-14C]leucine SA (0.88 ± 0.19 to 0.61 ± 0.09 μmol/kg·min, P < 0.02) (Table III).
During glycerol infusion, the rate of leucine disappearance which could not be accounted for by oxidation decreased but the change was not significant (Table III). During TG+H infusion, the rate of leucine disappearance not accounted for by oxidation decreased ($P < 0.05$) using the $[^{14}C]$KIC SA for the calculation of leucine oxidation, but did not change when the rate of leucine oxidation was calculated from the $[^{14}C]$leucine SA (Table III).

### Table I. Effects of Glycerol, Triglycerides, and Nicotinic Acid Infusions on Plasma FFA, Ketone Bodies, Glucose, Lactate, Pyruvate, Insulin, and Glucagon Concentrations in Dogs

<table>
<thead>
<tr>
<th></th>
<th>Glycerol</th>
<th>TG+H</th>
<th>Nicotinic acid</th>
<th>Nicotinic acid + TG+H</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before</td>
<td>After</td>
<td>Before</td>
<td>After</td>
</tr>
<tr>
<td>FFA (mM)</td>
<td>0.6±0.1</td>
<td>0.7±0.1</td>
<td>0.5±0.1</td>
<td>1.2±0.2*</td>
</tr>
<tr>
<td>Total ketone bodies</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>($\beta$OHB + AcAc)</td>
<td>140±33</td>
<td>153±31</td>
<td>89±8</td>
<td>92±10</td>
</tr>
<tr>
<td>$\beta$OHB/AcAc</td>
<td>0.6±0.1</td>
<td>0.5±0.1</td>
<td>1.1±0.2</td>
<td>1.5±0.3</td>
</tr>
<tr>
<td>Glucose (mg/dl)</td>
<td>111±6</td>
<td>114±8</td>
<td>95±4</td>
<td>98±4</td>
</tr>
<tr>
<td>Lactate (mM)</td>
<td>1.25±0.15</td>
<td>1.37±0.13</td>
<td>1.73±0.20</td>
<td>1.47±0.19</td>
</tr>
<tr>
<td>Pyruvate ($\mu$M)</td>
<td>55±19</td>
<td>77±27</td>
<td>42±13</td>
<td>63±15</td>
</tr>
<tr>
<td>Insulin ($\mu$U/ml)</td>
<td>8±2</td>
<td>6±1</td>
<td>12±2</td>
<td>10±1</td>
</tr>
<tr>
<td>Glucagon (pg/ml)</td>
<td>67±7</td>
<td>65±6</td>
<td>173±31</td>
<td>155±21</td>
</tr>
</tbody>
</table>

* $P < 0.01$. † $P < 0.05$. $\beta$OHB + AcAc, $\beta$-hydroxybutyrate plus acetoacetate.

### Table II. Steady State Plasma Leucine and KIC Concentrations, SA, and Rates of $^{14}$CO$_2$ Expired Before and After Infusions of Glycerol, Triglyceride, and Nicotinic Acid in Dogs

<table>
<thead>
<tr>
<th></th>
<th>Glycerol</th>
<th>TG+H</th>
<th>Nicotinic acid</th>
<th>Nicotinic acid + TG+H</th>
</tr>
</thead>
<tbody>
<tr>
<td></td>
<td>Before</td>
<td>After</td>
<td>Before</td>
<td>After</td>
</tr>
<tr>
<td>Isotope infusion rate (dpm $\times 10^{-3}$/kg·min)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>$[^{1}H]$Leucine</td>
<td>59.18±5.09</td>
<td>59.15±6.38</td>
<td>40.20±3.39</td>
<td>66.99±5.44</td>
</tr>
<tr>
<td>$[^{14}C]$-Ketoisocaprate</td>
<td>25.03±5.25</td>
<td>21.45±1.71</td>
<td>25.95±3.04</td>
<td>16.96±0.92</td>
</tr>
<tr>
<td>Leucine</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concentrations ($\mu$M)</td>
<td>130±9</td>
<td>134±11</td>
<td>137±8</td>
<td>123±9</td>
</tr>
<tr>
<td>$[^{1}H]$-SA ($dpm/nmol$)</td>
<td>12.1±1.2</td>
<td>12.3±0.8</td>
<td>13.4±1.2</td>
<td>15±1.1*</td>
</tr>
<tr>
<td>$[^{14}C]$-SA ($dpm/nmol$)</td>
<td>4.3±0.8</td>
<td>3.6±0.8</td>
<td>3.9±0.5</td>
<td>4.8±0.6*</td>
</tr>
<tr>
<td>$\alpha$-Ketoisocaprate</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Concentrations ($\mu$M)</td>
<td>22±3</td>
<td>16±1†</td>
<td>13±1</td>
<td>14±1</td>
</tr>
<tr>
<td>$[^{1}H]$-SA ($dpm/nmol$)</td>
<td>16.4±4.0</td>
<td>15.3±3.4</td>
<td>9.6±1.3</td>
<td>9.9±0.9</td>
</tr>
<tr>
<td>$[^{14}C]$-SA ($dpm/nmol$)</td>
<td>10.1±1.4</td>
<td>9.3±1.1</td>
<td>8.0±1.4</td>
<td>9.0±1.2</td>
</tr>
<tr>
<td>Expired $^{14}$CO$_2$ (dpm $\times 10^{-3}$/kg·min)</td>
<td>9.1±0.9</td>
<td>11.6±3.0</td>
<td>7.3±0.8</td>
<td>5.6±0.5§</td>
</tr>
</tbody>
</table>

Data are expressed as mean±SE, see Methods for details of the experimental design. * $P < 0.01$ cf before value. † $P < 0.05$ cf before value. § $P < 0.02$ cf before value.
TG+H. During both infusion studies, the $^3$H- and $^4$C-SA of leucine decreased ($P < 0.01$ with nicotinic acid; $P < 0.05$ with nicotinic acid plus TG+H, Table II); although the $[^3]$H- and $[^4]$CJIC SA decreased under both study conditions, these changes were not significant. The rate of $[^4]$CO$_2$ expired increased ($P < 0.01$) during both infusion periods (Table II), but this increase was blunted during the combined infusion of nicotinic acid plus TG+H.

Injection of nicotinic acid alone resulted in an increase ($P < 0.05$) in the rate of leucine entering the vascular space (3.20±0.19 to 3.75±0.37 µmol/kg·min, Table III). The rate of appearance of KIC increased, but the change was not significant (Table III). Although the entry of leucine and KIC increased, the changes only approached the level of significance ($0.05 > P > 0.10$).

During the combined infusion of nicotinic acid plus TG+H, leucine, KIC, and leucine plus KIC entry increased, but the changes were not significant (Table III). During the nicotinic acid infusion, leucine oxidation increased ($P < 0.01$) two- to threefold regardless of whether $[^4]$CJIC or $[^4]$Cleucine SA was used (Table III). During combined infusion of nicotinic acid plus TG+H, leucine carbon oxidation increased ($P < 0.05$ or <0.01 depending on the precursor used) twofold regardless of the model used (Table III).

The rate of nonoxidized leucine disappearance during nicotinic acid infusion alone decreased regardless of the precursor pool SA used for the calculation of leucine oxidation, but the decrease was only significant ($P < 0.05$) using the $[^4]$Cleucine SA (4.55±0.28 to 3.32±0.51 µmol/kg·min, (Table III). During infusion of nicotinic acid plus TG+H, rate of disappearance of leucine not accounted for by oxidation did not change significantly regardless of which precursor pool SA was used for calculating leucine oxidation (Table III).

In a limited number of experiments ($n = 3$) using a constant infusion of only H$^{14}$CO$_2$, a 5–20% increase in the rate of expired H$^{14}$CO$_2$ was observed following initiation of the nicotinic acid infusion. If this occurred during the studies of leucine kinetics employing nicotinic acid, this small increase in H$^{14}$CO$_2$ expired (or decrease in CO$_2$ fixation) due to nicotinic acid's effect on expired H$^{14}$CO$_2$ would result in a slight overestimation in the rate of leucine carbon oxidation; however, since leucine oxidation increased 250–300%, any error in the estimation of expired H$^{14}$CO$_2$ on this basis would be negligible.

**Correlations.** Inverse correlations were found between the changes in plasma FFA concentrations and estimates of rates of leucine carbon oxidation (using $[^4]$CJIC SA, $r = -0.806$, $P < 0.001$, or using $[^4]$Cleucine SA, $r = -0.762$, $P < 0.001$, Fig. 1). Changes in plasma FFA concentrations were inversely correlated with the rate of appearance of leucine ($r = -0.713$, $P < 0.001$) and with total leucine carbon (leucine + KIC) ($r = -0.612$, $P < 0.003$). Changes in FFA did not correlate with the rate of appearance of KIC or with the estimates of nonoxidized leucine disappearance using $[^4]$CJIC as the precursor pool for oxidation. A weak ($r = 0.451$) but significant ($P < 0.04$) inverse correlation was observed between $\Delta$FFA and nonoxidized leucine disappearance using the $[^4]$CJIC SA.

Changes in total ketone body concentrations correlated inversely with the rate of leucine oxidation using $[^4]$CJIC ($r = -0.473$, $P < 0.004$) or using $[^4]$Cleucine, as well as with both the rate of appearance of leucine ($r = -0.576$, $P < 0.05$) and of total leucine carbon ($r = -0.435$, $P < 0.05$). Again, changes in total ketone body concentrations did not correlate with the estimated rates of appearance of KIC or nonoxidized leucine disappearance. Changes in β-hydroxybutyrate/acetacetate ratio were inversely correlated with changes in leucine apparent oxidation ($r = -0.470$, $P < 0.05$).

However, when multiple regression analysis was performed between changes in FFA and total ketone bodies as independent variables and each of the above mentioned parameters of leucine metabolism as dependent variable(s) only changes in FFA con-
A Leucine Carbon Ra

+1.0

0

-1.0

B Leucine Rd via oxidation

+2.0

-2.0

C Non-oxidized Leucine Rd

+2.0

-2.0

△ Plasma FFA mM

△ Plasma FFA mM

Figure 1. The effects of changes in plasma FFA concentration on changes in leucine rate of appearance (Ra) and rate of disappearance (Rd) attributable to oxidative and nonoxidative processes. Changes in FFA concentrations were induced by infusions of saline, TG+H, nicotinic acid, or a combination of nicotinic acid and TG+H. (A) — o —, KIC Ra (r = -0.299, P = NS); — - - - , leucine Ra (r = -0.713, P < 0.001); — - - - , leucine plus KIC Ra (r = -0.612, P < 0.003). (B) — - - - , using [14C]KIC (r = -0.806, P < 0.001); — - - - , using [14C]leucine (r = -0.762, P < 0.001). (C) — - - - , using [14C]KIC for oxidation (P = NS); — o — , using [14C]leucine for oxidation (r = 0.451, P < 0.04).

Discussion

Fasting is associated with complex changes in the circulating concentrations of hormones and substrates, rates of substrate flux and interconversion, and energy expenditure. Therefore, it is difficult to determine the role and/or effects of alterations in availability of a single substrate, such as fatty acids, from fasting studies alone. The present studies demonstrate that acute changes in plasma FFA concentrations within the physiologic range observed with feeding and fasting via the infusion of triglyceride, heparin, and/or nicotinic acid are inversely related to estimates of whole body leucine carbon flux and oxidation, but not with estimates of leucine disappearance unaccounted for by oxidation (a possible indicator of the rate of leucine entering protein). Assuming that the only source of plasma leucine in the postabsorptive state is endogenous protein, and that there is no accumulation of intracellular free leucine, we conclude that FFA availability is inversely related to the rate of whole body proteolysis and of the oxidation of at least one essential amino acid, leucine. As a result, at least some of the changes in leucine metabolism observed in both the fasted (12, 13) and fed (28-30) states may be linked to changes in FFA availability.

During periods of fasting which are associated with increases in plasma FFA concentrations, decreased rates of whole body leucine and lysine flux have been observed in both man (13, 14) and dog (12, 28). These observations are consistent with a decrease in protein turnover and/or a fasting-induced protein sparing effect (15). A decrease in whole body leucine oxidation has been observed in some (12, 28), but not all (13) reports and is consistent with a decrease in urinary nitrogen excretion observed with fasting in man (2, 15). Adibi and co-workers (13) reported decreased leucine flux, but increased leucine oxidation in obese humans consuming 80 kcal/kg for 7 d; this would result in a greater fraction of the leucine flux being oxidized. Their results would not be consistent with the predominance of published data outlined above, demonstrating a nitrogen and/or an amino acid sparing effect of fasting.

Nicotinic acid, a potent inhibitor of lipolysis, has been reported to increase urea excretion and amino acid oxidation in fasted animals (31–34). These results are thought to be related to the effect of nicotinic acid on the availability of lipid-derived fuels, resulting in increased utilization of other substrates (e.g., amino acids and glucose) as metabolic fuels, and not a direct effect of the drug (32, 33). However, nicotinic acid, by direct or indirect effect, decreases the hepatic reduced nicotinamide-adenine dinucleotide/nicotinamide-adenine dinucleotide (NADH/NAD) ratio (35) and in the present study decreased the plasma \( \beta \)-hydroxybutyrate/acetacetate ratio (Table I). The activity of the branched chain \( \alpha \)-ketoadipic dehydrogenase complex (36, 37) and rates of proteolysis (38) have been linked to alterations in the NADH/NAD ratio; therefore, changes in cellular redox could account for the increased rates of leucine flux and oxidation observed in the present studies during nicotinic acid infusion, as well as during the infusion of triglyceride and heparin (39). Although the changes in the \( \beta \)-hydroxybutyrate/acetacetate ratio in the present studies are consistent with the purported effect of changes in NADH/NAD on leucine flux and oxidation, changes in the lactate/pyruvate ratio (an indicator of the cytosolic redox state) were not. In addition, neither the plasma concentrations of these substrates (\( \beta \)-hydroxybutyrate, acetacetate, lactate, pyruvate) nor their ratios (\( \beta \)-hydroxybutyrate/acetacetate or lactate/pyruvate) correlated by multiple regression analysis with changes in leucine flux or oxidation. However, since the plasma \( \beta \)-hydroxybutyrate/acetacetate and lactate/pyruvate ratios are only indirect indicators of cellular redox state, we cannot exclude the possibility that changes in cellular redox (whether a direct effect of either FFA and/or nicotinic acid) do not underlie, at least in part, the changes in leucine (and presumably protein) metabolism observed in the present studies. It should be pointed out that the addition of TG+H infusion to nicotinic acid infusion had no substantial effect on leucine carbon flux estimates when compared to nicotinic acid infusion alone (\( \Delta 0.68 \) vs. 0.77 \( \mu mol/kg \cdot min \), respectively, Table III), whereas the nicotinic acid–induced increase in estimates of leucine oxidation were decreased by \( \sim 50 \% \) with the addition of TG+H infusion.
As a result, we cannot establish with certainty that the changes observed in leucine metabolism during nicotinic acid infusion were directly attributable to changes in fatty acid availability as estimated by changes in the plasma concentration of FFA.

Regardless of the experimental perturbation, the changes in total leucine carbon flux (leucine plus KIC) are almost entirely attributable to changes in leucine flux alone since the KIC flux did not change significantly. However, in all cases the KIC flux changed in the same direction as the leucine flux. The reasons for this apparent discrepancy in the magnitude of change in the rate of appearance of leucine and KIC into the plasma space is not clear. The isotope model utilized assumes equal efficiency of removal of the leucine and KIC labels from the plasma space. Under the experimental conditions employed, changes in plasma substrate concentration might differentially affect the transport of the labels to the intracellular space; therefore the resultant changes in leucine and KIC SA and the flux estimates derived from these SA may not be proportional. However, the differential changes in leucine and KIC flux may suggest some fundamental differences in the metabolism of leucine and its α-ketoacid which remain to be elucidated.

The present studies demonstrate the potential importance of fatty acid availability in protein sparing, and are consistent with an increase in protein wasting (increased total leucine carbon flux and oxidation) under situations in which fatty acid fuel availability is limited. Since the present studies were carried out over short periods of time and nitrogen balance was not assessed, no conclusion can be drawn regarding the availability of FFA over either short or long term periods of time on whole body protein metabolism. Whether increasing plasma FFA via the infusion of triglycerides has advantages when simultaneously infused with other substrates such as glucose and amino acids, as occurs during parenteral nutrition, remains to be clearly established.

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P. Tessari, S. L. Nissen, J. M. Miles, M. W. Haymond

580


