131

M.L. Aubert, N.M. Gruaz^{*}, D.F. Carmignac^{*}, C.-D. Walker^{*}, and P.C. Sizonenko. Biology of Growth and Reproduction, Dept. of Ped., Univ. of Geneva Medical School, 1211 Geneva, Switzerland. MELATONIN-INDUCED INHIBITION OF SEXUAL MATURATION IN THE MALE RAT: EVIDENCE FOR CENTRAL INHIBITION OF GARH SECRETION.

We have shown that daily treatment with melatonin (MT) markedly delays sexual maturation in the male rat. The bservations that MT-treated rats have normal pituitary responsiveness to GnRH and low or absent pulsatile release of LH pointed out for a central inhibitory action of MT on GnRH release. The sensitivity of MT-treated rats to gonadectomy and sex steroid replacement was investigated: male rats receiving 100 µg MT daily or saline were castrated at 30 d and immediately treated with various doses of testosterone propionate (T), or oil until sacrifice at 40 d. In animals that received oil, if release was drastically reduced in MT-treated rats (46.0[±]7.3 ng/ml) as compared to controls (305[±]44) demonstrating that MT inhibits GnRH secretion in absence of sex steroid. Replacement with T showed that MT-treated rats were much more sensitive to T than controls, suggesting that MT potentiates the Θ feed-back action of T on LH release. Conversely, the ability of GnRH neurons of MT-treated rats to release GnRH in response to N-methyl-DL-aspartic acid (NMA), an excitatory amino acid analog known to cause LH release via GnRH, was found normal at 30 d MT-treated and in control rats. These data further indicate that MT-treated and in control rats. These data further indicate that MT induces central inhibition of GnRH release without impairing secretory processes of GnRH neurons and prolongs the high sensitivity to sex steroid Θ feed-back typical of infantile animals.

> J.M.B.Wennink*, H.A.Delemarre-van de Waal. Dept. of Pediatrics, Vrije Universiteit, Amsterdam, The Netherlands.

132 DIURNAL AND NOCTURNAL LH AND FSH SECRETION IN BOYS THROUGHOUT PUBERTY ESTABLISHED WITH A SENSITIVE IRMA.

LH and FSH values were studied in 30 healthy boys (CA:11.3-14.6 yrs, BA:10.3-16.2 yrs (TW2), Tanner stage G1-G5) Infor-med consent was obtained. Blood was sampled at 10' intervals from 12.00-18.00h and from 24.00-06.00h. An IRMA employing 2 monoclonal antibodies with a sensitivity of LH and FSH of 0.10 IU/1 and antibodies with a sensitivity of LH and FSH of 0.10 10/1 and 0.25 IU/1 respectively was used. Testosterone (T) was measured every hour applying a RIA with a sensitivity of 1.0 nmol/1. Results of mean FSH, of mean LH, of mean LH pulse amplitude (pA), of LH pulse number (pN), of mean Nadir Interval in min.(NI) and of mean T are presented as the median per Tanner stage.

	DAT							NIGHI					
	FSH	LH	pА	PN	NI	т	FSH	LH	pA	рN	NI	т	
Gl	0.69	<0.1	-	-	-	<1	0.77	0.15	0.35	2	120	<1	
G2	2.05	0.37	0.3	1		<1	2.01	1.77	1.55	4	70	<1	
G3	2.69	0.86	0.8	1	-	<1	3.05	3.42	1.77	4	80	1.8	
G4	2.76	0.87	0.5	2	130	1.4	3.28	3.93	1.68	5	65	10.5	
C5	4.82	1.88	1.1	2	120	8.6	4.30	3.89	2.96	3	110	13.2	
							berty(C t LH is						
							pA and p						

d. In C5 pN decreases, while pA still increases, presumable as a result of an increased negative feedback action of T. In contrast to LH, diurnal FSH is detectable in Gl, shows a clear increase already in G2 and has no sleep-wake pattern.

> Merete Jørgensen*, N.E. Skakkebæk, N. Keiding, C.T. Nielsen*, Janet A.B. Darling*, W.M. Hunter*, D.W. Richardson*.

133 Statistical Research Unit, University of Copenhagen, Denmark. Dept. of Paediatrics, Hvidovre Hospital, Den-mark. Dept. of Paediatric Biochemistry, Royal Hospital for Sick Children and MRC Reproductive Biology Unit, Edinburgh, Scotland.

TIME LAGS BETWEEN VARIOUS MATURITY MEASURES AND SPERMARCHE.

On the basis of biannual examinations of 40 normal boys followed On the basis of blannual examinations of 40 normal boys followed longitudinally for a period of up to 7 years (initially aged 8-11), the following maturity measures were studied; first occur-rence of a testis size of more than 4 ml., public hair stage gre-ater than Tanner stage one, broken voice, axillary hair greater than stage one, beard growth, occurrence of acne and the age of maximal height growth velocity and spermarche (onset of the release of spermatozoa).

Spermarche was estimated on the basis of occurrence of spermatozoa in quarterly collected 24-h urine samples, using the information of first occurrence of spermaturia and the frequency of intermittent occurrence of spermaturia after the first.

With confidence limits of about ± 4 months, the maturity measure: showed the following time lags. Prior to spermarche: testis size >4 ml (16 months), puble hair stage>1 ($10\frac{1}{2}$ months), first occur-rence of broken voice ($\frac{1}{2}$ month). After spermarche: max. height growth velocity ($5\frac{1}{2}$ months), axillary hair stage>1 (8 months), growth of beard (12 months) and occurence of acne (13 months).

The United Kingdom CnRH analogue Collaborative Study Group. (Introd. by C. G. D. Brook).

134

INTRANASAL (D-Ser⁶) GnRH ANALOGUE FOR THE MANAGEMENT OF CENTRAL PRECOCIOUS PUBERTY.

We have treated 61 children (51F, 10M) with central precocious puberty using intranasal (D-Ser⁶) GnRH analogue (Buserelin). Maximum duration of treatment was 3.7 yrs in the girls and 3.0 yrs in the boys. Mean dose (SD) of (D-Ser⁶) GnRH was 25.3 (9) μ g/kg in the girls and 18.6 (12) μ g/kg in the boys. No side effects were seen. 5 patients stopped treatment due to non-compliance. Of 27 children who had puberty stage 2 - 3 before treatment, 20 arrested or regressed and 17 advanced. Of 21 children treated at stage 4 - 5. 7 regressed and 14 had no

betroft treatment, 20 artested of regressed and 14 had no alteration. Mean testicular volume in the boys was reduced from 9.2 (6) mls to 5(1.4) mls after 1 year of treatment.
Mean basal serum LH became elevated during treatment, although

there was a decrease in peak stimulated LH concentrations.

 $_{ABA/aCA}$ fell during the first year of treatment to below 1.0. Height velocity in the girls reduced from 9.0 - 6.0 cms/yr during the second year of treatment. Height for bone age SDS increased from -1.86 (1.5) to -1.1 (1.5) during the first 1.5 yrs of treatment in the girls which was not significantly different from untreated controls. Intranasal (D-Ser 6) GnRH is an effective treatment for central

precocious puberty but does not alter height prognosis.

R. Kauli, L. Kornreich*, Z. Laron. Inst. Pediat. & Adolesc. Endocrinol. & Dept. Pediat. Radiol., Beilinson Med. Ctr., Sackler Faculty of 135 Medicine, Tel Aviv University, Israel. PUBERTAL DEVELOPMENT AND FINAL HEIGHT IN GIRLS WITH CENTRAL PRECOCIOUS PUBERTY (CPP) AFTER THERAPY WITH THE Gn-RH ANALOG D-TRP-6-LH-RH.

THE Cn-RH ANALOG D-TRP-6-LH-RH. Fifteen girls with CPP who have been treated with D-TRP-6-LH-RH (13 daily s.c.; 2 monthly i.m.) for 1 to 5 yrs were followed after therapy for 6 months to 4 yrs. Resumption of pubertal deve-lopment was evident clinically 2 to 4 months after therapy was stopped and menstruation appeared after 3-6 months in 13 and after 2 yrs in one. In all growth velocity increased in parallel to pubertal reactivation and bone maturation advanced at a moderate where G for R eight here reached their final height. rate. So far 8 girls have reached their final height.

Pt.	СА	ВА	of Therapy Predicted Final Ht. cm	Achieved Final Ht. cm
	y:n			
1	6:3	10:6	140.6	157.0
2	8:2	11:3	141.7	153.5
3	8:4	12:0	158.5	158.5
4	8:0	12:0	151.6	152.6
5	9:0	13:3	142.7	146.5
6	10:6	12:6	151.4	154.0
7	10:9	13:0	141.7	144.8
8	10:10	13:0	137.9	142.5
Our	data ind	icate t	hat Gn-RH analog thera	py in CPP has a benefi-
cial	effect (on fina	l height when started	at an early bone age.
				factor in final heigh

E. Frejaville*, C. Pifferi*, C. Cesari*, M.T. Tosi*, D. Tassinari*, A. Bergamaschi*, S. Salardi*, A. Cicognani, E. Cacciari. 136 Department of Pediatrics, S. Orsola - Malpighi Hospital, University

of Bologna, Italy. TREATMENT OF PRECOCIOUS PUBERTY BY INTRANASAL D-SER (TBU)6 LHRHI -9E A) (BUSERELIN).

GnRH analogues afford effective, selective, reversible inhibition of pituitary gonadotropin secretion. We administered Buserelin (300 ug x 6/die) to 18 girls affected by precocious puberty, for 6-24 months (onset of first symptoms before 8 years of age). We evaluated Buserelin's effects on pituitary-gonadal, pituitarythyroid, pituitary-adrenal function, ultrasonographic pelvic organs and thermographic mammary patterns, clinical and auxological features. Results: significant decreases (paired t test) were observed in: both LH and FSH reserves (LHRH test) and basal FSH since the first month (m) of treatment, E2 (3rd m), Progesterone (6th m); also ISH reserve (IRH test) since the 6th m, I3 (12th m) and fI3 (6th m) decreased significantly, though remaining in control range. 14, f14, PRL, ACIH and adrenal hormones did not change significantly. Ultrasonography of the pelvis showed significant decreases of uterine (6th m) and ovarian (12th m) volume; the ovarian structure did not progress and follicle diameter was always 9 mm. Clinical/thermographic evaluation of the breast showed a generalized arrest or regression of its maturational patterns. Auxological data (slower bone maturation and growth velocity, lower SD scores for chronological and bone age, adult height prediction tending to improve) suggested a good prognosis. We conclude that these results of intranasal Buserelin administration and the absence of serious side effects make it advisable in the treatment of girls with true precocious puberty.