

sites of oxidase. Thiouracil derivatives not only block iodination but also depress the formation of thyroxine from diiodotyrosine *in vivo* and *in vitro*. Drugs of this type interfere in some way with thyroxinogenesis and a new iodoprotein appears after a long treatment with propylthiouracil. The relative potency of antithyroid drugs was established by *in vivo* and *in vitro* methods. Relation between chemical structure and antithyroid action was studied. Derivatives of thiourea, mercaptoglyoxaline, thioglyoxaline and mercaptotriazole possess antithyroid properties. The mode of action of these drugs seems to be related to a blockade of oxidizing enzymes involved in organification of iodine. Reducing agents stimulate cathepsins and accelerate the rate of liberation of hormones; thyroxine inhibits deiodination of iodotyrosines preventing uptake of thyroidal iodide.

As iodine metabolism depends of TSH secretion, any compound acting directly or indirectly on the stimulation of TSH produces an effect on hormonogenesis. That is the case for thyroxine, 3:5:3'-triiodothyronine and also for the non-calorigenic 3:3':5'-triiodothyronine which are concentrated by the pituitary and block TSH secretion. Adrenaline decreases ¹³¹I uptake and produces thyroid hyperplasia.

Compounds such as 3:5-diiodo-4-hydroxybenzoylbutylester affect the rate of metabolism of iodohormones; the resulting changes in PBI level modify TSH secretion. Cortisone works by another way, it exaggerates excretion of iodide by kidneys; circulating iodide is reduced and then ¹³¹I thyroid uptake decreases.

Thus many chemical products active on peripheral iodine metabolism interfere with biosynthesis and release of thyroid hormones.

10a Action de Quelques Dérivés du 2-Amino-méthyl-Benzodioxan sur l'Équilibre Hormonal du Cycle, de la Pseudo Grossesse et de la Grossesse chez la Ratte. F. BOVET-NITTI et G. BIGNAMI (Italy).

Des recherches réalisées sur l'effet de différents dérivés sur la grossesse chez la ratte ont montré que le traitement au cours des premiers jours de la gestation peut provoquer des accidents dont les manifestations n'apparaissent parfois que d'une manière tardive.⁽¹⁾ Dans le cas d'un sympatholytique, le Piperoxan (933F.), l'on a observé que 100 mg/kg administrés par voie orale le jour 1 de la grossesse empêchent l'ovoimplantation chez 30 à 50 pour cent des rattes sans provoquer d'anomalies chez les autres animaux du lot. L'étude de dérivés voisins montre que cette action ne paraît pas liée aux propriétés sympatholytiques des ces produits.

Une activité sur la gestation a été également reconnue à certains dérivés du benzodioxan à fonction hydrazine, inhibiteurs des monoamino oxydases (MAO). Dans le cas du N'-1:4-benzo-

dioxan-2-méthyl-hydrazine (2596IS)⁽²⁾ l'administration de 100 mg/kg par voie orale le jour 1 de la grossesse provoque chez 50 pour cent des rattes soit une inhibition partielle ou totale de l'implantation soit, exceptionnellement, la résorption précoce d'une partie des nidations.

Diverses considérations font penser que l'effet du Piperoxan et du 2596IS est d'origine centrale et que ces substances sont susceptibles de modifier l'équilibre hormonal par l'intermédiaire des sécrétions de l'antéhypophyse. Il a été en effet remarqué⁽³⁾ que le 2596IS s'oppose dans le 100 pour cent des cas à la pseudogrossesse provoquée par la Réserpine et dans une proportion pouvant aller jusqu'à 40 pour cent à celle dérivant d'une stimulation mécanique.

1. BOVET-NITTI, F. and BOVET, D. (1959), *Proc. exp. Biol. Med.*, **100**, 555.
2. BOVET-NITTI, F., ORSINGER, O., LAND-VITTORE, R. and BOVET, D. (1961), *Comptes Rendus Acad. Scand.*, **252**, 614.
3. BIGNAMI, G., *Rendiconti Ist. Sup. Sanità.* In press.

10b Action of Certain Derivatives of 2-Amino Methyl-Benzodioxan on the Hormonal Equilibrium of the Reproductive Cycle, Pseudopregnancy and Pregnancy in the Rat. F. BOVET-NITTI and G. BIGNAMI (Italy).

Investigations on the effect of various derivatives on pregnancy in the rat have shown that treatment during the first days of the gestation period can cause accidents which sometimes only become apparent later.⁽¹⁾ In the case of the sympatholytic, Piperoxan (933F) it has been observed that 100 mg/kg administered orally on the first day of pregnancy prevent the implantation of the ovum in 30 to 50 per cent of the rats without causing any abnormalities in the other animals of the group. The study of related derivatives shows that this action does not appear to be related to the sympatholytic properties of these drugs.

Certain derivatives of benzodioxan with hydrazine properties, which inhibit monoamino oxydase (MAO), have also been found to affect gestation. In the case of N'-1:4-benzodioxan-2-methyl-hydrazine (2596IS) the administration of 100 mg/kg orally on the first day of pregnancy provokes either partial or total inhibition of implantation in 60 per cent of the rats or, in exceptional cases, the precocious absorption of the implant.

Various considerations lead to the hypothesis that the effect of Piperoxan and of 2596IS is of central origin and that these substances can modify the hormonal equilibrium through the intermediacy of the anterior hypophyseal secretions. It has, in fact, been observed⁽³⁾ that 2596IS counteracts the pseudopregnancies due to Reserpine in 100 per cent of the cases and in up to 40 per cent in cases where it is due to mechanical stimulation.