458 Abstracts

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To understand the early molecular events that lead to some of the effects of growth hormone on liver, a dose of $^{8}\text{H-HGH}$ (18×10^{6} cpm, $60~\mu\text{g}$) was given i.v. to hypophysectomized rats. After the injection, livers were homogenized and fractionated by centrifugation. Specific activity (cpm/mg protein) and total radio-activity expressed as percent of that in the homogenate were estimated (table). 3H-HGH in the homogenate was estimated by precipitation with TCA, and with Na₂SO₄ as ³H-HGH antibody complex with and without unlabeled hormone. Up to 10 min most radio-activity was protein bound and ³H-HGH in the homogenate was antigenically similar to the injected dose. In contrast, when ³H-HGH was added in vitro before homogenization most radioactivity (77%) appeared in the cytoplasm and the rest distributed almost equally among the other fractions. These results indicate (1) the early rapid uptake by the particulate fractions may be of functional significance and the important initial binding sites may be mitochondria and microsomes; (2) the movement of 3H-GH in liver cell may be from the microsomal and mitochondrial fractions into cytoplasm; (3) after 10 min 3H-HGH is degraded or deacetylated.

Fraction	2 min	5 min
Homogenate	1517 (100%)	2816 (100%)
Crude nuclear	866 (18.3%)	1232 (11.2%)
Mitochondrial	1638 (22.5%)	5043 (41.0%)
Microsomal	4776 (41.9%)	5841 (29.2%)
Cytoplasmic	1464~(27.5%)	2177 (23.2%)
Fraction	10 min	20 min
Homogenate	3246 (100%)	3901 (100%)
Crude nuclear	1707 (13.3%)	2042 (13.8%)
Mitochondrial	6308 (29.9%)	4622 (21.8%)
Microsomal	4754 (15.3%)	2954 (8.5 %)
Cytoplasmic	3294 (23.5%)	7186 (46.1%)
Fraction	40`min	, , , , ,
Homogenate	1997 (100%)	
Crude nuclear	845 (13.5%)	
Mitochondrial	1442 (16.1%)	
Microsomal	1302 (7.9%)	
Cytoplasmic	4665 (65.3%)	

89 Partial 3β -hydroxysteroid Dehydrogenase (3β -HSD) and 21-hydroxylase Deficiencies in a Family With Congenital Adrenal Hyperplasia, With Evidence for Increasing 3β-HSD Activities with Age. FREDERIC M. Kenny, John W. Reynolds and Orville C. Green, Univ. of Pittsburgh, Univ. of Minnesota and Northwestern Univ., Depts. of Ped.

In this family, two affected boys have perineal hypospadias and bifid scrotum, and two affected girls have slight clitoral enlargement with otherwise normal gen-italia. All are mild 'salt-losers' with spontaneous crises occurring late (3 months and 2 years) in the boys. The girls had negative Na+ balance only when stressed by salt deprivation at ages 2 months and 4 years. All had elevated 17-ketosteroid excretion when diagnosed and, in the 2 youngest, urinary DHA > androsterone. Cortisol production and/or 17-OH-corticosteroid excretions were normal. Steroid excretion patterns showed an increase of 3β -HSD activity with increased age, but with a persisting high excretion of △5-pregnanetriol (Δ^5 -p'triol). At 2 months, one girl excreted per 24 h: pregnanetriol (p'triol)-0.23 mg, △5-p'triol-1.2 mg, $17\alpha\text{-OH-pregnenolone-2.6}$ mg, $16\alpha\text{-OH-pregnenolone-8.2}$ mg, $16\alpha\text{-OH-DHA-5.4}$ mg, DHA-0.5 mg. Her sister, at age 10 years, during withdrawal of cortisone therapy, excreted per 24 h: p'triol-24 mg, Δ^{5} -p'triol-14 mg, DHA-1.2 mg, no 16α -OH-pregnenolone or 16α -OH-DHA. We conclude that these cases have partial 3β -HSD deficiencies, on the basis of the inadequate fetal virilization and persistent post-natal excretion of large amounts of Δ^5 -p'triol; and partial 21-hydroxylase deficiencies, on the basis of the high p'triol excretions. The low post-natal excretion of DHA, relative to Δ^5 -p'triol and 17α -OH-pregnenolone, may be due to a late fetal appearance of the 3β -HSD for DHA or may be due to an underactivity of the 17-20 desmolase (side-chain splitting) enzyme.

Sulfonylureas Amplify Cyclic 3',5' AMP Mediated Hormone Action. HANS H. BODE, BARBARA M. HAR-LEY, ALLEN M. SPIEGEL and JOHN D. CRAWFORD, Harvard Med. Sch., Massachusetts Gen. Hosp., Children's Service and Shriners Burns Inst., Boston, Mass.

The reports that the sulfonylurea compounds have a vasopressin-like action in some patients with diabetes insipidus (DI) and Mahoney and Goodman's observations of benefit in one patient with hypodipsia led us to a trial in three children with both deficiencies. Hitherto extraordinarily difficult management problems, these children have shown restoration of homeostatically appropriate thirst and facultative urinary concentrating ability during chlorpropamide (CPM) treatment.

Observations to provide insight into the mechanism of these clinical effects were as follows: much as in total diabetes mellitus, the sulfonylureas were without benefit in total vasopressin lack (Brattleboro rats) and in clinical nephrogenic DI Sulfonylureas, while inactive alone, augmented vasopressin stimulated water flux across toad bladder. Water flux induced by cyclic 3',5' AMP was not increased suggesting the compounds act neither by augmenting the influence of second messenger nor by phosphodiesterase inhibition. CPM treatment increased and prolonged parathyroid hormone provoked urinary excretion of cyclic 3',5' AMP in two patients with pseudohypoparathyroidism much as it extended and intensified antidiuresis due to exogenous vasopressin in water loaded normals.

The observations support the hypothesis that sulfonylureas act by amplifying the cyclic 3',5' AMP signal generated by hormones at their specific target tissues, possibly including CNS, where second messenger concentrations are high. If correct, the amplifier hypothesis implies extended applicability of sulfonylueras and their congeners in restoring homeostasis in states of

attenuated endocrine secretion.

Adverse Effects of Large Doses of Medroxyprogeste-rone Acetate (MPA) in Idiopathic Isosexual Precocity. Robert A.Richman, Louis E. Underwood, Frank S. French and Judson J. Van Wyk, Univ. of North Carolina Sch. of Med., Dept. of Ped., Chapel Hill.

Three girls and one boy with isosexual precocity were treated with large intramuscular doses of MPA, 200-300 mg q. 7-10 d., in an attempt to control the rapid advancement of skeletal maturation. Although secondary sexual development regressed, all four developed signs suggestive of corticoid excess. These included rapid weight gain, mild hypertension, plethoric