[S-6]

Effects of Pahs and Pcbs and Their Toxic Metabolites on Inhibition of Gjic and Cell Proliferation in Rat Liver Epithelial Wb-F344 Cells

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The liver progenitor cells could form a potential target cell population fore both tumor-initiating and -promoting chemicals. Induction of drug-metabolizing and antioxidant enzymes, including AhR-dependent CYP1A1, NQO-1 and AKR1C9, was detected in the rat liver epithelial WB-F344 "stem-like" cells. Additionally, WB-F344 cells express a functional, wild-type form of p53 protein, a biomarker of genotoxic events, and connexin 43, a basic structural unit of gap junctions forming an important type of intercellular communication.

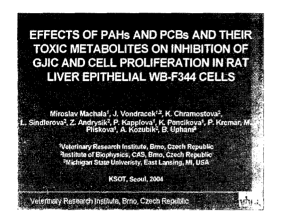
In this cellular model, two complementary assays have been established for detection of the modes of action associated with tumor promotion: inhibition of gap junctional intercellular communication (GJIC) and proliferative activity in confluent cells. We found that the PAHs and PCBs, which are AhR agonists, released WB-F344 cells from contact inhibition, increasing both DNA synthesis and cell numbers. Genotoxic effects of some PAHs that lead to apoptosis and cell cycle delay might interfere with the proliferative activity of PAHs. Contrary to that, the nongenotoxic low-molecular-weight PAHs and non-dioxin-like PCB congeners, abundant in the environment, did not significantly affect cell cycle and cell proliferation; however both groups of compounds inhibited GJIC in WB-F344 cells.

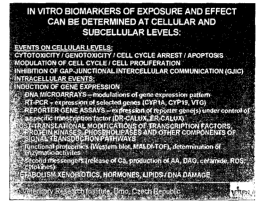
The release from contact inhibiton by a mechanism that possibly involves the AhR activation, inhibition of GJIC and genotoxic events induced by environmental contaminants are three important modes of action that could play an important role in carcinogenic effects

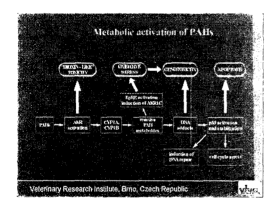
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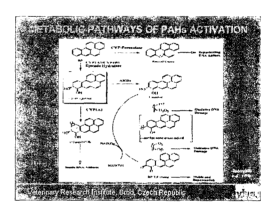
of toxic compounds. The relative potencies to inhibit GJIC, to induce AhR-mediated activity, and to release cells from contact inhibition were determined for a large series of PAHs and PCBs and their metabolites. In vitro bioassays based on detection of events on cellular level (deregulation of GJIC and/or proliferation) or determination of receptor-mediated activities in both "stem-like" and hepatocyte-like liver cellular models are valuable tools for detection of modes of action of polyaromatic hydrocarbons. They may serve, together with concentration data, as a first step in their risk assessment.

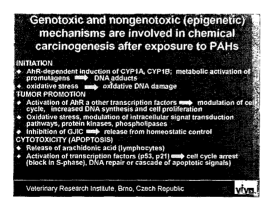
This work was supported by the Czech Ministry of Agriculture grant No. MZE00002716201.









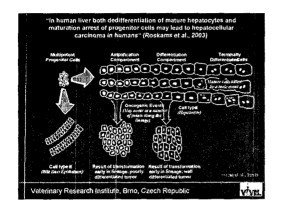


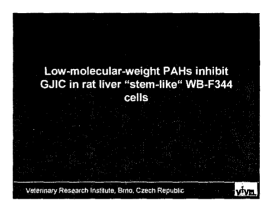
Two cell types used in the study:

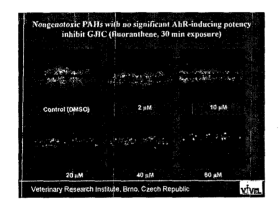
1) poorly differentiated "stem-like" (progenitor) cells (model: non-transformed liver epithelial WB-F344 cells)

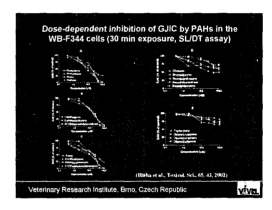
2) primary hepatocytes / hepatoma cell lines (e.g., rat hepatoma H4IIE cells)

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INHIBITION OF GJIC: CONCLUSIONS

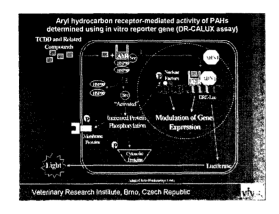
Low-molecular-weight PAHs incl. fluoranthene, pyrene, and phenanthrene inhibited significantly GJIC; most of high-molecular-weight PAHs with known strong carcinogenic properties possessed only weak (DBPyrenes) or no inhibition potency (DBakF, DBajF, N[23a]P).

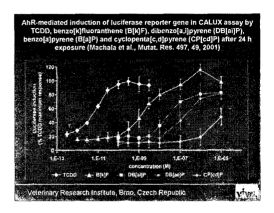
Using of selective inhibitors of protein kinases suggested mechanism of inhibition of GJIC after exposure to PAHs different from effects of EGF or TPA; possible roles of src kinase, PC-PLC and DAGlipase in this process.

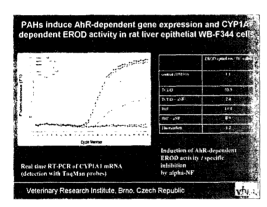
Receptor tyrosine kinases EGFR, ErbB-2, and NGFR, as well as mitogen-activated protein kinases ERK1/2, p38, PKC, Akt, phospholipases PI-PLC and PLA2 are probably not involved in the GJIC inhibition by PAHs.

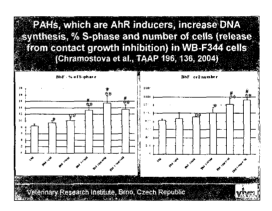
PAHs induce AhR-dependent
("dioxin-like") activity in hepatoma and
non-transformed liver epithelial cells

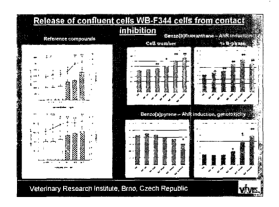
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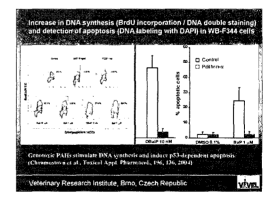


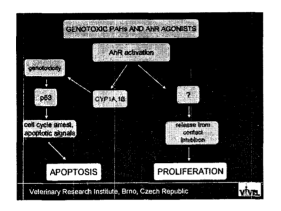


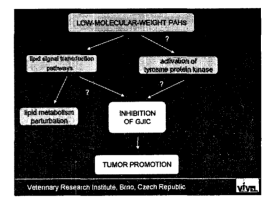


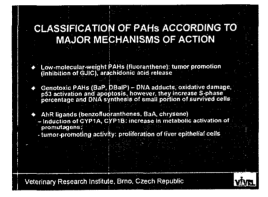




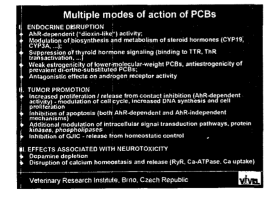


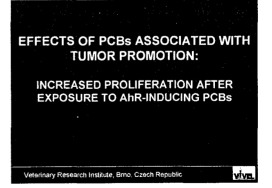


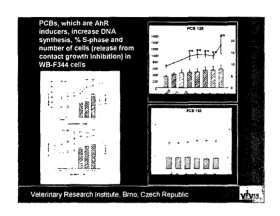


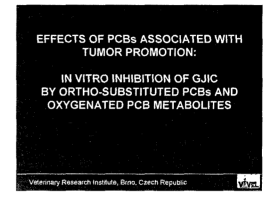


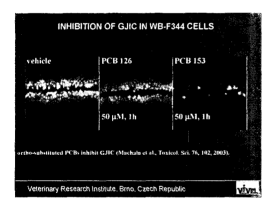
Classification of PCB congeners Coplanar PCBs Mono-ortho-substituted PCBs (156> 114,118> 189 > 74 >> 105...) Prevalent di-ortho-substituted PCBs (138,153,170,180 > 187 >>...) Episodic PCBs (18,49,149,...) OH-PCBs, MeSO₂-PCBs Veterinary Research Institute, Brno. Czech Republic

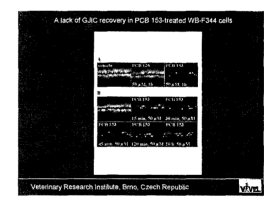


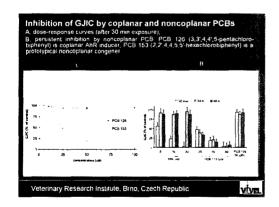


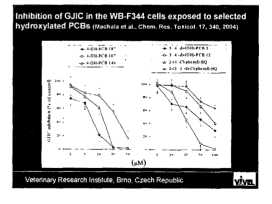


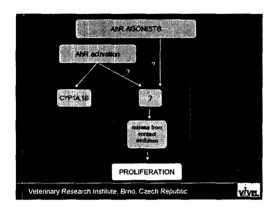


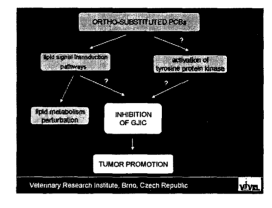












Di-ortho-substituted (noncoplanar), mono-ortho-chlorinated PCBs and OH-PCBs acted as potent <u>inhibitors of GJIC</u> in micromolar concentrations. Coplanar PCBs (strong inducers of aryl hydrocarbon receptor) possessed no acute inhibitory potency. Coplanar and mono-ortho-substituted PCB congeners increased % S-phase and total numbers of liver epithelial cells (<u>proliferative activity</u>).

