CATCH-UP GROWTH IS INTRINSIC TO THE EPIPHYSEAL GROWTH PLATE. J. Baron, K.E. Oerter, J.A. Yanovski, J.A. Novosad, J.D. Bacher, G.B. Cutler, Jr., Developmental Endocrinology Branch, NICHD, NIH, Bethesda, MD 20892, USA.

Catch-up growth is believed to result from a central nervous system mechanism that compares actual growth to a set point and adjusts growth velocity accordingly. Alternatively, we hypothesized that catch-up growth is intrinsic to the growth plate. To test this hypothesis, we infused dexamethasone phosphate for 4 weeks directly into one proximal tibial growth plate of 5-week old rabbits and infused vehicle into the contralateral growth plate addetermined radiographically. Dexamethasone decreased the growth velocity of the treated growth plate by 37 \pm 8 % (mean \pm SEM, p< 0.01) compared to the control, causing a growth deficit of 4.8 \pm 1.2 mm (p<0.01). After the end of the infusion, the growth velocity of the dexamethasone-treated growth plate, control and ultimately corrected 52 \pm 10 % of the deficit (p<0.01). Because the catch-up growth plate, we conclude that catch-up growth is intrinsic to the epiphyseal growth plate and cannot be explained by a systemic mechanism. Catch-up growth is believed to result from a central

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THYROID FUNCTION IN NON-GH-DEFICIENT SHORT CHILDREN DURING A PLACEBO-CONTROLLED DOUBLE-BLIND TRIAL OF GH THERAPY. G.M.Leong, S.R.Rose^{1,} K.M.Barnes, G.B.Cutler Jr. & the DEB GH Study Group,

<u>G.M.Leong.</u> S.R.Rose¹ · K.M.Barnes, G.B.Cutler Jr. & the DEB GH Study Group, NICHD, NIH, Bethesda, MD, 20892, ¹U. Tennessee. Memphis, TN, 38103, U.S.A. Growth hormone (GH) therapy has been reported to cause changes in thyroid function in GH deficiency including decreases in T4 and TSH and an increase in T3. We sought to determine if GH therapy alters thyroid function in non-GH-deficient short children. Twenty-two children (19 boys) were followed for 12 months while receiving either GH (Humatrope, Eli Lilly) 0.074mg/kg t.i.w. s.c. (n=10) or placebo (n=12). Total T4, free T4, T3 and TSH were measured every 3 months and in 13 children also at 1 and 2 months. A TRH test and nocturnal TSH surge were performed at baseline and 6 months in 19. There were no significant clinical differences at baseline between the placebo and GH groups. The results below are mean (SD) mean (SD).

	Months	0		1		2		3		6		12	
T 4	Placebo	8.3	(1.3)	8.0	(1.1)	8.3	(1.1)	7.6	(1.2)	8.1	(1.4)	7.4	(1.6)
(ng/dL)	GH	7.4	(1.2)	8.4	(1.3)	8.1	(1.6)	8.0	(1.5)	7.4	(1.5)	7.2	(1.3)
Free T4	Placebo	1.3	(0.2)	1.4	(0.2)	1.4	(0.2)	1.2	(0.2)	1.4	(0.2)	1.2	(0.1)
(pg/mL)	GH	1.3	(0.2)	1.3	(0.2)	1.3	(0.2)	1.2	(0.1)	1.3	(0.2)	1.3	(0.1)
T3	Placebo	181	(31)	170	(20)	177	(24)	171	(12)	162	(22)	175	(42)
(ng/dL)	GH	157	(21)	179	(12)	183	(22)	184	(24)	165	(22)	179	(31)
TSH	Placebo	2.8	(0.8)	2.2	(0.8)	2.2	(0.7)	2.6	(0.9)	1.8	(0.6)	3.1	(0.9
(uIU/mL) GH	2.5	(1.5)	1.8	(1.5)	1.9	(0.8)	1.8	(1.1)	2.8	(1.7)	2.2	(1.3
There we	re no sign	ifican	t diffe	rence	es bei	weer	h the	two g	roups	in a	iny of	the a	abov
measuren	nents or ir	1 TSF	I resp	onse	to TR	H or	in noc	lurna	I TSH	surg	ie. We	e con	clud
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that GH therapy over 12 months does not alter thyroid function significantly in non-GH-deficient children.

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LINEAR GROWTH IN THE RABBIT IS CONTINUOUS, NOT SALTATORY. K.E. Oerter, J.D. Bacher, G.B. Cutler, Jr., J. Baron, Developmental Endocrinology Branch, NICHD, NIH, Bethesda, MD 20892

A recent report (Science 258:801) suggests that human growth occurs in brief bursts separated by extended periods of stasis. We tested this hypothesis in an animal model in which growth rate could be measured with greater accuracy than that achieved in the human. Metal pins were inserted into the bone immediately adjacent to the proximal tibial and distal femoral growth plates of 6-week old rabbits. Growth rates were determined by measuring the change in distance between pins on serial radiographs taken daily for 10 days. All measurements were made in quadruplicate by an observer blinded as to the day and animal number. The error of measurement was small (0.05 mm/d) compared to the mean growth rate (0.27 mm/d). The individual growth curves revealed continuous growth. To analyze statistically the pattern of growth, we examined the frequency distribution of daily growth velocities. The model of saltation and stasis predicts a majority of daily growth velocities clustered around zero, and a minority of high daily growth velocities. The pattern we observed was significantly different (p<0.001), showing instead a single Gaussian distribution about an intermediate growth velocity, indicating continuous growth. We conclude that linear growth in the rabbit is continuous, not saltatory.

227 TREATMENT OF A BOY WITH GROWTH HORMONE INSENSITIVITY SYNDROME WITH rhIGF-I AND LHRH ANALOG. <u>D.R. Counts,</u> D.D., and G.B. Cutler, Jr., M.D., Developmental Endocrinology Branch, National Institutes of Health, Bethesda, Maryland, USA. The determine the effect of rhIGF-I treatment plus syndrome (GHIS), we treated an 18.5-year old mid-puber al boy with GHIS with rhIGF-I 100 ug/kg twice daily plus LHRH analog (deslorelin) 4 ug/kg/day. Pretreatment height was 111.2 cm (-10 S.D.), growth rate was 4.1 (m/year, bone age 14.5 years, peak plasma LH 32.1 LU/t, plus LHRH analog (deslorelin) 4 ug/kg/day. Pretreatment height was 111.2 cm (-10 S.D.), growth rate was 4.1 (m/year, bone age 14.5 years, peak plasma LH 32.1 LU/t, plus LHRH analog (deslorelin) 4 ug/kg/day. Dretreatment fGF-I increased from an undetectable level at baseline to 108 ng/mL at 1 h post administration. GH secretion, freat three months of therapy, growth rate had increased from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was unchanged, from a 4.1 to 8.7 cm/year, bone age was u

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THE RELIABILITY OF TWO RETHODS TO PREDICT ADULT HEIGHT IN 132 TALL GIRLS. <u>N.M.Draver</u> and L.Cox. Dept.of Paediatrics, Univ.of Groningen, 9713 EZ Groningen, the Netherlands and Institute of Child Health, Univ.of London, London WC1N 1EH, England.

How reliable is the ultimate height prediction (UHP) for tall girls? 78 Girls who were pre-menarcheal (pre-m) and 54 who were post-menarcheal(post-m) when initially measured during childhood, had their adult heights measured. All girls were then aged more than 17 years. They had all chosen not to receive treatment on the basis of the UHP.Height was measured according to the same standardised techniques using Karpenden stadiometers. X-rays of the left hand were assessed according to Greulich-Pyle (G-P) and Tanner-Whitehouse TWII RUS (RUS). For the UHP the Balley-Pinneau (B-P) and the Tanner-Whitehouse method for tall girls (TW) were used. pre-m (mean ±SD) post-m (mean ±SD)

	1	pre-m (mean ±SD)	post-m (mean ±SD)
nitial age	(yrs)	12.6 ±1.1	13.7 ±1.4
nitial height	(cm)	174.1 ±5.0	177.4 ±5.2
keletal age G-P	(yrs)	12.6 ±0.6	13.7 ±0.9
RUS	(yrs)	13.1 ±0.5	14.0 ±0.9
JHP B-P	(cm)	184.8 ±4.1	183.1 ±4.4
TW	(cm)	183.6 ±3.9	182.0 ±4.7
Adult age	(yrs)	20.4 ±2.4	21.7 ±2.7
Adult height	(cm)	184.5 ±3.6	182.4 ±4.2
B-P overpredicted	more	than 5.08 cm (2") for 4 pre-m and 2 po

B-P overpredicted more than 5.08 cm (2-) for 4 pre-m and 2 post-m girls. TW for 2 pre-m girls. B-P underpredicted more than 5.08 cm for 6 pre-m girls. TW for 11 pre-m and 1 post-m girl. The adult height for tall Dutch girls as a group can be predicted using methods developed more than 30 years ago for the USA and UK $= 10^{-1}$ cm s $= 10^{-1}$ populations.

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INSULIN-LIKE GROWTH FACTOR BINDING PROTEIN 3 (IGEBP-3) CONCENTRATION IN INSULAVLIKI GROWTH FACTOR BINDING PROTEIN 3 (GFBF-3) CONCENTRATION IN CEREBROSINAL FLUID (CSF) OF CHLDREN WITH BRAIN TUMORS OR LEUKEMIA. <u>HLMüller</u>, Y.Oh, S.E.Gargosky, R.L.Hintz, R.G.Rosenfeld, Department of Pediatries, Stanford University School of Medicine, Stanford, CA 94305, USA The major IGBPs in CSF are IGEPP-2 and IGEPP-4, whereas IGFBP-3 is a minor component in CSF of healthy subjects. We investigated IGFIP-3 levels in CSF from patients with brain tumors,

in CSF of healthy subjects. We investigated IGFBP-3 levels in CSI⁺ from patients with brain tumors, leukenia or meningitis. IGFBP-3 was measured by radioimmuno-assay with dIGFBP-3g1, a rabbit polycional antibody. Further, as proteolysis of IGFBP-3 is part of the modulation of IGF activity IGFBP-3 fragmentation was quantified by densitometric analysis of [¹²⁵1]IGFBP-3 protease assays. We examined CSI's of 24 children with malignant brain tumors, 18 children with leukemia and 13 children with meningitis. CSI's of 38 children, who received a lumbal puncture in order to exclude meningitis, were used to define a normalative range for IGFBP-3 concentrations were found in CSI⁺ of 17 of all 24 (71%) brain tumor patients and 7 of 8 (87%) brain tumor patients who had microscopically detectable malignant cells in CSF. 13 of 14 (93%) patients with medultoblastoma or ependymoma and 17 medultoblastoma/ependymoma patients with malignant cells in CSI⁺ fad elevated IGFBP-3 concentrations in CSI⁺. IGFBP-3 protease activity in CSI⁺ was clevated in 15 of 17 (88%) patients with histological high grade (WHO III/9/W) brain tumors. 5 of 6 (83%) patients with acute leukemia and microscopically detectable malignant cells in CSI⁺ at the time of diagnosis showed elevated IGFBP-3 to CSI⁺ that CSI⁺ that CSI⁺ that CSI⁺ that chevenia patients with acute leukemia and microscopically detectable malignant cells in CSI⁺ at the time of diagnosis showed elevated IGFBPI-3 concentrations in CSI⁺ that Tormatilized under chevotherapy. Leukenia patients withoutany to the CSI⁺ that CSI⁺ that Tormatilized under chevotherapy. Leukenia patients withoutany to the table to the tormatilized under chevotherapy. Leukenia patients withoutany the two tormatice activity in CSI⁺ that Tormatilized under chevotherapy. Leukenia patients withoutany the two tormatilized under chevotherapy. Leukenia patients withoutany to the table table malignant cells in CSI⁺ that Tormatilized under chevotherapy. Leukenia patients wi (JGFIP)-3 concentrations in CSF that normal IGFIP-3 concentrations in CSF. We conclude that detectable malignant cells in the CSF had normal IGFIP-3 concentrations in CSF. We conclude that in CSF of children with high grade malignant brain tumors or CNS leukemia, IGFIP-3 is elevated. In CSF of children with high grade maniform train to have so COS reducting, IGFMF-3 is elevated. We hypothesize that this phenomenon could be caused by local production of IGFBF-3 by the brain tumor tissue and secretion into CSF or by local secretion of IGFBF-3 by maniform that spread into CSF. This hypothesis is supported by the observation that only 2 of 13 (15%) manifolds patients had slightly elevated IGFBF-3 concentrations in CSF while high numbers of non malignant inflammatory cells were present in all cases. Further studies will analyse the origin of elevated IGFBF-3 levels in CSF of tumor patients and will show whether these findings are of diagnostic or presented will be the studies of the studies of the spread of the studies of the stud prognostic value.

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