299 MACROPROLACTINOMA IN ADDLESCENTS: THEEE CASES THEATED WITH BROWOCRIPTINE. Reves, M.C., Campusano, C., Cattani, A. Decartments of Pediatrics and Endocrihology, Catholic University, Santiago, Chile. Macroprolactinomas (M) seem to be infrequent in individuals below 18 years of ace. In acults Bromocriptine (BC) therapy has been successful, but few data have been reported in addlescents. We report the clinical course and hormonal response to BC in three addlescents with M. (Case 1] A 14-year-old cirl with secondary amenorihea, calactorrhea and headache, Height (M:1-20); Cf scan : infræsellar mass with schenoldal sinus invasion, 20 mm in djameter: Goldmann's perimetry (GP); normal. She received 7.5 mg of BC peri-crimer stage IV. PRI 20 morporolactinemic with 20 mc factor (adv), and after 4 months PRI and menses were normal. After two years of follow-up, she is normoprolactinemic with 20 mc factor of the de-commeter resolution of timer (Davis) and 10 second 10 fb. Symmetr stage III et 4.5 mg/ml; fb ey off, 15 ey off, 10 km promet stage III et 4.5 mg/ml; fb ey off, 15 ey off, 10 km strange (608) and normalization of visual fields (Cases do 193 pyrini, but the emenormed has persisted. She experienced tumor strinkage (608) and normalization of visual fields (Cases 4) an 18-excess adominal fat and Tamer stage III. PRI 1518 ng/ml; GF strinkage (608) and normalization of visual fields (Cases 4) and 8-excess adominal fat and Tamer stage III. PRI 1518 ng/ml; GF strinkage (608) and normalization of visual fields of the resting MC, Alter 1.5 years he presented catch-up-croact, progression of tumor. After 1.5 years he presented catch-up-croact, progression of the umor with delay in growth and puberty. Height of 8C (10 tumor. After 1.5 years he presented catch-up-croact, progression of the umor with visual field deficits or monoprolactinemic with 7.5 m 6 daily of 8C to conclusions I in our experience 8C was effective in the treatment with strinkage upproves suroical ageproach, even if hormo

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6.21.8 1.11.2 2.611.2 3.212.5 1.41.5 1.11.1 2.011.0 1.81.6 ANOVA, 2 factors: px0.01:a vs a.c vs c.d vs d. px0.0510 vs b.e vs e.f vs f.g vs g. In conclusion: During the first year of life serum LH levels were clearly higher in povs than in girls. However, we observed that serum FSH levels were significantly higher in females than in males after the onset of puberty. These sex differences could be due to different inhibitory effects of gonadal steroids and/or petides on the concotation or to sevual differential secretion in the GRH pulse generator, we found very low serum LH concentrations in the propubertal period (these levels, cuid only be detected by IFMA), tollowed by an abruct increase in LH secretion (30- to 70-Iold) at the orset of puberty. Genum FSH levels, clearly higher during the propubertal period, did not change substantially during the propubertal period, and not change substantially during puberty, curing childhood may be less effective on FSH than on LH secretion.

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COMPARISON OF GROWTH HORMONE (GH) LEVELS MEASURED BY IMMUNORADIOMETRIC (IPMA) AND IMMUNOFLUORIMETRIC ASSAYS (IPMA) IN CHILDREN WITH SHORT STATURE. MATUL S. Lemos, M.M. Cassina, C., Sorio, G., Oliveira, S.R., SIVA, E., Satista, M.C., Nicolai, W., Arphold, I.J.P., Mendonca, B. Department of Endocrinology and Laboratory of Redioimmunoassay - HCFWISP, Sao Faulo, Brazil. Traditionally, the diagnosis of GH Ceficiency is based on maximal GH values after two stimulation tests under / ng/ml measured through IBMA. There are now more sensitive methods to measured through IBMA. There are now more sensitive methods to measured by IRMA and IFMA. In this study we compared the GH response measured by IRMA and IFMA in a group of 9 children (SM, SF) with short stature (SD for stature between -1.2 and -3.7), who were considered normal through two pharmacological tests for GH measured by IRMA. We also assessed by IRMA the GH responses to stimulation tests in patients with proven GH deficiency (transection of picultary stalk associated with propolastic picultary deficiencies) of Strong Clinical evidence [SD for neight <-C1.06] and low values of 10671. The IRMA and IFMA detection limit was 0.25 ng/ml and 0.1 ng/ml respectively.

SEX	AGE	HEIGHT	SDS	TEST	Bas IR	A Peak	Bas IF	MA Peak
FMM	445	102.5	-2.6	CLO CLO ITT	(ng/ml) 0.5 2.2 1.1_	17.3 18.5	0.24 0.72	17.9 10.7 13.6
والالالال	10.3	109.7 122.0 118.5 126.0		CLO LTT	0.47 0.344 8.1	16.02 11.22 14.3	0.28 0.1 2.7	947.9
ň	11.9	130.7	-2:9	CLO	0.8	²⁰ .3 7.3	0.3	8.5

In the GH deficient group, the patients had, no response to stimulation tests (< 0.1 to 0.2 pg/ml at all times): in two patients the maximal values were 0.8 and 1.7 ng/ml. We observed a positive correlation (r=0.899, p< 0.0001) among the 37 GH samples measured by both methods (GH values ranging from 0.31 to 35.1 ng/ml in IRVA, and from 0.1 to 17.9 ng/ml in IRVA) in the group with normal responses. We conclude that the GH values measured by IrVA are lower relaxed to avoid misdiagnosis of GH deficiency.

32 RAT GH RECEPTOR/GH-BINDING PROTEIN MENAS WITH DIVERGENT 5' -UNTRANSLATED REGIONS ARE EXPRESSED IN A TISSUE- AND TRANSCRIPT-SPECIFIC MANNER. Domené, H.M., LeRcith, D., Roberts, Jr. C.T., Cassorla, F. Developmental Endocrinology Branch, NICHD, and Diabetes Branch, NIDDK, NIH, Bethesda, Maryland, U.S.A. In the rat, the growth hormone receptor (GH-R) gene generates two transcripts, one that encodes for the GH-R, and a shorter one that encodes for the GH-binding protein (GH-RP). The mRNAs encoding for these transcripts present a high degree of heterogeneity in the 5'-untranslated regions (5'-UTR). It seems likely that some of the exons encoding 5'-UTR variants may be flanked by distinct promoter regions. The activity of different promoters could result in the tissue-specific expression of these variants. To assess this possibility, we used PCR amplification to characterize the 5'-UTR variants of rat GH-R mRNA, and by using 5'-UTR-specific probes, we determined their pattern of expression in several tissues in the trat. In addition to two previously described variants (VI and V2), three new 5'-UTR variants were identified, extending 56 nt. (V3), 135 nt. (V4), and 209 nt. (V5) upstream of the ATG translation initiation codon. The study of tissue distribution revealed that variant VI and V5 exhibited a pattern of expression resembling that variant VI and V5 exhibited a pattern of expression resembling that variant VI and V5 exhibited a pattern of expression resembling that variant VI and V5 exhibited a pattern of expression resembling that variant VI and V5 exhibited a pattern of expression resembling that variant VI and v5 exhibited a pattern of expression resembling that suppressed at low levels. These findings support the concept that different 5'-UTR variants are the result of different promoters expressed at low levels. These findings ruports that generates GH-BF in the rat might be controlled by the 5'-flanking region driving specific leader exons.

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REGULATION OF INSULIN DEGRADING ENZYME.

REGULATION OF INSULIN DEGRADING ENZYME. <u>Pérez, A.</u> Cambercs, M.C., Zuazquita, A., Cresto, J. C. CEDIE, Ricardo Gutiérrez Children's Hospital, Buencs Aires, Argentina. The main enzyme that triggers and controls insulin degradation is the insulin degrading enzyme (IDE). Many mechanisms have been postulated for IDE regulation but none has been conclusively proven. Highly purified rat liver cytosolic IDE was prepared by: 1) precipitation with ammonium sulphate, 2) DEAE-Sephadex with NaCl gradient, 3) pentylagarose with ammonium sulphate gradient, 4) chromatofocussing in FBE94. Insulin degradation by IDE was inhibited with ATP (0.05-4 mM) and GTP (1-8 mM) in cose/degendent fashion. AlF² (0.05-40 mM) had the same effect in the presence of Mg⁴+, but not NaF. Mg⁴+ suppression does not change AIF, inhibition. G-protein participation in this inhibition was excluded since these are activated with AIF₂, only if Mg⁴+ is present.

Since these are activated with Air, only if more is present. We conclude: 1) ATP inhibits IDE at physiological concentrations, while GTP acts as a phosphate donor at the concentrations used: 2) the G-protein participation in IDE inhibition could not be demonstrated in our system.

54 EPIDEMIOLOGY AND IMMUNOGENETICS OF INSULIN-DEPENDENT DIABETES MELLITUS IN VENEZUELAN CHIDREN. Gungzler, P., Lanes, R., Layrisse, Z., Balducci, P., Esparza, B., Salas, R., Arnaiz-Villena, A. Clinicas Hospital Caracas. Scientific Research and Hygiene Institutes, Caracas, Venezuela and 12 of Octubre Hospital, Spain. We evaluated 91 newly diagnosed IDEM children mean age 7.8 ± 4.5 yrs; 56.7% had had an upper respiratory infection prior to diagnosis and 12.7% had had either mumps or varicella. Peak incidence of disease was found in February and March and August to October. Eighty seven percent had HLA-DR3 and/or DR4 vs 37% of the Venezuelan general population, 81.6% were HLA-DQW2 and/or HLA-DQW8.Studies of oligonuclectid hybridization showed the presence of arginine in position 52 of the DQ alpha chain and absence of aspartic acid in position 57 of the DQ beta chain, with an increased prevalence of IDR2 and especially with DQB1 OGC2 which has been associated with protection. We found 55.9% to have positive islet cell antibodies (ICA) with 4 of these having a positive complement fixation test. Three patients (7.9%) were found to have positive insulin autoantibodies. No positive serotypes for enterovirus (Coxsackie-B) were found in our patients, but we detected 11 cases with elevated titers for cytomegalovirus antibodies. Positive antibodies for immanogenetics of insulin dependent diabetes mellitus in Latin-American children.