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MALE REPRODUCTIVE HEALTH, CHEMICALS AND ENVIRONMENTAL FACTORS, A MEDITERRANEAN WORKSHOP

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Organizing Committee : Bernard JEGOU (Rennes) Pierre JOUANNET (Paris) Alfred SPIRA (Bicêtre)

Sponsors:

Direction Générale de la Santé (DGS), Secrétariat d'Etat à la Santé International Program on Chemical Safety (IPCS) European Environmental Agency (EEA)

INTERACTION OF XENOBIOTICS WITH SEX HORMONE BINDING GLOBULIN

Florence Le Gac, Martine Ollitrault.

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Sex steroid hormones circulate in the blood and extracellular compartments mainly bound to albumin and specific Steroid Binding Proteins (SBP). Xenobiotics may interfere with this endocrine equilibrium by occupying the binding sites of the transport protein modify the ratio of bound/free endogenous testosterone (T) and estradiol (E2) and therefore the biological activity or the metabolic clearance of these natural hormones (Aldercreutz et al. 1985, 1990). Moreover, binding to SBP may influence the bioavailability of the xenohormone itself: it may protect the exogenous molecules against metabolism and promote their accumulation inside the organism, while the absence of binding make them more available for action on their target cells.

We have set up a sensitive and practical in vitro screening method to detect chemicals that modify SBP-sex steroid binding. We incubate increasing concentrations of xenobiotics (10⁻⁷ to 10⁻⁴ M) in competition with 10⁻⁹ M of labelled testosterone or estradiol (-3H-T or 3H-E2), for binding to fish or human blood plasma SBP in (Fig.1). vitro

30 active pesticides used in French agriculture were screened (exemples in table 1): four of them had significant affinity for SBP with ED50 $= 0.5 - 5 \times 10^{-6} M$ (Parathion-methyl > Bifenox > Dodemorphe = Triadimefon, in decreasing

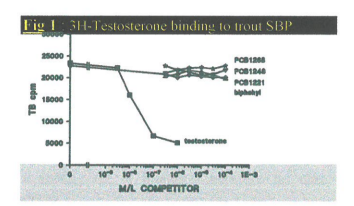
order of activities) and should be concidered as

potential endocrine disruptors.

Among estrogen-like compounds, phytoestrogens equal and, to a lesser extent, genistein competed with E2 and T for binding to SBP, as well as Nonylphenol4 and its ethoxyl derivatives (ED50: 2 à 10 10⁻⁶ M). None of the polychlorinated derivatives of biphenyl (PCBs) tested revealed activity.

anabolic molecules used in Most of the farming, or their metabolites, are able to bind to vertebrate SBP and to inhibit natural hormone binding to their specific blood transport proteins. Exposition of either human or wild vertebrates to one of these compounds could therefore consequences have estrogens/free androgens equilibrium or disrupt endogenous steroid hormone action on target tissues, especially if the xenobiotic compound considered is susceptible to bioaccumulate in the organism.

The case of **Zeranol** and its metabolites is of special interest as their very high estrogenic activity is not "buffered" by binding to an high affinity protein in the extracellular compartment (table2): this could make it more available for action on target cells (as proposed for the action of Diethylstilbestrol on the human foetus).



Chemicals	INHIBITION	OF		
	Charles and the second	BINDING TO		
	% inh.			
	at 10 -5 M	ED 50 M/L		
Biphenyl	0			
Captan	0			
Chlorothalon	30%	3 x 10 E-5		
Dodemorph	60%	3 x1 0 E -6		
Pentachlorophen	50%	2 x 10 E -5		
Prochloraz	50%	2 x 10 E -5		
Triadimefo	85%	3 x 10 E -6		
Carbofura				
Carbosulfan	20%	> 10 E -4		
Deltaméthrin	0			
Lindan	50%	2 x 10 E -5		
Parathion-	100%	5 x 10 E -7		

Chemical	Inhibition T-SBP binding	E2 Recept. Activation at 10 ⁷ M
Ethynyl-estradiol Diethylstilbestrol	100 0	137% 100 %
Zearalenone	0	108 %
α Zearalanol	0	146 %
β Zearalanol	20	62 %
α zearalenol	100	20 %
B zearalenol	15	134 %