xxii Abstracts

3. Diabetes

Urinary total estrogens were proven to be a useful tool in managing diabetic pregnancies. However, in cases of minor decreases more useful information was obtained from serum unconjugated estriol levels. The other parameters were found to be without diagnostic value.

4. Intrauterine death

In all cases of fetal death the event was preceded by a sharp decrease of serum unconjugated estriol. The indication of fetal death by the other parameters was less stringent.

Conclusion: Of the steroids (and proteins) investigated, unconjugated serum estriol values proved to be the biochemical parameter most closely correlated to "fetal well-being".

45. In vitro and in vivo adrenal cortical steroid production by foetal sheep – effect of angiotensin II, sodium deficiency, ACTH., E. M. WINTOUR, E. H. BROWN, K. J. HARDY, J. G. McDOUGALL, C. J. ODDIE and G. T. WHIPP, Howard Florey Institute of Experimental Physiology and Medicine, Dept. Physiology, Dept. Surgery, University of Melbourne, Victoria, Australia

In 1974 we reported that the ovine foetal adrenal cortex was capable of secreting aldosterone, corticosterone, and cortisol as early as the 40th day of a 145-150 day gestation period. ACTH was shown to be a potent stimulus to all three steroid secretions from adrenals incubated in vitro, particularly in the <90 day old animals. The experiments have been further extended by studying the effects of angiotensin II, and sodium deficiency on in vitro steroid production, and by the study of ACTH infused into chronically-catheterized foetuses 100-150 days. ACTH infused into chronicallycatheterized foetuses (5 I.U/h for 90 min) produced an approximate doubling of peripheral blood aldosterone, corticosterone, and 11-deoxycorticosterone concentrations, with no change in 11-deoxycortisol concentrations. From 110 days → term, control cortisol values increased from $0.05 \rightarrow 0.5 \,\mu\text{g}/100 \,\text{ml}$, and acute ACTH infusion induced 3-10-fold elevations on this baseline. Forty-one pregnant ewes provided foetuses for the angiotensin II study. Angiotensin II (2.5 µg/ml) added to the incubation medium increased the production rates of aldosterone $(1\frac{1}{2}-2\text{-fold})$, corticosterone (2-9-fold) and cortisol (2-8-fold) from adrenals of foetuses up to 100 days gestation. After 120 days angiotensin II was not a significant stimulus to steroid production in vitro. When adrenals of foetuses, 125-127 days gestation were incubated in low sodium (130 mol/l) buffer aldosterone production was not increased. 8 ewes were made severely sodium deficient by uncompensated loss of parotid saliva for 10 days. The adrenals of their foetuses, when incubated in vitro, did not produce substantially increased quantities of aldosterone. Despite demonstrated steroidogenic capacity foetal adrenal cortical cells younger than 80 days contained insigificant amounts of agranular endoplasmic reticulum.

J. Steroids in late pregnancy, ARNOLD KLOPPER, University of Aberdeen, Scotland

Although the foeto-placental unit produces a great array of steroids, clinical interest lies mainly with progestagens and oestrogens. This review will be confined to these two groups, — their precursors, the active hormones and their metabolites. Urinary steroid assays have been done for many years, plasma measurements are new; attention will be directed to plasma assays. The concentration of a steroid in blood is a different concept from urinary

steroid excretion. These differences will be examined and models for the control of steroid hormone concentration proposed.

Data concerning the range of steroid concentration in normal subjects in late pregnancy will be produced. These show that plasma concentration, as with urinary excretion may vary greatly from one healthy woman to another. So large is the normal range that there is a considerable overlap with the values found in a variety of obstetric diseases. It will be demonstrated that steroid assays have little diagnostic value; they cannot be used to diagnose the presence of retarded foetal growth or other obstetric complication. In this event the main clinical application for steroid assay is to delimit changes of steroid concentration with time in the same subject. Day-to-day variability of steroid concentration in the same subject and the factors which may affect this, becomes the central criterion in the application of hormone assays and it is intended to present some evidence concerning the time-to-time variability of plasma steroid concentration.

The use of plasma hormone estimations is based on the assumption that the maternal plasma concentration of a steroid reflects its rate of production by the foetoplacental unit, i.e., the activity of a variety of biosynthetic enzymes in the foetus and placenta. Evidence will be presented concerning the activity of such placental enzyme systems as 3β -hydroxysteroid dehydrogenase and ring A aromatase when precursors such as dehydroepiandrosterone sulphate or pregnenolone sulphate, are injected intravenously into the mother in late pregnancy.

Accurate information concerning the range of steroid concentration at various stages of pregnancy is an essential prerequisite to the application of steroid assays in the assessment of foeto-placental function. The normal levels of a variety of steroids will be reviewed and the order of change in obstetric pathology demonstrated. The changes with time in the same patient will be explored and an attempt made to correlate these with changes in the pathological state and the outcome of the pregnancy.

The evidence to be presented will tend to show that none of the steroid assays presently in use are wholly satisfactory and some speculations will be offered concerning particular steroid assays with a larger potential.

46. Identification and measurement of three oestetrols and two oestriolones in late pregnancy urine, N. F. TAYLOR and C. H. L. SHACKLETON, Division of Clinical Chemistry, Clinical Research Centre, Harrow, Middlesex HA1 3UJ, England

Two oestriolones have been identified by gas chromatography-mass spectrometry in extracts of late pregnancy urine. Sodium borohydride reduction of these steroids gave compounds with mass spectra identical to 15- and 18-hydroxy-oestriol respectively. It was concluded that they had the structures 3,15\xi216\xi225-trihydroxy-3,16\(\xi\$,18-trihydroxyoestratrien-17-one and oestratrien-17-one and therefore might be intermediates in the placental conversion of foetal 3β , 16α or β , 18-trihydroxy-5-androsten-17-one and β , 16 α -trihydroxy-5-androsten-17-one to the oestetrols which occur in pregnancy urine (Taylor N. F. and Shackleton C. H. L.: Steroids 24 (1974) 185). In order to assess the importance of these new oestrogens they have been quantified in extracts of late pregnancy urine. Steroids were recovered from urine by enzymic hydrolysis, Amberlite XAD-2 extraction and Sephadex LH-20 chromatography (Taylor N. F. and Shackleton C. H. L.: J. Endocr. 64 (1975) 8P). Following derivatization the oestetrols and oestriolones were quantified by gas chromatography and mass fragmentography. See Table.