Adaptation of the Growth Hormone and Insulin-like Growth Factor-I Axis to Chronic and Severe Calorie or Protein Malnutrition

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Abstract

The hierarchy of diet components (e.g., protein, carbohydrate, vitamins, and minerals) influencing growth hormone (GH), insulin-like growth factor-I (IGF-I), and their binding proteins (BP) is not well defined. Young adult rats were fed diets for 1 mo that included low protein or 60% and 40% of carbohydrate calories. We hypothesized that levels of both hormones, their dominant BPs and liver IGF-I mRNA would fall, and that part of the mechanism for decreasing serum IGF-I would be enhanced IGFBP-3 protease activity. By day 30, caloric deprivation to 40% lowered serum GH, GHBP, IGF-I and IGFBP-3, and liver IGF-I mRNA. This was the only condition resulting in body weight loss (−15%) vs 39% gain in controls. Restriction to 60% calories had no impact on BP levels, slightly lowered IGF-I (−12%) in the face of a 95% inhibition of GH levels, while allowing a modest 9% body weight gain. Protein deprivation lowered serum GH, IGF-I and IGFBP-3, and liver IGF-I mRNA, while GHBP levels were normal. The reduced total IGF-I under these dietary conditions could not be explained by an increase in IGFBP-3 protease activity, or a decrease in the association of IGF-I with IGFBP-3 and the acid labile subunit. (J. Clin. Invest. 1995; 95:2258-2265.) Key words: binding proteins • growth hormone • insulin-like growth factor-I • malnutrition

Introduction

Nutritional status is a comorbidity variable in a multitude of diseases (e.g., AIDS, cancer) and time to death is predicted by the magnitude of body cell mass depletion (1, 2). The anabolic growth hormone (GH)/insulin-like growth factor I (IGF-I) axis, including a host of circulating binding proteins (BP), is a complex endocrine system that promotes statural growth during development and directs substrate availability toward lean body mass vs fat throughout the lifespan. Pituitary-derived GH stimulates IGF-I synthesis and export from the liver, the primary source of circulating IGF-I; this coupling of GH to IGF-I production appears optimal only with adequate dietary intake and general health (3). Insulin-like growth factor-I, in turn, targets a wide variety of cells, where it stimulates multiple cellular responses, including glucose and amino acid uptake as well as synthesis of DNA, RNA, and cellular protein (reviewed in reference 4).

Given this understanding of the system biology, the GH/IGF-I axis must sense and adjust to reductions in essential nutrient availability. Indeed, a generation ago Grant et al. (5), and Hintz et al. (6) first observed that serum IGF bioactivity was decreased in protein-malnourished children. Soliman et al. (7) later reported low serum IGF-I bioactivity and high serum GH levels in children with combined protein, energy, and micronutrient deficits; these hormones returned to normal after nutritional rehabilitation (7).

Since these studies, an effort to understand the metabolic and nutrient controls of the GH/IGF-I axis has evolved (3). In animal models, a spectrum of micronutrient (e.g., potassium [8], magnesium [9], zinc [9, 10]) and macronutrient (carbohydrate [11, 12], protein [6, 7, 13-22], fat [23]) deficiencies have produced variable GH responses, with no change or decreased liver GH-binding and consistently diminished serum IGF-I and IGFBP-3 levels. While these studies describe nutritional influences on the axis, most were limited to pubertal animals studied for a few days and did not control for equivalent micronutrient intake when deficient macronutrient diets were prepared. One report demonstrates that protein deficiency in the weaning rat reduces serum IGF-I significantly more than in the adult rat (24). This suggests a reduction in IGF-I sensitivity to protein deficiency with age.

Under normal conditions both GH and IGF-I are associated with specific BPs in the circulation; the plurality of GH is bound to a high affinity GHBP, and the majority of IGF-I is bound in a 150-kD complex consisting of IGF-I, IGFBP-3, and an acid-labile subunit (84-86 kD). The remaining IGF-I is either free (<5%) or bound to five other IGFBPs in binary complexes. Free GH and IGF-I are considered to be the active forms of the hormones. The effects of restricted protein and energy intake on GHBP are unknown. In contrast, protein restriction was reported to decrease serum IGFBP-3 in pubertal, 4-wk-old rats (19). This was speculated to occur due to diminished IGFBP-3 protein production or stability, perhaps via IGFBP-3 proteolysis, an activity correlated with severe catabolic conditions and late pregnancy (25-28). The influence of malnutrition alone (in the absence of trauma) on this protease or the state of the 150-kD heterotrimer is unknown.

The purpose of these experiments was to define in young adult rats the response of the GH/IGF-I axis during the progressive metabolic adaptation to discrete and chronic protein or energy deficiency. The primary postulate was that these dietary restrictions would dramatically suppress serum IGF-I via reduc-
tion of both liver IGF-I mRNA and IGFBP-3 availability. We attempted to discover: (a) if loss of GHBP explains the reported reductions in circulating GH levels during food restriction (12, 23, 29); (b) the threshold and time at which limited calories or protein, without micronutrient, protein (low calorie diets) and fat undernutrition, reduce total serum IGF-I; (c) whether the mechanism for this IGF-I response involves reductions in IGFBPs. IGFBP-3-specific protease activity, the 150-kD complex, or liver IGF-I mRNA.

Methods

Design

Pilot study. A pilot study was performed to establish the level of caloric restriction needed to reduce serum IGF-I levels in young adult rats. Male Sprague-Dawley (Harlan Sprague-Dawley, Indianapolis, IN) rats were housed individually in hanging wire-mesh cages. Rats were fed isocaloric, defined diets (3.90 calories/g) for 21 d as follows: ad lib. control (AIN-76A; control; n = 6), or restricted to 75% (n = 7) and 60% (n = 6) the calories consumed by control. The results of this study led to the 60% and 40% calorie diets in the study reported below.

Animals and groups. 8-wk-old male Sprague-Dawley rats were housed individually in hanging wire-mesh cages in a temperature (22°C±2) and light (12:12 h light-dark cycle; light off at 9:00 a.m.) controlled room. After adaptation to conditions, handling, and a defined diet (AIN-76A; Dyets, Inc.; Bethlehem, PA), rats were randomly placed into one of six weight-matched groups (282 g±2; mean±SE). Rats were fed isocaloric, defined diets (3.90 calories/g) for 30 d as follows: ad lib. control (AIN-76A; control; n = 7), ad lib. 4% casein (low protein diet [LoPro]; n = 8), or restricted to 60% (n = 7) and 40% (n = 8) the calories consumed by control. The 60% and 40% diets were formulated such that protein, fat, vitamin, and mineral consumption (grams) was equivalent to control intake to avoid protein and micronutrient undernutrition. Body weights and 24-h food intakes, corrected for spillage, were measured daily between 8:00 and 9:00 a.m. The amount of food fed to the 60% and 40% groups was calculated as that percentage of the control group’s previous day food intake (grams). All food cups were returned to the cages immediately before lights off. On days 0, 7, and 21, 0.3 ml of blood was collected from the tail vein of nonfasted rats and serum was frozen at −70°C. On day 30, rats were killed by CO2 and cardiac puncture. Serum samples were collected and frozen at −70°C until assayed for IGF-I, IGFBPs, IGFBP-3 protease, and GH. The pulsatile pattern of GH secretion can also be measured to assess GH pulse height or frequency. However, the necessary cannulation of the maldistributed animals and intense serial blood sampling would have added significant surgical and handling stress. Accordingly, as GH status was not the most important variable and given the added stress, we chose to measure serum GH concentrations from samples taken at a single time-point.

Liver samples were frozen at −70°C or fixed in 10% neutral buffered formalin. As an indicator of carcass adiposity (30), retroperitoneal fat pad was weighed. Thymus, gastrocnemius, and adrenals were weighed and expressed as the fraction of body weight. This protocol was approved by the Protocol Review Committee and Institutional Animal Care and Use Committee of our AAALAC-approved vivarium.

Analytical Procedures

Hematology, blood chemistry, and liver histology. Hematology (hemoglobin, hematocrit) was performed on a Serono Baker 9000 unit (Serono Baker Diagnostics, Inc., Allentown, PA) with veterinary discriminator settings. 20 serum chemistry parameters were measured on a Hitachi analyzer (Tokyo, Japan). Sections of liver were harvested from each animal, fixed in 10% neutral buffered formalin, processed routinely to paraffin blocks, sectioned at 4 μm, and stained with hematoxylin and eosin by standard procedures.

Serum GH. Serum GH concentrations (ng/ml) were measured utilizing a double antibody ELISA assay. Briefly, ELISA plates were coated with a goat anti-rat GH antibody. After incubation and washing, National Institutes of Health standard rat GH-RP-2 or 100 μl of serum (1:50 dilution) was applied to wells and incubated at room temperature with gentle agitation for 2 h. Sheep anti-rat GH antibody was then applied and plates were incubated at room temperature for 2 h. After washing, Kirkegaard and Perry 2-component tetramethyl benzidine substrate solution was applied. The reaction was stopped with H2PO4, and the optical density read at 450 nm.

Serum GHBP. Rat GHBP was assayed using an RIA method (31) and values are expressed as nanograms rhGHBP per milliliter serum.

Serum IGF-I concentrations. Before measurement of serum total IGF-I, high molecular weight IGFBP’s were precipitated by incubating in acid-ethanol (12.5% 2 N HCl, 87.5% EtOH) at 4°C for 30 min (32). There was incomplete recovery of IGF-I if lesser ratios than 1:15 serum to acid-ethanol were used. Samples were then centrifuged for 5 min at 10,000 g. Supernate was neutralized with 1 M Tris base, and diluted in PBS pH 7.4 with 0.1% gelatin, 0.05% Tween 20 and 0.01% thimerosal. Serum IGF-I concentrations were determined that same day by radioimmunoassay (33). The percentage recovery of recombinant human IGF-I (Genentech, Inc.) added to samples extracted as above was 91.±9.9% (mean±SE; n = 4).

SDS-PAGE and Western ligand blotting. Serum IGFBP profiles were measured on 7-, 21-, and 30-d samples. Serum (2.5 μl) was diluted with nonreducing SDS sample buffer (0.5 M Tris, pH 6.8; 69% glycerol; 4% SDS), applied to a 4% stacking gel and electrophoresed through 11% polycrylamide gel. Size-fractionated proteins were then transferred to nitrocellulose according to Towbin et al. (34). Western ligand blotting was performed following the method of Hossennopp, et al. (35) as previously described (25), using −10×106 cpm 125I-IGF-I and 125I-IGF-II. Densitometry was performed using the BAS2000 phosphorimager (Fuji, Inc., Tokyo, Japan). Relative quantities of IGFBP-3 were determined after scanning the bands corresponding to 40–45 kDa. At least three separate control sera were run on each gel and IGFBP-3 values are reported as percentage of control±SE.

IGFBP-3 molecular distribution. It is possible that nutritional state will change the 150-kD trimer complex that carries the majority of IGF-I in the circulation. For example, more dimer or free IGF-I could result in faster clearance of IGF-I and thereby be a mechanism to lower total serum IGF-I levels. To determine the molecular weight distribution of endogenous IGFBP-3, pooled serum samples (200 μl) were size fractionated at neutral pH on a Super-Dex 200 column (Pharmacia LKB Biotechnology; Uppsala, Sweden) at a flow rate of 0.5 ml/min and 0.5-mL fractions were collected. Fractions 7–20 (~300–20 kD) were analyzed by Western ligand blotting as described above.

Serum IGFBP-3-specific protease activity. Recombinant 125I-IGFBP-3 (30,000 cpm in 5 μl 20 mM Hepes+1 mM CaCl2, pH 7.2; Diagnostic Systems Laboratory, Inc., Webster, TX) was incubated for 5 h at 37°C with 20 μl diluted test serum (1:10 vol/vol serum: 20 mM Hepes +1 mM CaCl2, pH 7.2). The reaction was terminated by the addition of nonreducing SDS sample buffer, the radiolabeled fragments separated by SDS-PAGE on a 15-cm gel under nonreducing conditions and the bands visualized via autoradiography. To control for spontaneous degradation of 125I-IGFBP-3 during the incubation period, a sample containing 30,000 cpm 125I-IGFBP-3 in buffer alone was included.

Liver IGF-I mRNA isolation. Liver total RNA was isolated from 200 mg of frozen liver homogenized in 1 ml of 5 M guanidium thiocyanate buffer containing 1% beta-mercaptoethanol. Samples were phenol–chloroform (1:1) extracted and centrifuged at 14,000 g for 10 min. The aqueous phase was removed and precipitated with 8 M LiCl overnight at −20°C. Samples were centrifuged at 14,000 g for 25 min at 4°C. The supernate was removed and the pellet was resuspended in 3 M LiCl, centrifuged at 14,000 g for 10 min at 4°C. The supernate was removed and the pellet resuspended in 1×SET (1% sodium dodecyl sulfate, 10M Tris, pH 8.0, and 5 mM EDTA, pH 8.0), treated with 200 μg/ml proteinase K (Boehringer Mannheim Biochemicals, Indianapolis, IN) for 1 h at 45°C, phenol–chloroform (1:1) extracted.

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and precipitated with 1 vol isopropanol and 1/10 vol 3 M NaOAc at −20°C until assayed.

Liver IGF-I and cyclophilin mRNA, solution hybridization protection assay. Liver IGF-I and cyclophilin mRNA were quantified following procedures described previously (36). A 209-bp segment of the rat IGF-I gene, containing 41 bases of exon 3 and all of exon 4 was subcloned into the phagemid pBluescriptISK (+). A 110-bp segment of the rat cyclophilin gene was subcloned into the phagemid pBluescriptKS (+) (Stratagene, La Jolla, CA). Sense RNA standards and antisense RNA probes (labeled to 10^6 cpm/μg with [32P]UTP) were generated using commercially available transcription reagents and protocols (Stratagene). Samples were reconstituted in 1 × TE (10 mM Tris, pH 8.0 and 1 mM EDTA, pH 8.0). The concentrations of RNA in each sample was determined from a measurement of the absorbance of OD_{260}. 10 μg of each RNA sample (in 5 μl 1 × TE) was assayed. The sense RNA standard was quantified by absorbance at OD_{260} and diluted in 1 × TE for addition into the assay in 5 μl 1 × TE; standards were run in the range from 0.5 to 600 pg/tube. 25 μl of hybridization buffer (36) containing 1 ng of each antisense RNA probe (quantitated by incorporation of 32P and resuspended in hybridization buffer) were added and the reactions were heated to 85°C for 5 min, then incubated overnight at 45°C.

Hybridizations were digested with 0.2 ml RNAase digestion mix (36) for 1 h at 30°C, followed by treatment with sodium dodecyl sulfate and proteinase-K for 15 min at 37°C (36). The protected RNA:RNA hybrids were phenol–chloroform extracted, precipitated with 100% ethanol, and electrophoresed through nondenaturing polyacrylamide gels. The gels were dried and exposed to x-ray film, and the bands formed by the protected RNA:RNA hybrids were visualized on a BAS2000 phosphorimager for quantitation. The concentration of RNA was computed by comparing the density of the samples to a standard curve generated by the linear regression of density vs picogram of sense standard (36).

Statistical analyses. Results are given as means±SE. Data were analyzed by one-way ANOVA and subsequent post-hoc tests were performed using Fisher’s PLSD test. For all analyses, a P value of <0.05 was considered significant.

Results

Pilot study. Compared to the circulating IGF-I measured in ad lib. fed controls (365±18 ng/ml), restriction to 60% of normal intake for 21 d insignificantly decreased serum IGF-I (316±21 ng/ml) and diet restriction to 75% had no effect (408±32 ng/ml). Given this surprising result, the final experiment imposed a more severe and prolonged calorie restriction on these young adult animals.

General health, food consumption, and body weight. No animals died or were judged clinically sick during the course of this protocol. Cumulative food intake in grams was as follows: control, 590±12; LoPro, 561±18; 60%, 357±1; 40%, 241±0.3. Control rats consistently gained weight from day 0 (281±6) to day 30 (391±7). Reduction to 60% of control intake yielded a modest weight gain from day 0 (281±4) to day 30 (306±4). In contrast, a reduction to 40% of control intake resulted in steady body weight loss from day 0 (281±4) until day 20, followed by maintenance of body weight to day 30 (238±2). The rats fed the LoPro diet (day 0: 282±4) initially lost weight and then tracked with the 60% group to day 30 (304±4). In summary, by day 30 the controls had gained 39% body weight, while 60%, 40%, and LoPro body weights were +9, −15 and, +8% of day 0, respectively.

Serum GH, GHBp, and IGF-I concentrations. The serum GH concentrations were markedly reduced in the 40% and 60% groups, whereas the serum GH concentrations of LoPro group was halved but not significantly different from controls at day 30 (Table I). Serum GHBp concentrations of the 40% group were lower, while those of the 60% and LoPro groups were similar to controls (Table I). Table II lists the changes in serum IGF-I concentrations over the 30-d period. These were similar among the groups on day 0, but by day 7, IGF-I declined in the restricted groups, particularly in the LoPro rats. On day 21, IGF-I levels were similarly low in the 40% and LoPro rats. On day 30, serum IGF-I levels were slightly lower in the 60%, and further reduced in the 40% and LoPro rats compared to controls.

Serum IGF binding proteins. Serum IGFBP-3 levels of the 60% rats were unchanged through the month while 40% and LoPro rats had a 20–30% reduction in IGFBP-3 (Fig. 1).

Serum IGFBP-3 molecular distribution. While it was evident that there was decreased total IGFBP-3 in the 40% and LoPro groups, energy and protein deprivation had no effect on the overall molecular distribution of IGFBP-3 in serum (Fig. 2). The concept was that a change in this distribution pattern would suggest another mechanism for lowering total IGF-1 via an alteration in either the binding affinities and/or availability of IGFBP-3 and/or the acid-labile subunit. Regardless of diet, the majority of IGFBP-3 was bound to the acid-labile subunit and IGF-I in the 150-kD trimer, while a smaller proportion of serum IGFBP-3 was found in the 55-kD complex.

Serum IGFBP-3 protease activity. All rats had measurable IGFBP-3 protease activity, as illustrated in Fig. 3. When compared to the amount of IGFBP-3 protease activity present in pregnant rat serum, none of the diet treatments had increased IGFBP-3 protease activity as witnessed by little change in the amounts of low molecular weight IGFBP-3 fragments.

Table I. Serum GH and GHBp concentrations

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>60%</th>
<th>40%</th>
<th>LoPro</th>
</tr>
</thead>
<tbody>
<tr>
<td>GHBp (ng/ml)</td>
<td></td>
<td></td>
<td></td>
<td></td>
</tr>
<tr>
<td>Day 0</td>
<td>21.0±0.7</td>
<td>18.3±1</td>
<td>12.2±1</td>
<td>22.2±0.8</td>
</tr>
<tr>
<td>Day 7</td>
<td>54.3±11.2</td>
<td>25.3±10.7</td>
<td>29.5±8.9</td>
<td>12.2±4.7</td>
</tr>
<tr>
<td>Day 21</td>
<td>66.3±35.2</td>
<td>31.7±9.4</td>
<td>5.5±2.3</td>
<td>12.2±8.3</td>
</tr>
<tr>
<td>Day 30</td>
<td>67.8±31.9*</td>
<td>3.5±1.7</td>
<td>4.8±2.1</td>
<td>32.6±15.2*</td>
</tr>
</tbody>
</table>

The mean±SEM (n = 7–8) is represented for serum collected on given days. Superscripts denote significant differences between treatment groups; means not sharing the same superscripts are different at P < 0.05; means without superscripts are not different.

Table II. Total Serum IGF-I Concentrations (ng/ml)

<table>
<thead>
<tr>
<th>Day</th>
<th>Control</th>
<th>60%</th>
<th>40%</th>
<th>LoPro</th>
</tr>
</thead>
<tbody>
<tr>
<td>0</td>
<td>593±36</td>
<td>610±20</td>
<td>590±14</td>
<td>576±19</td>
</tr>
<tr>
<td>7</td>
<td>575±23*</td>
<td>512±244</td>
<td>443±15</td>
<td>362±15</td>
</tr>
<tr>
<td>21</td>
<td>554±22*</td>
<td>502±14*</td>
<td>396±31*</td>
<td>396±23*</td>
</tr>
<tr>
<td>30</td>
<td>582±34*</td>
<td>513±19*</td>
<td>339±13*</td>
<td>391±15*</td>
</tr>
</tbody>
</table>

The mean±SEM (n = 7–8) is represented for serum collected on given days. Superscripts denote significant differences between treatment groups; means not sharing the same superscripts are different at P < 0.05; means without superscripts are not different.
Liver IGF-I mRNA. Liver IGF-I mRNA expressed as picogram/microgram total RNA was decreased by 25 and 33% in the 40% and LoPro groups, respectively, (control, 5.8±0.5; 60%, 5.2±0.7; 40%, 4.3±0.2*; LoPro, 3.9±0.1*; *P < 0.024). Liver cyclophilin mRNA was used to normalize the IGF-I measurement, as cyclophilin was considered a constitutive cell component. We found that liver cyclophilin mRNA content was similar between control, 60%, and 40% groups, but was increased in the LoPro rats (pg cyclophilin/μg total RNA: control, 2.8±0.1; 60%, 2.5±0.2; 40%, 2.8±0.1; LoPro, 3.7±0.3*; *P < 0.001). The ratio of liver IGF-I mRNA to cyclophilin mRNA was decreased to a greater extent in the LoPro than in the 40% group (Fig. 4).

Serology. Hemoglobin, hematocrit, blood urea nitrogen and serum albumin, and total protein values are listed in Table III. Although there were statistical differences in hemoglobin concentrations and hematocrits, these group means are within normal ranges. The similar albumin and total protein concentrations of the energy deprived groups (60% and 40%) compared to controls, demonstrate that these malnourished groups were adequately protein nourished. Caloric restriction to 40% of normal caused some impairment of liver function as measured by in-

Figure 1. (A) Autoradiograph of Western ligand blot of serum IGFBPs. Rats were fed control diet ad lib., a low protein diet ad lib. (LoPro), or restricted to 60 and 40% the calories consumed by controls for 30 d. (B) Time course of the nutritionally induced alterations in serum IGFBP-3 concentrations is shown in the graph. All IGFBP-3 concentrations are expressed as a percent of control values. Data are presented as means±SE (n = 6–8).

Figure 2. Autoradiograph of IGFBP-3 molecular distribution. Rats were fed control diet ad lib., a low protein diet ad lib. (LoPro), or restricted to 40% the calories consumed by controls for 30 d.

Figure 3. Autoradiograph of IGFBP-3 protease activity in rats fed control diet ad lib., a low protein ad lib. (LoPro), or restricted to 60 and 40% control calories for 30 d. (Pregnant rat serum) Serum from pregnant female Sprague-Dawley rat. Each lane represents a different animal.

Figure 4. Quantitation of liver IGF-I mRNA expressed as picogram/picogram cyclophilin (CYC) mRNA is shown for rats fed control diet ad lib., a low protein diet ad lib. (LoPro), or restricted to 60 and 40% control calories. Means±SE. *, †, P < 0.05 vs other groups.
Table III. Selected Serum Chemistry and Hematology Values on Day 30

<table>
<thead>
<tr>
<th></th>
<th>Control</th>
<th>60%</th>
<th>40%</th>
<th>LoPro</th>
</tr>
</thead>
<tbody>
<tr>
<td>Hgb (g/dl)</td>
<td>14.5±0.2*</td>
<td>15.1±0.1†</td>
<td>16.1±0.1‡</td>
<td>13.9±0.2†</td>
</tr>
<tr>
<td>Hematocrit (%)</td>
<td>39±1.0*</td>
<td>40±1.0*</td>
<td>43±0.4†</td>
<td>39±1.0*</td>
</tr>
<tr>
<td>Albumin (g/dl)</td>
<td>3.2±0.04*</td>
<td>3.3±0.04*</td>
<td>3.3±0.02*</td>
<td>3.0±0.1†</td>
</tr>
<tr>
<td>Total Protein (g/dl)</td>
<td>6.8±0.1*</td>
<td>6.7±0.1*</td>
<td>6.8±0.1*</td>
<td>6.1±0.2†</td>
</tr>
<tr>
<td>BUN (mg/dl)</td>
<td>14±1*</td>
<td>15±0.4*</td>
<td>19±2†</td>
<td>5±1†</td>
</tr>
</tbody>
</table>

The mean±SEM (n=7–8) is represented for serum collected on day 30. Superscripts denote significant differences between groups; means not sharing the same superscripts are different at P < 0.05. BUN, blood urea nitrogen.

creased serum glutamic pyruvic transaminase levels (40%, 42±2 IU/liter vs control, 28.6±2 IU/liter) and direct bilirubin concentration. Consumption of the LoPro diet caused liver function to decline as serum albumin, total protein, globulin, and blood urea nitrogen concentrations decreased and alkaline phosphatase activity, total and direct bilirubin concentrations increased. All animals were euclorhyemic at time of death. The following serum chemistries were similar across all groups: serum glutamic oxaloacetic transaminase, creatine phosphokinase, creatinine, bicarbonate, calcium, sodium, chloride, potassium, and indirect bilirubin. Phosphorus concentrations were low in all diet treatment groups compared to controls.

Liver histology. To describe the effects of energy and protein deficiency on the liver, histology sections were evaluated (Fig. 5). In the 40% group, there was no marked difference in morphology compared to controls with the exception of reduced intracellular mass. The LoPro-fed rats had macrovesicular lesions throughout the liver with a greater severity close to the hepatic triad, consistent with the development of a nutritionally induced steatotic liver. In a subsequent study, using Oil Red O staining for fat and Periodic acid Schiff reaction with and without diastase digestion for glycogen content, we confirmed that this pathology was associated with triglyceride accumulation (data not shown).

Organ weights. As listed in Table IV, calorie restriction resulted in a graded increase in fractional adrenal weight, whereas there was a comparable but reciprocal thymic involution. The low protein diet yielded responses in these tissues similar to the 60% calorie condition. The retroperitoneal fat pad contribution to body weight was substantially reduced by limited calories, but unaltered by the low protein diet. The relative gastrocnemius weight was preserved, irrespective of treatment.

Discussion

While a broad spectrum of macro- and micronutrient deficiencies are capable of suppressing serum IGF-I concentrations, the IGF-I response may vary with age, type or degree of inadequacy, and length of malnutrition. Although serum IGF-I levels were low in protein energy-malnourished children and adults (5, 6, 7, 37), there appeared to be a critical threshold before IGF-I concentrations declined (3). The level of energy intake which supports normal circulating IGF-I was indicated in a study by Kupfer, et al. (38). Humans 22–47 yr of age were restricted to ~ 7/3 ad lib. intake (i.e., 20 calories per kilogram ideal body weight with 1 g protein per kilogram). Subjects lost

![Figure 5.Comparison of liver sections of rats fed control diet ad lib. or a low protein diet ad lib. (LoPro). (Top) ×20; (Bottom) ×100.](image-url)
Table IV. Retroperitoneal Fat, Gastrocnemius Muscle, Adrenal, and Thymic Wet Weight Fraction on Day 30

<table>
<thead>
<tr>
<th>Tissue Weight/Body Weight × 10⁻³ g</th>
</tr>
</thead>
<tbody>
<tr>
<td>Control</td>
</tr>
<tr>
<td>RP fat</td>
</tr>
<tr>
<td>Muscle</td>
</tr>
<tr>
<td>Adrenal</td>
</tr>
<tr>
<td>Thymus</td>
</tr>
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</table>

The mean±SEM (n = 7–8) is represented. Superscripts denote significant differences between groups; means not sharing the same superscripts are different at P < 0.05. RP, retroperitoneal.

an average of 3 kg during 1 wk of restriction with no change in serum IGF-I levels. The results of these human studies appear similar to the minimal IGF-I decline (12%) observed in rats restricted to 60% calories, with greater decreases associated with the more severe diets (Table II).

We controlled multiple nutritional variables so that the effects of energy and protein deficiency per se on circulating GH and IGF-I levels could be directly compared, without micronutrient or fat undernutrition. This is the first report of protein and carbohydrate restriction impact on molecular distribution of IGFBP-3, IGFBP-3-specific protease activity, and GHBP levels. In the rodent, GHBP is derived by alternative splicing during the processing of the full-length GH receptor pre-mRNA, and GHBP levels can be correlated with hepatic GH receptor binding in adult animals (39, 40). Although the primary source of GHBP is thought to be the liver, the broad tissue distribution of GHBP mRNA expression suggests that GHBP may have autocrine/paracrine roles in modulating GH action or in regulating GH presentation to or removal from target tissues (40).

While there are no adult animal studies that specifically limit dietary carbohydrate calories, 65% restriction of overall food and micronutrient intake suppressed pulsatile GH release and GH levels in adult rats (29). In the current study, the groups limited to 60% and 40% of ad lib. caloric intake had markedly reduced serum GH by day 30. This suggests that GH status is ultimately sensitive to carbohydrate calories independent of protein, fat, and micronutrient availability. Though pulsatile GH was not measured with intense blood sampling, there is a strong and significant trend suggested from the single samples. The 40% diet induced an order of magnitude inhibition of GH by day 21, while the 60% diet preserved higher ambient GH levels throughout at least 3 wk. This observation in 40% vs 60% groups was independently confirmed in a recent 21 day study (data not shown). The 40% caloric group was also the only set of rats that had reduced GHBP and lost body weight throughout the month.

In a previous study, 1-mo-old male Sprague-Dawley rats fed 70, 60, or 50% of ad lib. intake for 10 d had 39, 58, and 80% reductions in liver IGF-I mRNA, respectively (11). In that report, liver IGF-I mRNA levels were determined by Northern blot analysis using a cDNA probe which included the entire region protected by our exon 3 and 4 riboprobe. In contrast, we found that a reduction of energy intake to 60% did not significantly decrease liver IGF-I mRNA, either when expressed per microgram RNA or per picogram cyclophilin mRNA. In these 3-mo-old rats, a severe energy restriction to 40% significantly reduced liver IGF-I mRNA. The dissimilarity between our findings and those of Straus and Takemoto (11) may reflect a differential sensitivity of liver IGF-I gene expression to dietary restriction as a function of age. Such a finding has already been described for protein malnourished rats (24).

Inhibition of GH may be necessary but is not sufficient to trigger the alteration in IGF-I and IGFBP-3 with severe energy restriction. Both of these proteins continued to fall after the first week, and then stabilized over the latter part of the month, indicating the value of longer protocols to understand the timing and degree of hormonal adaptation to calorie undernutrition. We conclude that there is a clear threshold of carbohydrate availability, between 40 and 60% of ad lib. calories, necessary to maintain body weight, serum GHBP, and IGF-I levels in adult rats.

In the protein-deficient state, decreased IGF-I production (13, 14, 18) and increased IGF-I clearance (17) with normal GH (14, 19), reduced GH (20, 41) and normal GH binding (14, 18) have been described. For example, Lemonzy et al. (22) recently found in pubertal female rats a 50% reduction of serum IGF-I after 8 d of a 5% (vs 15% control) protein diet. In our study, serum GHBP was surprisingly unchanged at day 30 in the low protein group (unlike severe calorie restriction) while serum GH only trended toward lower values. The IGF-I levels were down by a third at day 7, and then maintained at this level, while hepatic IGF-I mRNA was significantly lowered with low protein availability, confirming shorter studies in younger animals (19, 23). Because liver is the major source of serum IGF-I and IGFBP-3, steatosis, a well-documented result of protein malnutrition (42, 43), may have hindered synthesis of IGF-I and IGFBP-3. Alternatively, limited essential amino acid availability from this diet could explain these responses, as suggested by diminished IGF-I gene transcription in cultured hepatocytes exposed to reduced low amino acids (44). However, normal GHBP levels at day 30 in the low protein group challenges this concept of a generalized hepatic dysfunction, as liver is thought to be the predominant source of circulating GHBP (40).

A role for IGFBP proteases has been postulated to contribute to the decline in IGFBP-3 (19). The hypothesis we tested for the first time was that these enzymes participate in the response to catabolic conditions, possibly promoting tissue availability or clearance of decomposed IGF-I and/or IGFBP-3. Increased IGF-I clearance was noted in young protein-restricted rats (17), though the mechanism was not ascertained. Furthermore, proteolytic activity directed at IGFBPs was reported in pregnancy (25, 26), surgery (28), and severe illness (27). Given these published data, we predicted that an IGFBP-3 protease would be more active in severe protein and/or calorie malnutrition. It was also expected that there would be a conversion of the 150-kd complex containing IGF-I into a smaller form. Contrary to these speculations, at day 30 there was a change in neither IGFBP-3 protease nor the distribution of the 150-kd complex. We conclude that IGFBP-3 proteolysis does not contribute to the decline of serum IGF-I or IGFBP-3 during marked protein or calorie deficiency in this model. We now postulate that the fall in serum IGFBP-3 is via a drop in its transcription or translation. This would be supported by the single data point indicating lower liver IGFBP-3 mRNA in rats suckled on dams fed at 50% of normal (45).

The greatly diminished adiposity of the caloric- vs protein-restricted state (Table IV) confirms that the compensatory
mechanisms reacting to each deficit are fundamentally different. In adult rats, protein deficiency results in reduced protein synthesis, which leads to a decrease in muscle mass due to the destruction of muscle proteins. Protein restriction results in the redirection of amino acids to other tissues, such as the liver, for energy production, further contributing to muscle wasting.

In young rats, protein restriction results in growth retardation, decreased circulating somatomedins, and reduced growth hormone output. These changes are associated with reduced circulating levels of insulin-like growth factor-I (IGF-I) and reduced growth hormone binding protein-3 (IGFBP-3). These changes are likely due to the decreased availability of amino acids for protein synthesis, leading to decreased growth hormone production and reduced circulating levels of IGF-I. These changes are associated with reduced growth and increased fat deposition.

In summary, the comparative adaptation of the somatotrophic axis to chronic, substantial, and specific protein or calorie deficiency was evaluated. Over a 30-day study in adult rats, serum GH was markedly reduced in the 40% calorie group by day 21, but was more modestly lowered in the 60% calorie group and low protein animals. Interestingly, serum GHBP was significantly decreased by about half only in the 40% calorie group. More consistent across groups was the finding that restricted calories or protein depressed serum IGF-I levels on day 30, in the following rank order: 60% (~12%), LoPro (~33%), 40% (~42%). Chronic, severe energy (40% calorie group only) and protein deficits were required to observe a reduction in serum IGFBP-3, liver IGF-I mRNA, serum total protein, albumin, and blood urea nitrogen. These dietary regimens had no measurable impact on the association of IGFBP-3 with the circulating 150-kD complex, which includes IGF-I and the acid-labile subunit. Reduced IGFBP-3 concentrations were also not due to a substantial increase in IGFBP-3-specific protease activity, characteristic of other conditions of lowered IGF-I and IGFBP-3.

We conclude that severe calorie and protein restriction of young adult rats produces only a modest fall in circulating IGF-I (32–43% at day 30), which is likely mediated by reductions in both liver IGF-I mRNA and serum IGFBP-3. These and other data indicate that there are broad ranges of nutrient intake and dietary composition which support normal circulating IGF-I and IGFBP-3 in young adult rodents and humans.

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References


