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DISORDERS OF RYPOTHALAKO-PITUITARY FUNCTION AFTER MEAD INJURY IN CHILDREM. A. Russzzyńska-Moiska, I.E. Romer, N. Ginalska-Halinowska, B. Rymkiewicz-Kluczyńska, Department of Embocrimology, Child Meath Center, Warsaw, Poland Mead trawas can demage the hypothalamus as well as pituitary gland, which can in effect lead to disorders in secretion of tropic hormones or to disfunction of the pituitary isself. In Se aim of the study was evaluate frequency and types of hormonal disorders after head injury in children. The study was conducted on 77 patients (0.8-8 yr.) with collection of auxological data and hormonal stimulation tests (the insulin-hypoglicemia, the GR 1-29, the IRM and GmBH test and serum and urine osciality) which were carried out from 1 month to 7 yr. after the injury was sustained. Results are presented on tables.

1ab. 1. Type and nouter of hormonal disorders.

1ab. 2. GM response to GMRBH 1-29 stimulation in 37 patients with GM response to GMRBH 1-29 stimulation in 37 patients with GM response (10 mg/al in the insulin-hypoglicemia test).

13

29.8

42.8

5,5

9,1

2.6 63,6

Number of cases I of effected patients < 5 ng/el | 10

(5-10)ng/ml 23

2

Total 33

GH peak response

LH FSH

GHU LHY FSHY

GHL Cort. L

	Number of patients	Percentage		
GH) 10 ng/ml	31	83,8		
GH < 10 ng/m1	6	16, 2		

Tab. 3. The number of hormonal disorders in relation

Type of injury	Total number of cases	Number of disorders		
Serious injury with prolon- ged loss of consciousness	47	36	76,6	
Injury without loss of of conciousness	30	13	43,3	

Only 8/49 patients with hormonal disturbances had clinical symptoms - 3 showed height deficiency, 3 precocious puberty and 2 - symptoms of diabetes insipidus. Fourty one patients showed biochemical evidences of hormonal disfunction only. Our data confirmed that the regulation of 6d secretion is the most sensitive to injury - mainly on the hypothalamic level, and indicates that routine hormonal diaposatic tests are justified, especially in children who have suffered serious head trauma with prolonged loss of consciousness.

ESSENTIAL HYPERNATREMIA AS A RARE CAUSE OF CHILDHOOD OBESITY

C. Livieri, M. Bozzola, R. Lorini, F. Severi Department of Pediatrics, IRCCS Policlinico S.Matteo, University of Pavia, Pavia, Italy

University of Pavia, Pavia, Italy
We describe two unrelated patients, a boy and a girl both referred at the age of 4 yrs because of sudden onset obesity, polyphagia, hypodipsia, hypersomnolence, mood alteration with outbrusts of hysterical laughter or cry, episodes of muscular weakness. They were found to be suffering from an hypotalamic syndrome of unknown origin. Both showed no sense of thirst even with chronic hypernatremia and hyperosmolality, severe acrocyanosis, profuse sweating, episodes of enuresis with polyuria and excretion of inappropriately dilute urine. ADH determination, performed when the patients were in good metabolic control, was in the lower normal limits. Other endocrinological investigation showed hyperprolactinemia and low GH response to provocative stimulation in the two patients. EEG revealed nonspecific slow wave changes in the boy and multifocal high amplitude spikes and sharp transients in the girl. X rays of skull CT scan and MNR were normal. A defective osmoreceptor function is suspected in both patients.

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PIT-1 GENE EXPRESSION IN THE ANTERIOR PITUITARY IS MODULATED BY CHANGES IN CIRCULATING LEVELS OF TESTOSTERONE.

S. González-Parra* 1, JA. Chowen* 2, L. M. García-Segura* 2 and J. Argente*.

Autonomous University. The Hospital of Niño Jesús. Division of Growth, Endocrinology & Metabolism*. Cajal Institute, C.S.I.C. 2. Madrid, Spain.

Pin-1, or GHF-1, is a transcription factor specific to the anterior pituitary and is involved in the expression of the GH, PRL and TSH-0-subunit genes. The expression of these genes is modulated by changes in the steroid environment. Therefore, we asked the question as to whether this effect could be mediated, at least in part, by changes in Pit-1 expression. Adult male rats were castrated and implanted subcutaneously with a Silastic capsule that was either empty or that contained testosterone (T), Intact controls were sham-operated. Four days later, animals were sacrificed and pitularies were removed and processed for in situ hybridization to detect Pit-1 messenger RNA (MRNA). A S35 labeled riboprobe was prepared by in vitro transcription of a 915 bp insert of the coding region of the Pit-1 cDNA. This probe was applied to 12 µm sections of the pitularies and in situ hybridization methodologies were followed. Specific labeling was found only in the anterior pitultary.

PIT-1 mRNA in the anterior pitultary

Conclusion: Changes in circulating levels of sex steroids modulate the expression of Pit-1. Hence,

animals.

Conclusion: Changes in circulating levels of sex steroids modulate the expression of Pit-1. Hence the effects of sex steroids on the synthesis of certain anterior pituitary hormones may be mediated, at least in part, through the modulation of this specific transcription factor.

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ACTIVATION OF NICOTINIC CHOLINERGIC RECEPTORS MODULATES THE SOMATOSTATINERGIC SYSTEM IN THE RAT HYPOTHALAMUS Y. Barrios. 1, S. González-Parra 1 and E. Arilla 2 (Introduced by J. Argente), Autonomous University. The Hospital of Niño Jesús. Division of Growth, Endocrinology & Metabolism 1, Department of Biochemistry & Molecular Biology, University of Alcalá de Henares, Madrid 2, Spain.

Acetylcholine potentiales the excitatory effect of somatostatin (SS) on brain neurons and nicotine, a cholinergic drug, modifies catecholamine turnover in the hypothalamus. The hypothalamus shows a high concentration of nicotinic cholinergic receptors and a dense innervation of SS-positive nerve terminals. In light of these findings, we studied the effect of intravenous (i.v.) nicotine injection (0.3 mg/kg) on SS peptide levels and receptor binding in the hypothalamus of male Sprague-Dawley rats. A second experimental group was pretreated with mecamylamine (5 mg/kg), a centrally acting antagonist of nicotinic cholinergic receptors, in order to evaluate whether the effects of nicotion on the studied system involved the activation of these receptors. Control rats received an i.v. saline injection. The rats were killed 4 min after i.v. administration, and the hypothalamus was dissected to isolate SS and its membrane receptors.

Results: Nicotine produced an increase in somatostatin-like immunoreactivity (SL1) and in SS receptors. When the rats were pretreated with mecamylamine, the effects of nicotine were inhibited. Mecamylamine alone did not influence either parameter.

SLI (ng/mg protein) SS receptors

•	SLI (ng/mg protein)	SS receptors			
Groups		Bmax (fmol/mg protein)	Kd (nM)		
Saline	19.65 + 2.14	152 + 34	0.97 + 0.13		
Nicotine	160.76 ± 16.86 *	347 ± 26 *	0.95 ± 0.20		
Mecamylamine plus nicotine	17.55 ± 2.06	163 ± 14	0.94 ± 0.17		
Mecamylamine plus saline	18.98 ± 1.54	173 ± 10	1.04 ± 0.07		

Conclusions: 1. These results suggest that the rat hypothalamic somatostatinergic system is regulated by nicotine-like acetylcholine receptors. 2. The somatostatinergic system may be involved in some of the neuroendocrine effects of nicotine.

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NEONATAL TESTOSTERONE MODULATES THE NUMBER AND RESPONSIVITY OF GROWTH HORMONE-RELEASING HORMONE (GHRH) NEURONS.

A. Chown 1. S. González-Para* 2. LM. García-Segura 1, and J. Argenta².

Cajal Institute, C.S.I.C. 1. Autonomous University. The Hospital of Niño Jesús². Madrid, Spain.

Exposure to sex steroids during the fetal and neonatal periods plays an important role in the organization of the hypothelalamus. Since the rate of growth of the pubertal and adult animal is also affected by neonatal sex steroid exposure, we asked the question as to whether this phenomenon is due to an effect of these steroids on those hypothalamic neurons involved in stimulating GH, e., GHRH neurons. To address this question, the following experimental groups were studied:

Group name	IM					MCTAT		FNOAT	FNTAO	FNTAT
Treatment on day 0	and the second	Sham castrated			Castrated - TIni	Castrated - Tini	Oil	Oil	Tlnj	Tlnj
Treatment on day 60	Sham	Castrated	Slmp	Tlmp	Simp		SImp	TImp	SImp	TImp

lay 60 <u>bastrated Simp</u> <u>land 10 gold.</u> Tinjainjection with 250 µg T in 100 µg oil; Simp-empty Silastic capsule; Timp-Silastic capsule containing T. Growth was charted throughout development and was significantly affected by these treatments. Animals were sacrificed (day 75) and the brains removed and processed for in 3ru hybridization for GHRH mRNA. The total number of GHRH cells and the relative level of GHRH mRNA (analyzed by an automated image analysis system) were assessed in anatomically matched slides. Neonatal T had a significant effect on the number of detectable GHRH neurons in the hypothalamus (ANOVA: p-0.0001). Animals exposed to neonatal T had significantly more GHRH neurons than those that were not. Adult T-treatment did not affect

Group	IM	MAC	MCOAO	мсоат	мстао	MCTAT	FNOAO	FNOAT	FNTAO	PNTAT
Number of GHRH neurons		530 ±	315 ±	345 ±	417±	623 ±	297±	331 ±	404 ±	442 ±
Grains/cell	102 + 7	75 + 2	67 + 5	86 + 3	72+6	105 + 8	58 + 1	75 + 4	73 + 2	110 + 12

[Grains/sct] | 102 ± 7 | 75 ± 2 | 67 ± 5 | 86 ± 3 | 72 ± 6 | 105 ± 8 | 58 ± 1 | 75 ± 4 | 73 ± 2 | 104 ± 12 |
the number of detectable GHRH neurons, but significantly influenced levels of GHRH mRNA (ANOVA p.C.0001). Furthermore, adult T-treatement had a significantly greater effect in those animals that had received neonatal T when compared to those animals that did not receive neonatal T (2-way ANOVA p.C.05). These results suggest that one way in which exposure to sex steroids during the neonatal period affects the growth axis is by increasing the number of hypothalamic GHRH neurons, as well as to modulate the ability of these neurons to respond to changes in circulating levels of testosterone.

MOLECULAR ANALYSIS OF THE PROOPIOMELANOCORTIN (POMC) GENE IN 3 CASES OF CONGENITAL ISOLATED ACTH DEFICIENCY. J.-C. Carel, I. Tardivel, X. Bertagna, P.F. Bougnères and J.-L. Chaussain, INSERM U342 and Pediatric Endocrinology, Hôpital Saint Vincent de Paul, Paris, FRANCE.

We studied the POMC gene in 3 cases (1 boy, 2 girls) of isolated ACTH deficiency with manifestations of hypocortisolism before 6 months of age, undetectable ACTH after stimulation with LPH and/or CRF, normal secretion of the other pituitary hormones and normal appearance of the pituitary on C.T. scan or M.R.I.. One patient was born to consanguineous parents and one girl had an affected brother who died in the neonatal period. DNA from the 3 patients digested with EcoRI, Billi and PstI revealed a normal pattern after hybridization with POMC-genomic probes encompassing exons 1 and 3. After digestion with Sacl and hybridization with an exon 1 probe, a 10/15 kb polymorphism was detected and compatible with linkage of the disease to the POMC gene in the two families studied. PCR amplification of exons 1, 2 and 3 using In the two families studied. PCH amplification of exons 1, 2 and 3 using primers in the flanking intronic sequences gave products of the expected size in the 3 patients. Direct sequencing of exon 2 which contains the transcription initation site and 15% of the coding sequence revealed no difference with controls and with the published sequence. Sequencing of exon 1 and 3 is under progress. We conclude that these 3 cases of congenital ACTH deficiency are not due to deletions in the POMC gene or point mutations in exon 2.