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GROWTH OF CHILDREN AFTER LIVER TRANSPLANTATION (LT):
LONGITUDINAL AND CROSS-SECTIONAL STUDY
Aims: to study the growth of children with liver disease before
and after LT. Methods: longitudinal study (LS): 16 infants (9 M, 7 F);
age 0.73 to 2.38 years (\$\frac{1}{2}\$ 1.39) at LT; mean height (Ht) SDS at LT:
2.02 (SD 1.25). Post-LT medication: cyclosporine in tapering doses:
prednisone, by daily regimen (DP) for the first 6-12 months, with
tapering doses depending on clinical course, then on alternate day
regimen (ADP). Cross-sectional study (CSS): 95 children (55 M, 40
F) aged 0.17 to 14.88 (\$\frac{2}{3}\$.83), observed before LT, after LT on DP,
and on ADP (same medication). Results: LS: pre-LT height velocity
(HV) SDS (\$\frac{2}{4}\$.59 -0.60\pm 1.31) significantly lower (p>0.01) than ADPHVSDS (2.84\pm 1.59) but not significantly different from DP-HVSDS (1.37\pm 1.66). CSS: DP-HtSDS (\$\frac{2}{4}\$.5D -1.91\pm 1.32) significantly lower
(p>0.01) than pre-LT-HtSDS (-1.07\pm 1.06) and ADP-HtSDS (0.98\pm 1.23). Similar results for sitting height (SH) SDS and
subischial leg length (SLL) SDS: pre-LT and ADP-SLLSDS
significantly greater than pre-LT and ADP-SHSDS (p=0.01); DPSLLSDS not significantly different from DP-SHSDS. Significant
improvement of head circumpherence SDS and skinfold SDS on
ADP. Conclusions: growth in children with liver disease does not
improve after LT on DP while shows catch-up growth on ADP;
therefore, it mainly depends on clinical course and corticosteroid
regimen.

### 261

CONCORDANCE BETWEEN BIOLOGY AND A VISUAL MOTOR PSYCHOLOGICAL TEST IN SHORT CHILDREN. A. Fiellestad-Paulsen, A. Andronikof-Sanglade, S. Ricard-Malvoir, D. Evan-Brion. Department of Pediatric Endocrinology, Höpital Robert Debré, Paris, France.

We explored in 40 prepubertal short children without organic disease (age 5 to 12 y., mean height - 2.6 ± 0.6 SD), the possible expression of an abnormal GH-secretion in a neuropsychological figure drawing test analysed with an original methodology. Nocturnal GH-profile, L. Dopa test and GHRH test were performed. The children underwort the psychological assessment, including the Rev. Observed. methodology. Nocturnal GH-profile, L-Dopa test and GHRH test were performed. The children underwent the psychological assessment, including the Rey-Osterrieth test and an IQ evaluation before medical investigation. The Rey figure was scored anonymously according to the boundary type errors defined as non developmental severe distortions or disrespect of boundaries. 23 children had normal and 17 had abnormal GH-secretion. Among those, 4 children had complete and 13 had partial GH deficiency but all of them had low nocturnal GH-secretion (AUCb < 85 ng/mV12h). 17 children had a normal Rey figure and 23 had boundary type errors Mean IQ was normal in both groups. The biological and the psychological analysis were in agreement in 32 children ( $\rho < 0.0005$ ). Night GH-secretion was lower ( $\rho < 0.006$ ) in children with boundary type errors (AUCb mean  $\pm$  SEM 88.5  $\pm$  8 ng/mV12h) as compared to children with normal Rey figure (AUCb 140.7  $\pm$  17.9). In conclusion, these results point to particular anomalies, independant of the IQ, at a visuomotor psychological test in short children with abnormal GH-secretion.

## 262

GROWTH HORMONE RESPONSES TO GROWTH HORMONE RELEASING PEPTIDE (GRP) AND TO GROWTH HORMONE RELEASING HORMONE (GRF) IN GROWTH HORMONE DEFICIENT CHILDREN (GHD). V. Mericq, F. Cassorla, H. García, A. Avila, C. Bowers and G. Merriam. Institute of Maternal and Child Research, University of Chile, Santiago, Chile; DEB, NICHD, Bethesda, MD; University of Washington, Seattle, WA; and Tulane University,

GRP is a potent and specific stimulator of growth hormone (GH) secretion. It is a 6 aminoacid peptide with lower molecular weight and longer half life than GRF. To clarify how this peptide acts, we administered separately and together 1 ug/kg bolus doses of GRP and GRF iv to 22 children (13 M, 9 F), ages 3-16 years, with previously documented GHD. Tests were separated by at least 1 week. Bone ages ranged from 1-10 years and growth velocities were less than 3 cm/year. GH was measured by RIA with an intraassay cv of 5 %. A positive response was defined as a GH increase greater than 4 cv's. We observed 12 (55%) positive responses to GRP, and 15 positive responses to GRF (68 %). Nineteen (86 %) patients responded to both peptides administered together. Out of the 12 GRP responders only 9 patients had a positive response to GRF, and out of the 15 GRF responders only 9 had a positive response to GRP. GH peak levels were observed between 5 and 60 min (mean 30 min) after GRF, and between 10-45 min (mean 24 min) after GRP. We conclude that GRP is a potent secretagogue of growth hormone in a substantial proportion of GH deficient children. The dissociated response to GRP and GRF suggests that they stimulate GH release through different mechanisms.

## 263

EFFECT OF GROWTH HORMONE (GH) ADMINISTRATION ON TOTAL AND SEGMENTAL BODY COMPOSITION IN CHILDREN WITH GH DEFICIENCY. Osorio, MGF; Mendonca, BB; Segura, TC; Estefan, V; Arnhold, IJP; Nicolau, W; Bianco, AC, Division of Endocrinology, Hospital das Clinicas, University of Sao Paulo, and Unidade de Densitometria Ossea, Sao Paulo, S.P., Brazil.

Sao Paulo, and Unidade de Densitomerría Ossea, Sao Paulo, S.P., Brazil. To investigate the GH-induced changes in body composition we studied the lean and fat tissues, bone mineral content (BMC) and density (BMD) of GH-deficient children by DEXA (Lunar, WI), before and after they were 174±14 days on hGH (0.075 U/Kg/day). Eight pre-pubertal children (4 boys), aging 6.2-14.1 yr (bone ages: 2-9 yr), were enrolled. All had peak GH responses to clonidine and insulin <5 ng/ml. BEFORE GH was given, body fat (4.2±2.5 kg) was distributed as 31±15 % in arms, 25±10 % in legs and 20±10 % in abdomen (abdm), while body lean tissue (13.7±3.1 kg) was 47±3 % in abdomen (abdm), while body lean tissue (13.7±3.1 kg) was 0.64±0.16 kg, of which 47±7 % was in the head, 25±4 % in legs and 9±1 % in arms. Total BMC was 0.64±0.16 kg, of which 47±7 % was in the head, 25±4 % in legs and 9±1 % in arms. Total BMD (g/cm2) was 0.8±0.04; it was higher in the head (1.4±0.1), followed by legs (0.6±0.074), pelvis (0.6±0.05), spine (0.6±0.01) and arms (0.5±0.03). GH treatment resulted in (i) height increase (4.6±1 cm; p>0.05) and no change in body weight (+0.9±1.4 Kg; p>0.05); (ii) decreased body fat ( $^{2}$ 3±16 %; p<0.05:  $^{2}$ 42±32 % in arms,  $^{2}$ 30±10 in abdomand  $^{2}$ 9±20 % in legs); (iv) increased BMC ( $^{2}$ 66 %; p<0.05:  $^{2}$ 54 % in the head; +21±21 % in arms; +18±9 % in legs); (v) decreased BMC (2.3±2.9 %; p<0.05: -6.4±4.3 % in the head, +6.9±6.6 % in pelvis, +4.3±3.1 % in arms, +3.7±2.2 % in legs). In conclusion, GH treatment caused body fat to decrease and lean mass to increase; because BMC increased significantly, the lower BMD is probably related to increased bone surface.

# 264

GONADAL STEROIDS ARE CAPABLE OF ACTING WITHIN THE HYPOTHALAMUS TO MODULATE SOMATOSTATIN (SS) GENE EXPRESSION IN PUREBRIAL ASED RATS. P.M. Martha, Jr. J.A. Brewer and E.O. Reiter, Department of Pediatrics, Baystate Medical Center, Springfield, MA 01199, USA.

P.M. Martha, Jr. JA. Brewer and F.O. Reiter, Department of Fediatrics, Baystate Modical Center, Springfield, MA 01199, USA.

Due to the highly complex nature of neuroendocrine-target organ "feedback" relationships, the primary site(s) at which gonadal steroids exert effects on the Gd axis remain largely unknown. Therefore, we sought to test the hypotheses that testosterone (T) and estradiol (E2) can exert direct cult-specific regulatory effects in vivo on hypothalamic neurons which control Gd secretion. The Meret Steroid of the parameters of the provided of the parameters of the provided in the periventricular nucleus (PaN) at the level of the parametricular nucleus (PaN) at the nucleus of the level of the parametricular nucleus (PaN) at the nucleus of the level of the panametricular nucleus (PaN) at the nucleus of the level of the panametricular nucleus (PaN) at the nucleus of the nucleus (PaN) at the nucleus of the level of the panametricular nucleus (PaN) at the nucleus of the nucleus (PaN) at the nucleus (PaN) a

### 265

LONGITUDINAL ASSESSMENT OF IGFBP-3 CONCENTRATION IN NORMAL BOYS.

A LONGITUDINAL ASSESSMENT OF IGFBP-3 CONCENTRATION IN NORMAL BOYS. P.M. Martha, Jr., R. M. Blizzard and A. D. Rogol, Departments of Pediatrics, Baystate Medical Center, Springfield, MA and University of Virginia, Charlottesville, VA, USA.

Recent data indicate the serum concentration of IGFBP-3 (the major post-natal IGF binding protein) may be clinically important in some pathophysiologic states. Existing cross-sectional data suggest an ontogeny exists for IGFBP-3, with levels increasing with age and peaking during puberty. To investigate the pattern of change in IGFBP-3 within individuals, and expand understanding of this potentially important component of the human GH axis, the IGFBP-3 level was determined in 309 serum samples obtained from 23 boys over 4.1-6.4 yrs as they progressed 'hrough puberty (assayed at Endocrine Sciences). IGFBP-3 conc. ranged from 1.7 to 7.0 mg/L. Though there was a general trend of rising IGFBP-3 conc. with age (intrasubject r=0.33, p<.001), values within individuals often fluctuated unpredictably. However, intrasubject variability was signif. less than that present in the larger population (CV=19.8±1%ys26\*, p<.001). Within individuals, IGFBP-3 level correlated with growth velocity (r=0.38, p<.001). For all data, IGFBP-3 conc. correlated with BA (r=0.43), testosterone conc. (r=0.31) and height SDS (r=0.26; all p<.001). We conclude that the serum concentration of IGFBP-3 follows the general trend of an increase with age and pubertal stage suggested by previous cross-sectional studies. There is significant fluctuation over time within individuals suggesting some caution when using this as a diagnostic tool though the range of this intra-individual variation is tighter than that of the larger "normal" range.