Fyziologie působení farmak a toxických látek

Přednáška č.5 Proteiny PAS a jejich úloha v organismu

Per-Arnt-Sim - PAS superfamily of proteins

environmental sensors, which mediate transcriptional responses to various types stimuli:

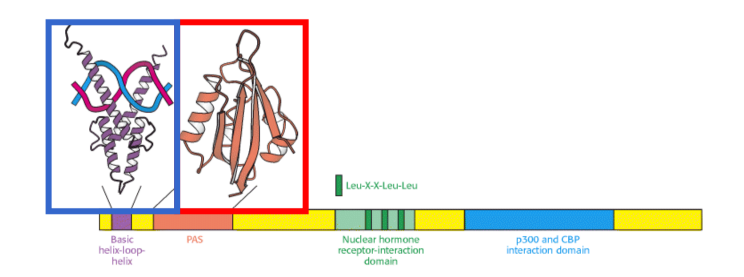
- circadian rhythms;
- oxygen sensing;sensing of toxicants;
- developmental role/cancer;

These proteins enable adaptation to rapid changes in the environment.

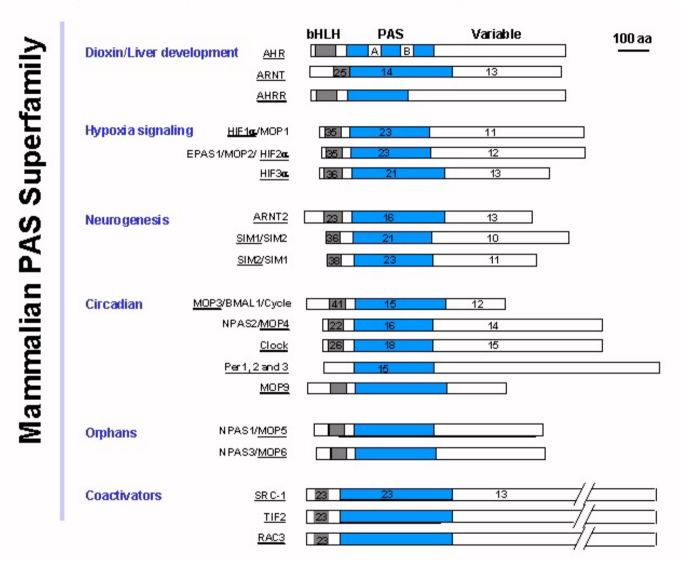
PAS proteiny jsou součástí širší rodiny bHLH proteinů:

There are three main sub-families of bHLH proteins:

- (a) those containing only the bHLH domain; and those where the bHLH domain is contiguous with a second dimerisation domain, either
- (b) the leucine zipper (Zip) or
- (c) the PER/aryl hydrocarbon receptor nuclear translocator (ARNT)/single minded (SIM) (PAS) homology domain.



PAS proteiny (rodina transkripčních faktorů):



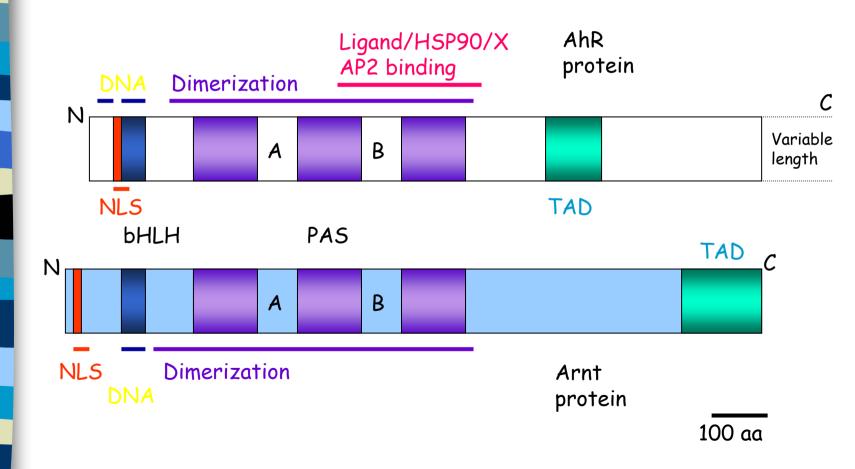
PAS domain

The PAS region consists of two adjacent degenerate repeats of ~130 amino acids, PAS A and PAS B. The domain is an ancient signalling device conserved through evolution, having been identified in proteins throughout the animal kingdom, in bacteria, fungi and yeast in addition to mammals and flies, where the most commonly studied bHLH/PAS proteins originate.

Many bacteria contain PAS-like proteins that detect light and oxygen (Dos, Aer, FixL, PYP).

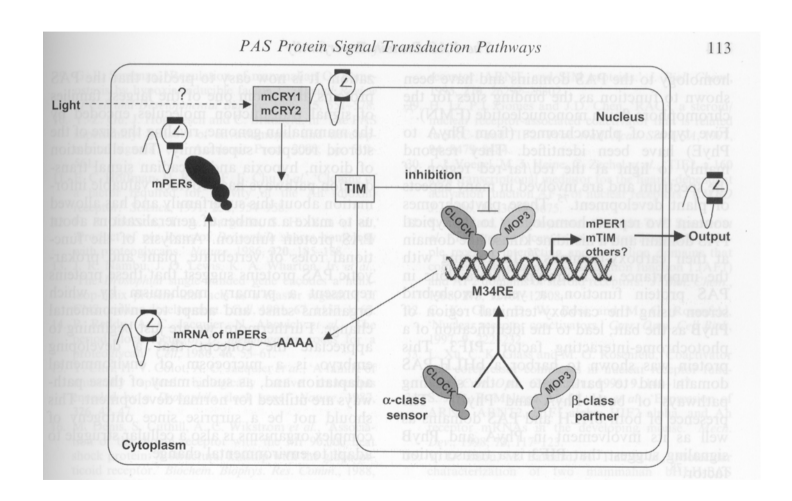
Similar proteins sense light in plants (phytochromes PhyA-PhyE, NPH1; phytochrome interacting factor PIF3).

Domain structure and function of PAS proteins:



(Gu et al., Annu Rev Pharmacol Toxicol. 2000;40:519-61.)

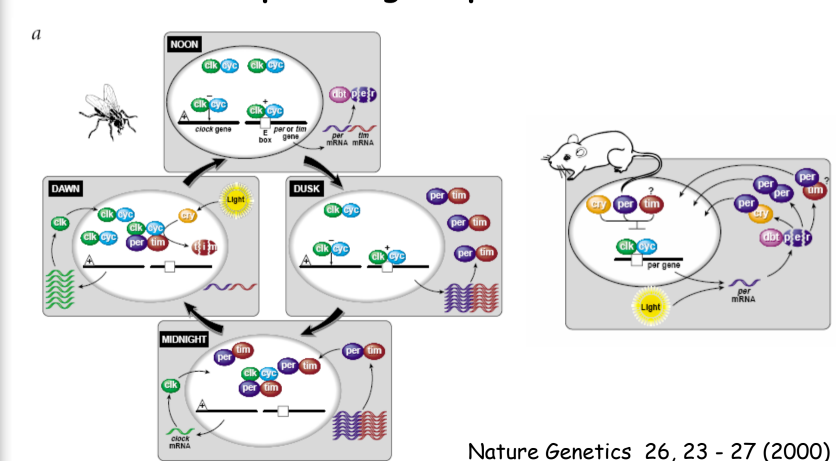
The circadian response pathway



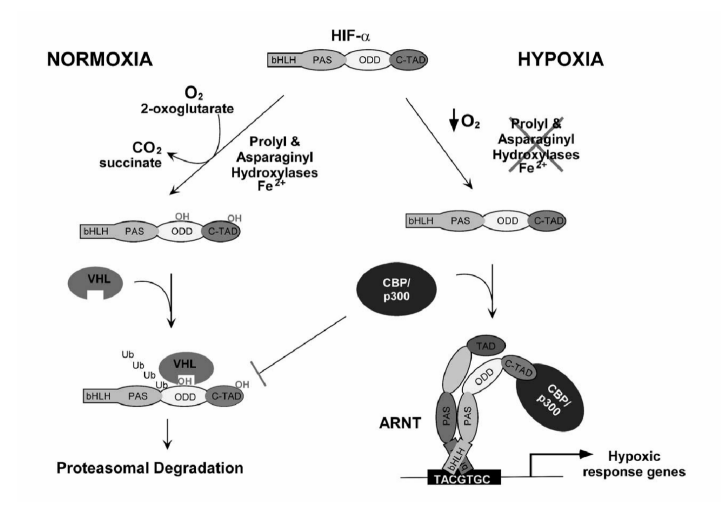
(Comprehensive Toxicology, vol. 14)

Daily changes in light/dark require physiological and behavioral adaptation:

✓ CLOCK/MOP3 heterodimer controls expression
of circadian responsive gene products - PER, TIM;



The hypoxia response pathway



The International Journal of Biochemistry & Cell Biology 36 (2004) 189-204

The ability to maintain O2 homeostasis is essential for survival of mammals. The hyperoxic state, or high O2 tension, can result in the generation of reactive

The hyperoxic state, or high O2 tension, can result in the generation of reactive oxygen intermediates and potentially lethal damage to membranes and DNA. The hypoxic state, or low O2 tension, can result in levels of ATP insufficient to maintain essential cellular functions. The hypoxic state occurs in a number of medical conditions, such as cancer and ischemias, inspiring research into understanding the cellular mechanisms for detecting and responding to low levels of oxygen. Responses to hypoxia are mediated by three bHLH/PAS proteins, HIF-1a, HIF-2a (also known as Endothelial PAS domain protein 1, HIF-like factor and member of PAS family 2), and HIF-3a.

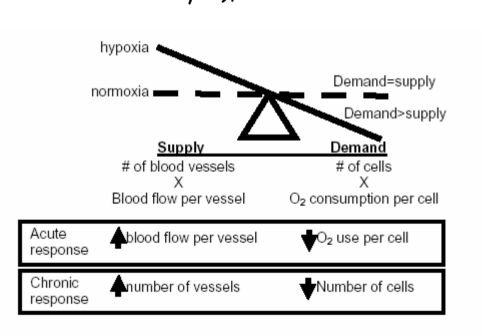
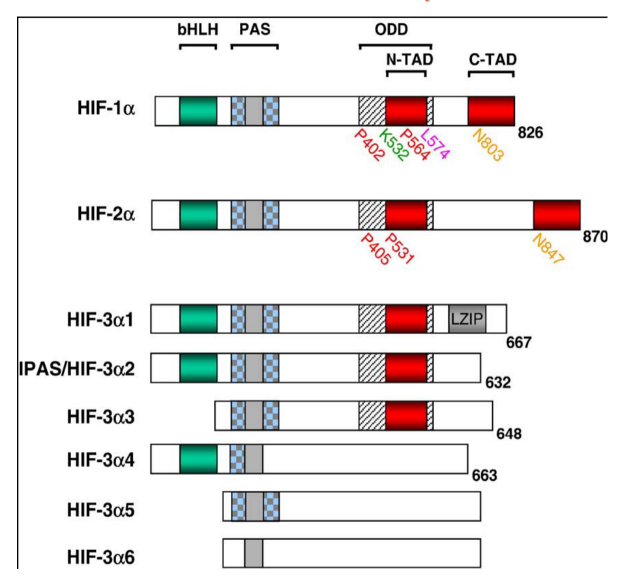


Fig. 1. Supply and demand governs oxygen availability.

HIF subfamily



Biochimica et Biophysica Acta 1755 (2005) 107 - 120

Hypoxia-inducible factor (HIF-1 α):

Hypoxia-inducible factor-1 (HIF-1), composed of HIF- α and HIF- β (ARNT) subunits, is a heterodimeric transcriptional activator. In response to hypoxia, stimulation of growth factors, and activation of oncogenes as well as carcinogens, HIF-1a is overexpressed and/or activated and targets those genes which are required for angiogenesis, metabolic adaptation to low oxygen and survival of cells. HIF-1 is critical for both physiological and pathological processes.

Several dozens of putative direct HIF-1 target genes have been identified on the basis of one or more cis-acting hypoxia-response elements that contain an HIF-1 binding site. A variety of regulators including growth factors, genetic alterations, stress activators, and some carcinogens have been documented for regulation of HIF-1 in which several signaling pathways are involved depending on the stimuli and cell types. Activation of HIF-1 in combination with activated signaling pathways and regulators is implicated in tumour progression and prognosis.

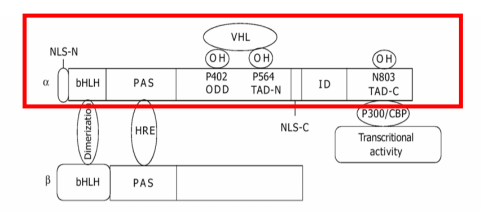


Figure 1 Molecular structure of HIF- 1α and HIF- 1β . bHLH domain mediates dimerization of the two subunits. PAS domain is responsible for DNA binding. Proline residues of 402 and 564 at ODD domain are hydroxylized by proline hydroxylase and recognized by VHL and then targeted to the ubiquitin proteasome pathway. Asn803 at the C-terminal transactivation domain (TAD-C) is hydroxylized by FIH-1 (factor inhibiting HIF-1) with a result of inhibition of HIF- 1α interaction with co-activator p300 and consequently inhibits transcriptional activity. The nuclear location signal at C-terminal functions in HIF- 1α translocation into nuclei.

World J Gastroenterol 2004;10(8):1082-108

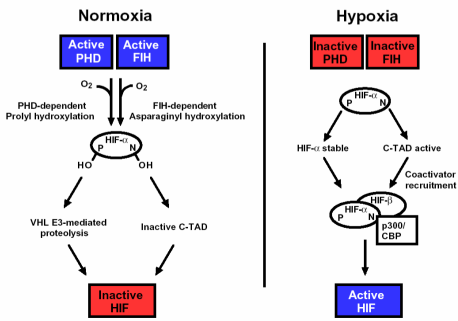


Fig. 1. Two independent hydroxylation pathways regulate HIF activity in response to cellular oxygen level. In normoxia, oxygen availability enables PHD-dependent prolyl hydroxylation of the HIF-α ODD. This prolyl hydroxylation allows binding of the VHL E3 ligase leading to ubiquitylation and degradation of HIF-α subunits. Oxygen availability also enables FIHdependent asparaginyl hydroxylation of the C-TAD, blocking interaction with the p300/CBP co-activator. In hypoxia, the PHD and FIH enzymes are inactive and the lack of hydroxylation results in stable HIF-α able to form a DNAbinding heterodimer with HIF-β and recruit p300/CBP at the C-TAD.

HIF-dependent responses to O2 may be modulated by the cellular environment:

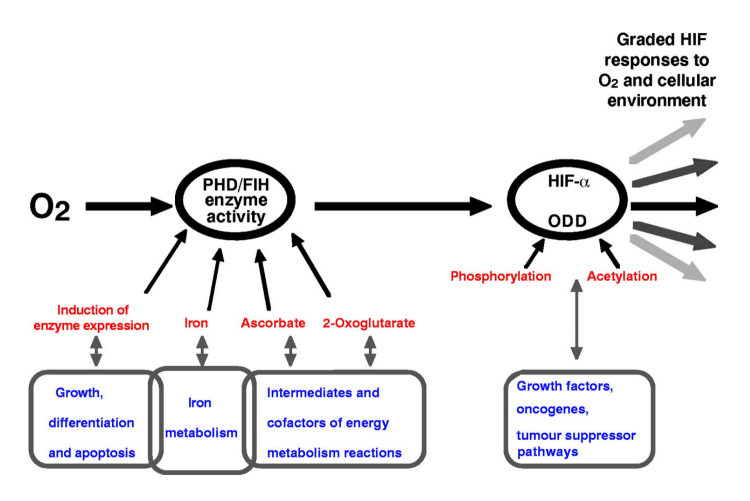


Table 2. HIF-1 target genes.

F 41	Constallarities) Before			
Function	Gene (abbreviation)	Reference		
	Erythropoietin (EPO)	(Semenza et al., 1991)		
	Transferrin (Tf)	(Rolfs et al., 1997)		
	Transferrin receptor (Tfr)	(Bianchi et al., 1999)		
	Ceruloplasmin	(Lok and Ponka, 1999)		
Angiogenesis	Vascular endothelial growth factor (VEGF)	(Levy et al., 1995)		
	Endocrine-gland-derived VEGF (EG-VEGF)	(LeCouter et al., 2001)		
	Leptin (LEP)	(Grosfeld et al., 2002)		
	Transforming growth factor-beta3 (TGF- β3)	(Scheid et al., 2002)		
Vascular tone	Nitric oxide synthase (NOS2)	(Melillo et al., 1995)		
		(Lee et al., 1997)		
	Endothelin 1 (ET1)	(Hu et al., 1998)		
	Adrenomedulin (ADM)	(Nguyen and Claycomb,		
		1999)		
	α1B-adrenergic receptor	(Eckhart et al., 1997)		
Matrix metabolism		(Ben-Yosef et al., 2002)		
	Plasminogen activator receptors and inhibitors	(Kietzmann et al., 1999)		
	(PAIs)			
	Collagen prolyl hydroxylase	(Takahashi et al., 2000)		
	Adenylate kinase-3	(O'Rourke et al., 1996)		
	Aldolase-A,C (ALDA,C)	(Semenza et al., 1996)		
	Carbonic anhydrase-9	(Wykoff et al., 2000)		
Glucose metabolism	Enolase-1 (ENO1)	(Semenza et al., 1996)		
	Glucose transporter-1,3 (GLU1,3)	(Chen et al., 2001)		
		(Graven et al., 1999)		
	(GAPDH)			
	Hexokinase 1,2 (HK1,2)	(Mathupala et al., 2001)		
	Lactate dehydrogenase-A (LDHA)	(Semenza et al., 1996)		
	Pyruvate kinase M (PKM)	(Semenza et al., 1994)		
	Phosphofructokinase L (PFKL)	(Semenza et al., 1994)		
	Phosphoglycerate kinase 1 (PGK1)	(Semenza et al., 1994)		
	6-phosphofructo-2-kinase/gructose-2,6-	(Minchenko et al., 2002)		
	bisphosphate-3 (PFKFB3)			
	Insulin-like growth factor-2 (IGF2)	(Feldser et al., 1999)		
Cell proliferation/	Transforming growth factor-a (TGF-a)	(Krishnamachary et al.,		
survival		2003)		
	Adrenomedullin (ADM)	(Cormier-Regard et al.,		
		1998)		
Apoptosis	Bcl-2/adenovirus EIB 19kD-interacting protein 3	(Carrero et al., 2000)		
	(BNip3)			
	Nip3-like protein X (NIX)	(Bruick, 2000)		

Molecular Pharmacology Fast Forward. Published on August 3, 2006 as doi:10.1124/mol.106.0 27029

ARNT - základní dimerizační partner:

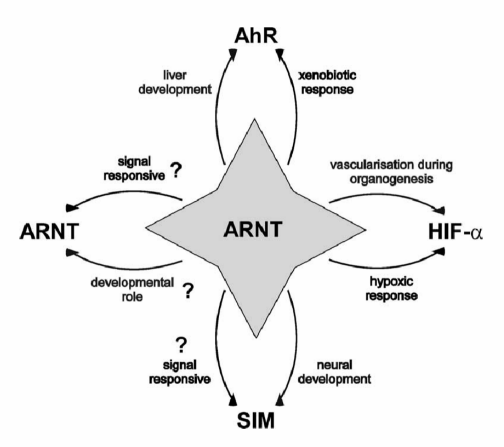
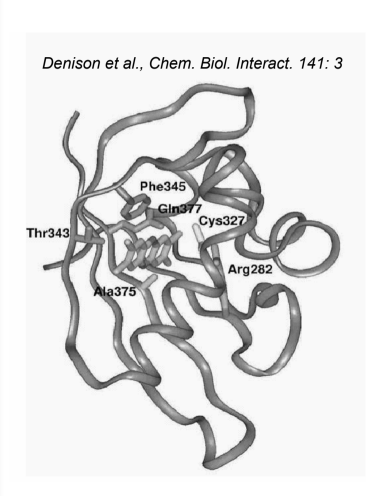


Fig. 4. ARNT is central to transcriptional regulation within the bHLH/PAS family of proteins. ARNT forms both homodimers and heterodimers with the AhR, HIF- α and SIM which play roles both during mammalian development and in response to environmental stimuli in mammals. Symbol '?' indicates where these roles have yet to be characterised.

Jak HIF-1α, tak ARNT představují proteiny nezbytné pro přežití – KO myši odumírají již v průběhu embryonálního vývoje.

The Ah receptor pathway



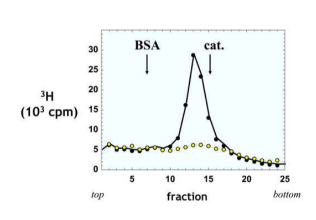


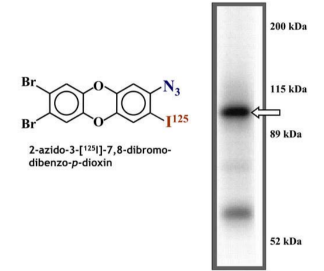
AhR =

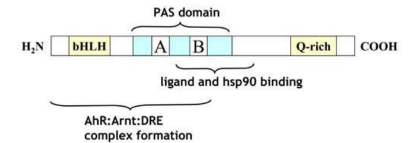
- · ligand-activated transcription factor;
- · important mediator of toxicity of POPs;
- regulator of xenobiotic metabolism and activation of promutagens.

AhR discovery

- different sensitivity of inbred mouse strains to TCDD and 3-MC inducers of CYP1A activity in liver microsomes;
- · autosomal dominant Mendelian trait;
- isolation of protein; cloning







Molecular Toxicology, 2nd ed.

Overview of aryl hydrocarbon receptor and dioxin-like toxicity:

- · what is AhR;
- evolution perspective;
- · activation of AhR; AhR-dependent genes
- · toxic effects associated with AhR activation;
- · AhR interactions
- · the role of AhR in cell cycle regulation

AhR domain structure:

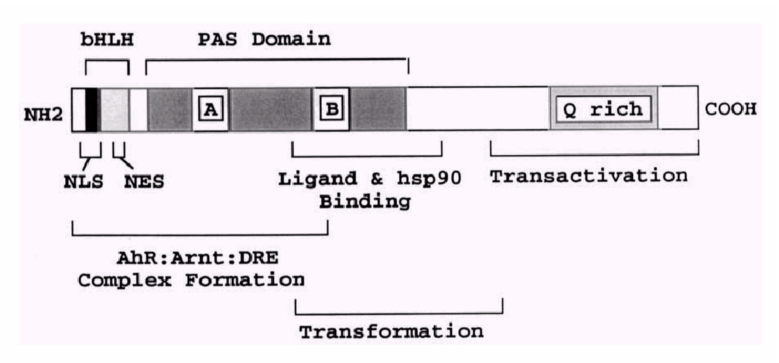
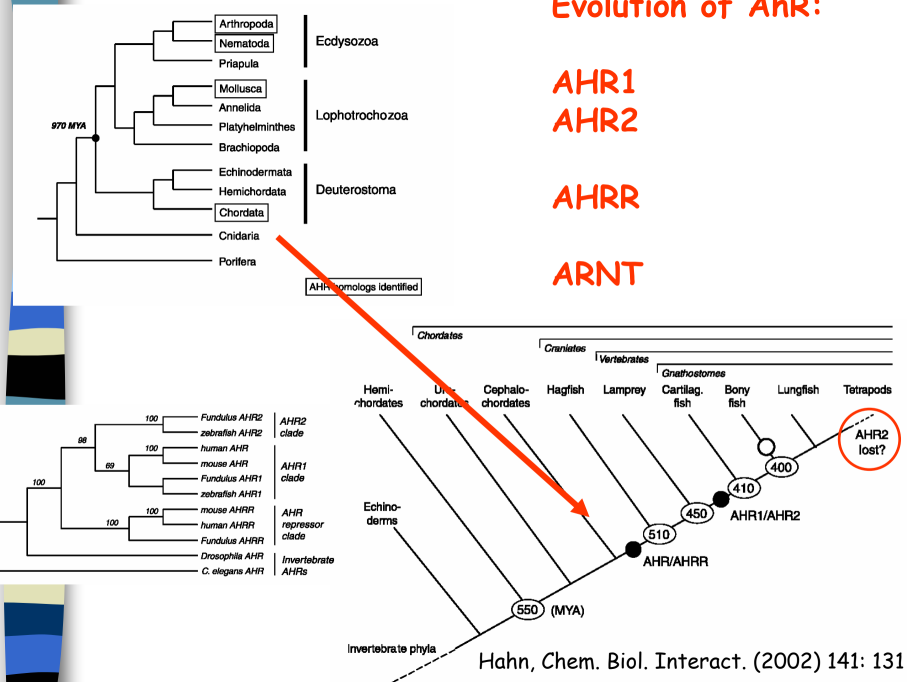


Fig. 2. Domain structure of the AhR.



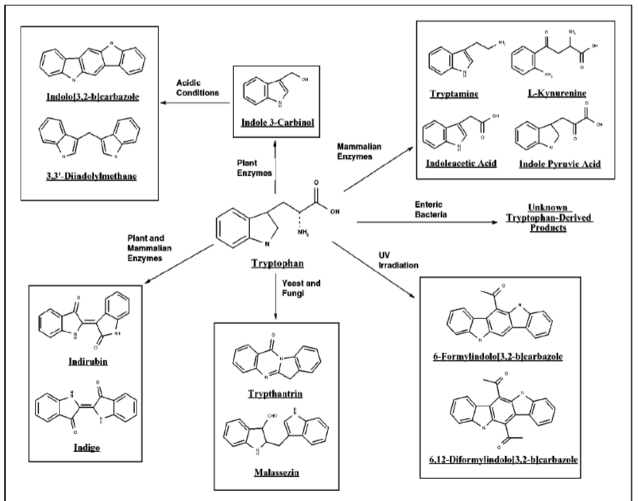
Evolution of AhR:

Organism:	Name:	Ligand-binding:	Physiological function:
Nematodes:	AHR-1	No	Neuronal development;
Caenorhabditis elegans			Behavioral effects.
Insects:	Spineless (Ss)	No	Development;
Drosophila melanogaster			Regulation of homeobox genes and dendrite morphology.
Vertebrates:	AhR	Yes	Toxicity mechanisms;
	(AhR1, AhR2)		Liver and kidney development;
			Neuronal differentiation?
			Circadian rhytms?

Natural ligands of AhR???????????



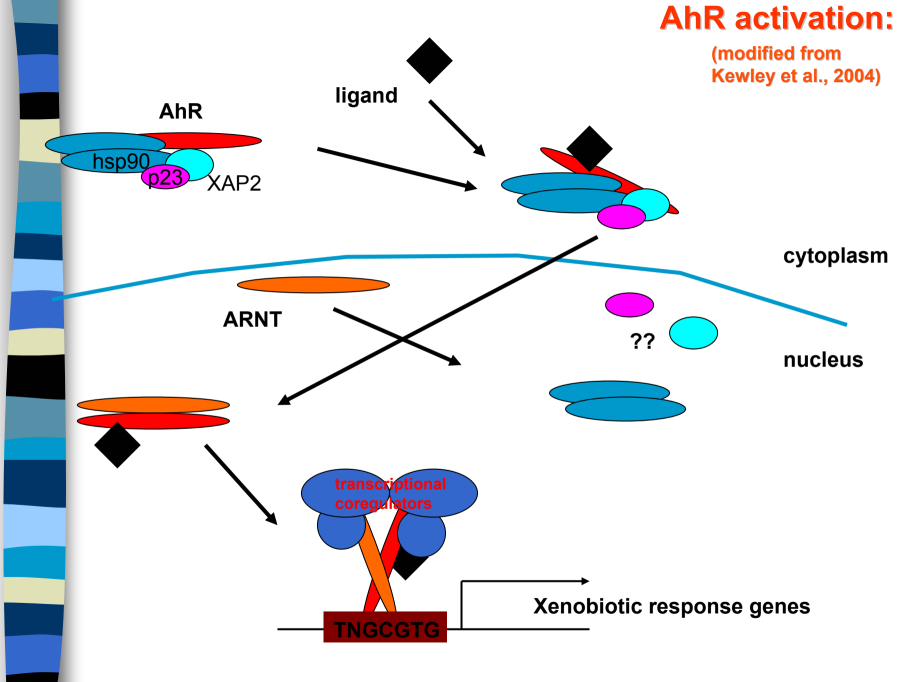
light hypothesis - tryptophane derivatives



9 Name Amour Day Dlamman J Tayling 1 42,20

Natural ligands of AhR??????????





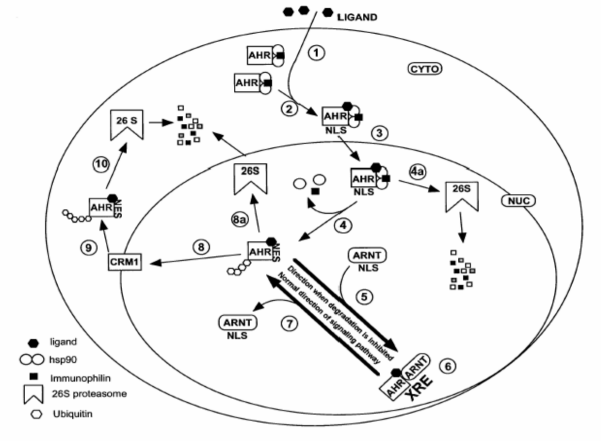
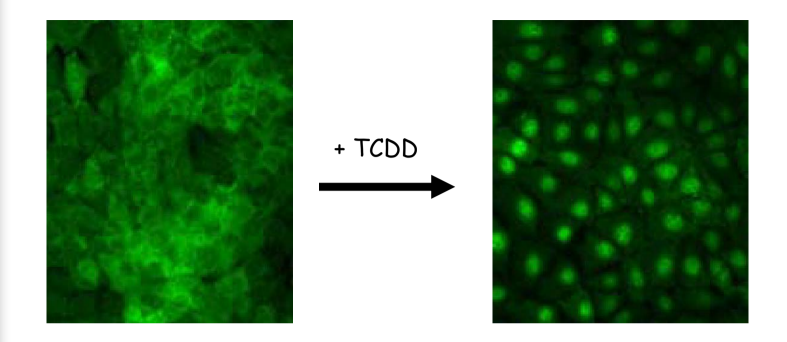
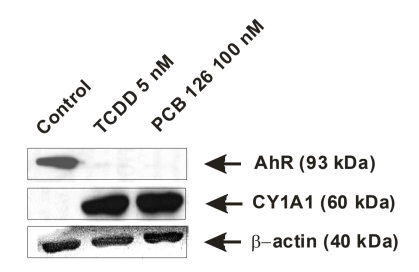


Fig. 1. Model of AHR-mediated signal transduction pathway. (1) Ligand enters cell. (2) Ligand binds to AHR-hsp90-immunophilin complex causing conformational change and exposing the NLS domain. (3) AHR complex is actively imported into the nucleus via NLS and nuclear import receptors. (4a) If receptor complex is in a misfolded conformation, it may be proteolytically degraded. (4) AHR dissociates from hsp90 and immunophilin exposing HLH/PAS domain and NES. (5) AHR dimerizes with ARNT-blocking NES sequence. (6) AHR-ARNT complex binds to XRE regions in DNA. (7) AHR-ARNT complex dissociates from DNA and ARNT exposing NES. (8a) AHR is ubiquinated in the nucleus and degraded or (8) AHR is exported from nucleus via CRM-1 export receptor. (9) AHR is ubiquinated in cytoplasm and (10) targeted to 26S proteasome for degradation. Note that the pathway is linear and also note the degradation of the AHR terminal step regardless of whether it occurs within the nucleus or cytoplasm. NLS, nuclear localization signal; CRM-1, chromosome region maintenance protein 1; 26S, 26S proteasome.



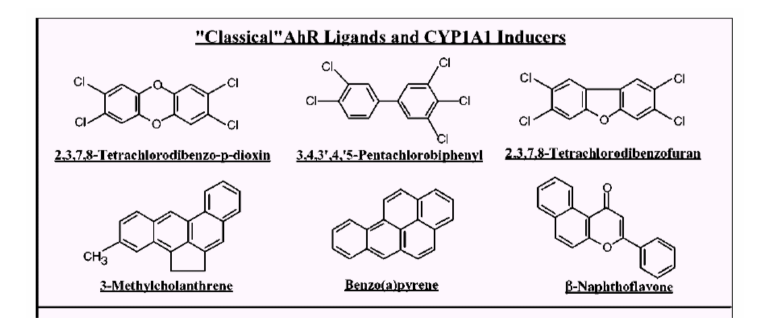


AhR regulated genes:

contain <u>xenobiotic response elements</u> (XRE) or dioxin responsive elements (DRE) in their promoter region:

- · phase I enzymes CYP 1A1, CYP 1A2, CYP 1B1;
- phase II enzymes UDP-glucuronosyltransferase, GST-Ya, NADP(H):oxidoreductase;
- · other genes Bax, p27^{kip1}, Jun B, TGF-b regulation of cell cycle and apoptosis;
- · AhRR.

AhR toxicants:



Toxic effects of dioxins:

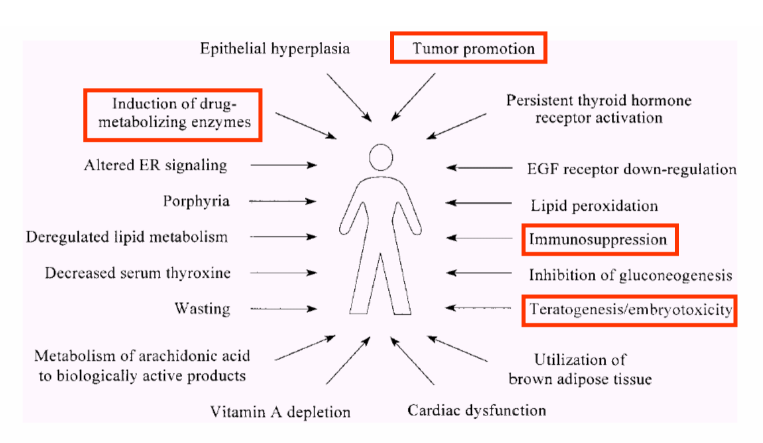


Figure 1 Biological responses to TCDD. A wide variety of cellular processes have been shown to be affected by TCDD.

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"Non-classical" AhR ligands

M.S. Denison et al. | Chemico-Biological Interactions 141 (2002) 3-24

Physiological role for AhR - AhR-deficient mice:

- ✓ significant growth retardation;
- ✓ devective development of liver and immune system;
- ✓ retinoid accumulation in liver;
- ✓ abnormal kidney and hepatic vascular structures.
- ✓ resistant to BaP-induced carcinogenesis and TCDDinduced teratogenesis;
- \checkmark no inducible expression of CYP 1A1 and 2.

Liver defects:

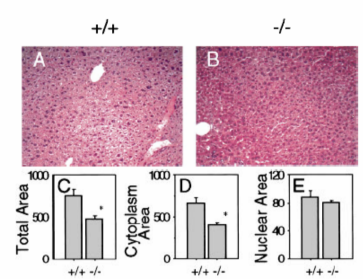
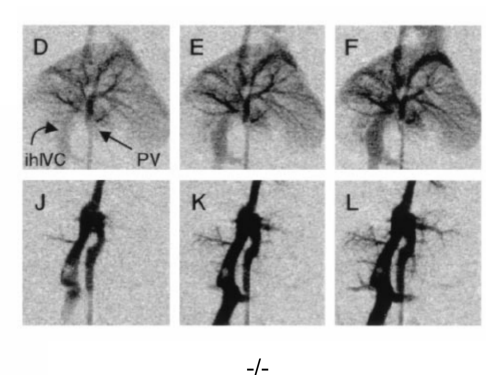


Fig. 1. Ah -/- mice have smaller hepatocytes than wild-type mice. Livers of 1-year-old mice were fixed in formalin, and $6-\mu m$ sections were examined after staining with hematoxylin/eosin. (A and B) Thin sections from wild-type (A) and age-matched Ah knockout (B) mice are shown, and results of morphometric analyses follow. (C) There is a significant decrease in the total area of the hepatocytes of Ah -/- mice. (D and D) Whereas the cytoplasmic area of D0, the nuclear areas of D1 mice and D2. Mean and standard errors generated from comparison of six 1-year-old male D3 male D4 mice are shown; asterisks indicate significance (D4 o.05).



A

O'

P

+/+
-/-

PNAS 2000 vol. 97:10447

BaP není karcinogenní v AhR KO myších:

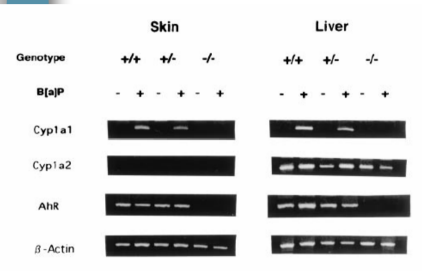


Fig. 1. *Cyp1a1*, *Cyp1a2*, and AhR gene expression in the skin and liver of AhR(+/+), AhR(+/-), and AhR(-/-) mice, with and without B[a]P treatment. One-microgram aliquots of RNA extracted from skin and liver of control and B[a]P-treated mice of the three genotypes were reverse-transcribed and analyzed by PCR using specific primers for the *Cyp1a1*, *Cyp1a2*, and AhR and β -actin genes.

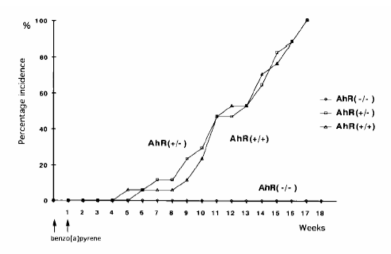


Fig. 2. Subcutaneous tumor induction in wild-type (\triangle) and AhR-deficient male mice (+/-, \Box ; -/-, \bigcirc) injected with B[a]P.

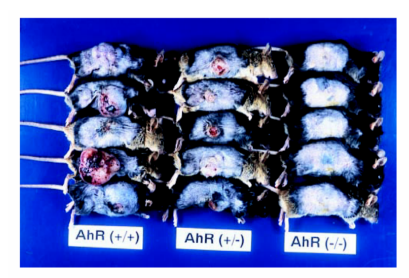
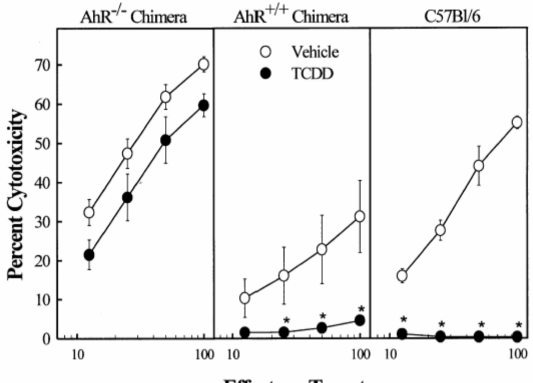


Fig. 3. Gross appearance of flank skins in AhR-wild-type mice (+/+), AhR-heterozygous mice (+/-), and AhR-deficient mice (-/-) injected subcutaneously with B[a]P.

AhR je nezbytný pro imunotoxické účinky TCDD:

N.I. Kerkvliet / International Immunopharmacology 2 (2002) 277-291



Effector: Target

CTL response

Interactions of AhR with other proteins

TABLE 1. Interactions Between Signal Transduction Pathways and $\mathsf{AhR}^{a,b}$

Interactions	References
Direct interactions with AhR	
HSP90	[79]
XAP2	[80-82]
ER, ERRα	[24]
NFkB (RelA/p65)	[39]
Rb	[44-46]
RIP 140, p300/CBP	[41,51,53]
SRC-1, NCoA-2, pCIP	[41,54]
ERAP 140, SMRT	[49,50]
COUP-TF1	[24]
pp60'stc	[70,71]
tyrosine phosphorylation	[69]
Direct interactions with AhR complex proteins	
HIF-1α, PAS proteins (ARNT)	[32,35]
p300/CBP (ARNT)	[52]
SRC-1, NCoA-2 (ARNT)	[54]
SHP (ARNT)	[78]
AhRR (ARNT)	[20]
ARNT Repressor (ARNT)	[21]
CK2 (XAP2)	[74]
p23 (HSP90)	[76]
XAP2 (HSP90)	[80]
Indirect interactions (cross talk) with AhR	t,
ER	[8,25,29]
hypoxia	[33,36]
NFκB	[40-42]
PKC	[59–66]
tyrosine kinases/phosphatases	[69,72,73
c-myc, AP-1, CK2	[72]
ТGF-в	[7]
p27 (Kip 1)	[43]
NF-1	[27]
C2-ceramide	[47]

J.R. Petrulis, G.H. Perdew / Chemico-Biological Interactions 141 (2002) 25-40

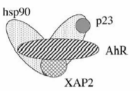


Fig. 4. Model for the arrangement of proteins found in the unliganded AhR complex.

O. Hankinson | Archives of Biochemistry and Biophysics 433 (2005) 379-386

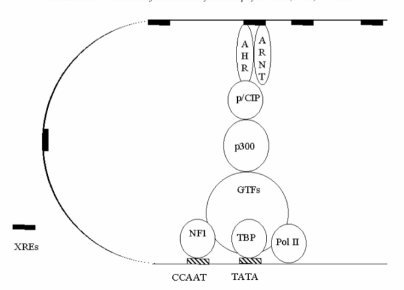


Fig. 3. Hypothetical model of coactivator recruitment at the Cyp1a1 gene.

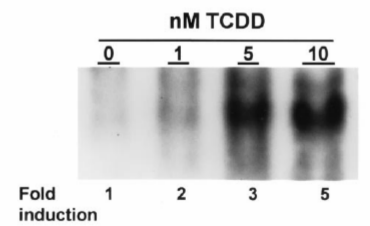
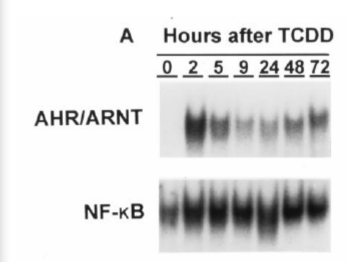
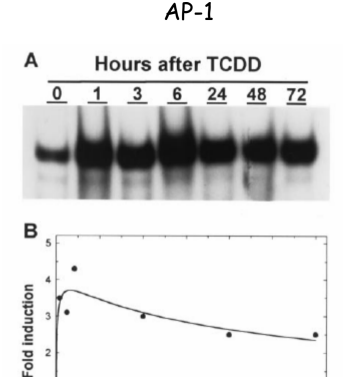


FIG. 2. Induction of c-Jun mRNA by TCDD. Hepa-1 cells were treated for 24 hr with TCDD in 0.05% DMSO at the indicated concentrations. Total RNA was extracted from these cells, fractionated in agarose–formaldehyde gels, and transferred and hybridized to a mouse c-jun probe as described in the Methods Section. Fold induction, determined by densitometry, is indicated below each lane.





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Hours after TCDD treatment

AhR-ERa crosstalk

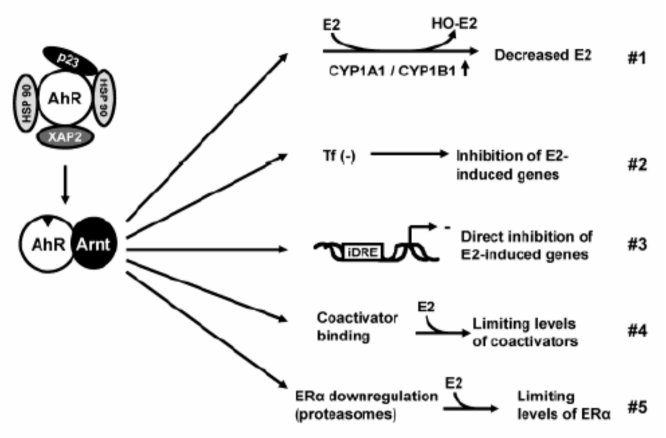
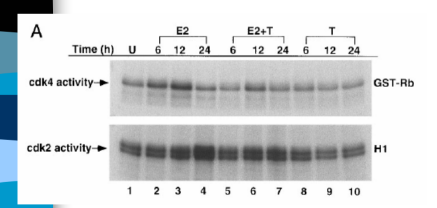


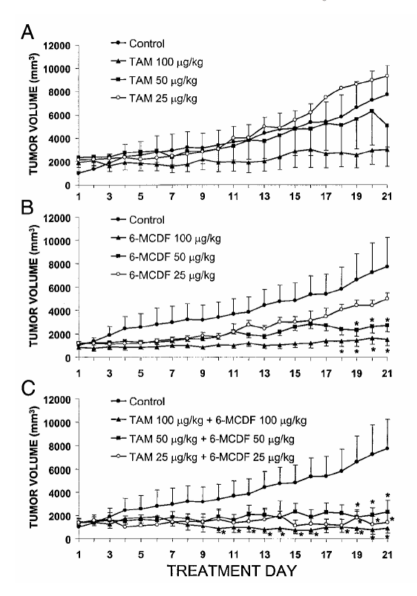
Figure 3. Proposed mechanisms of inhibitory AhR–ERα crosstalk (123–126).

Využitií AhR-ERa crosstalk v nádorové terapii?

TABLE I Effects of 17β -Estradiol and TCDD on Cell Cycle Distribution of MCF-7 Human Breast Cancer Cells a

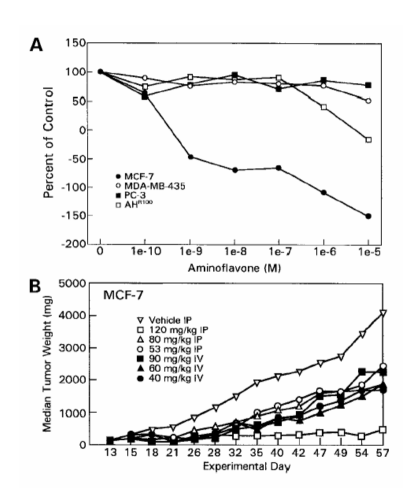
Т	Cell cycle phase (%)			
Treatment (time, h)	G_0/G_1	S	G_2/M	
Control	89.9 ± 2.1	4.9 ± 1.6	5.2 ± 0.6	
E2 (12)	87.7 ± 2.1	6.0 ± 1.4	4.4 ± 0.7	
E2 + TCDD (12)	87.2 ± 0.2	7.9 ± 0.7	4.9 ± 0.5	
TCDD (12)	89.1 ± 0.8	6.7 ± 0.8	4.2 ± 0.2	
E2 (24)	75.1 ± 0.6^{b}	23.4 ± 1.7^{b}	1.5 ± 1.2	
E2 + TCDD (24)	81.0 ± 1.3^{c}	15.8 ± 1.8^{d}	3.2 ± 0.7	
TCDD (24)	90.8 ± 0.6	5.2 ± 0.5	4.0 ± 0.9	





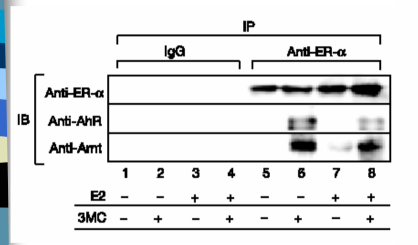
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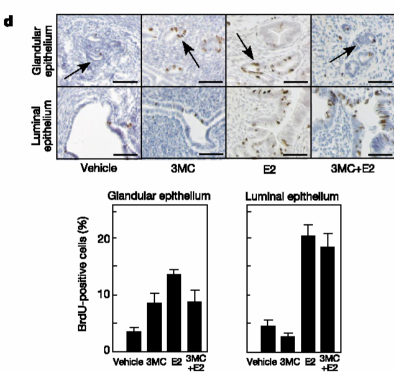
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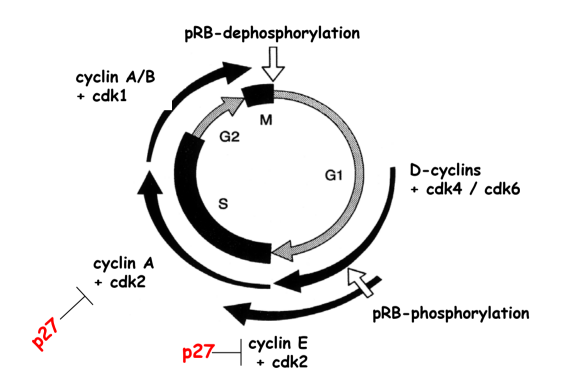
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Direct AhR-ER interaction?

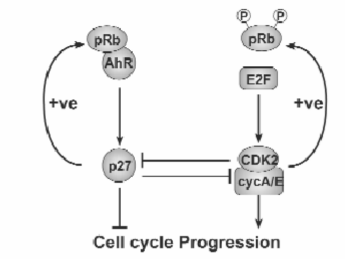




Regulation of the eukaryotic cell cycle



pRB = retinoblastoma protein cdk = cyclin-dependent kinase



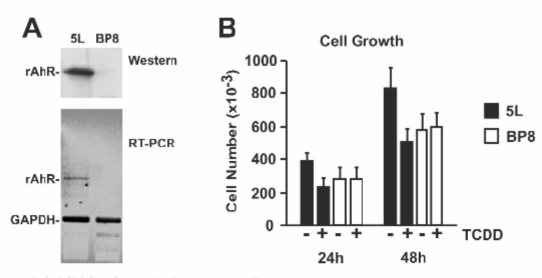


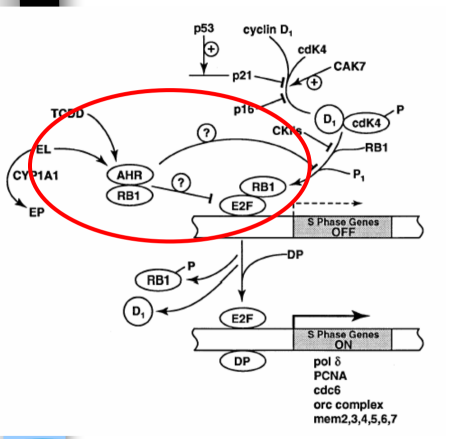
Figure 2.. TCDD induces growth inhibition in rat 5L hepatoma cells.

Panel A, total protein from 5L and BP8 cells was fractionated by SDS-PAGE and probed for AhR protein with an anti-AhR antibody (Western). Analysis of AhR expression was also performed by RT-PCR on total RNA from 5L and BP8 cells using primers specific for rat AhR (rAhR) and GAPDH (as a control for RT-PCR).

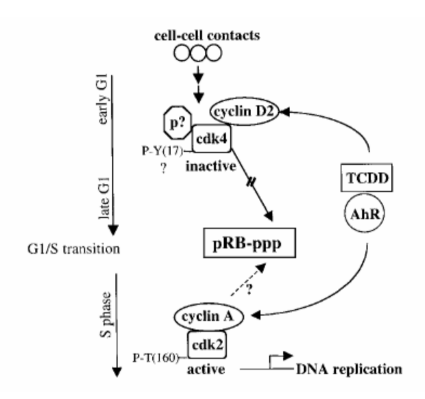
Panel B, 5L (solid bars) and BP8 (open bars) cells (2x10⁵) were grown in the presence of 10 nM TCDD (+) or absence of TCDD (-) for 24h or 48h and counted. The values presented are the mean ± S.D. of three independent experiments.

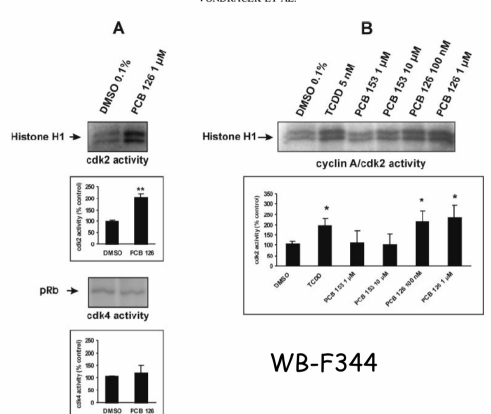
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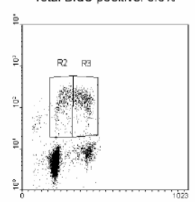


Úloha AhR v regulaci buněčného cyklu je pravděpodobně složitější

MCF-7

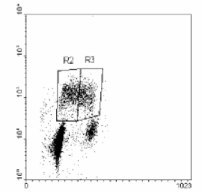


Early S-phase (left): 2.6% Late S-phase (right): 3.7% Total BrdU positive: 6.6%

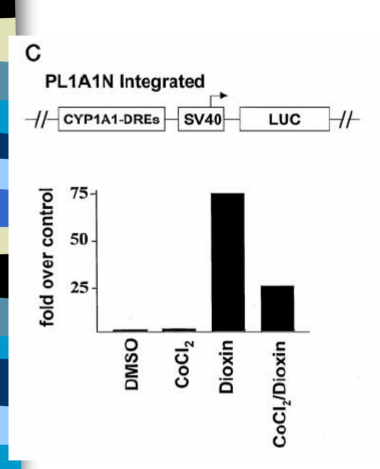


BaA

Early S-phase (left): 7.6% Late S-phase (right): 7.3% Total BrdU positive: 14.9%



? AhR-HIF-1a crosstalk?



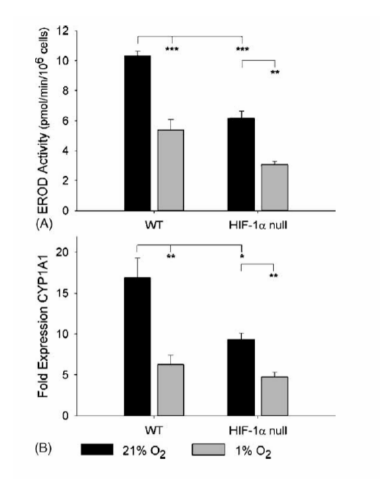


Fig. 3. Enzymatic activity (A) and gene expression (B) of CYP1A1. Rate of conversion of ethoxyresorufin. (A) was assayed in WT and HIF-1 α null cultures under normoxia (21% O₂, black bars) or hypoxia (1% O₂, grey bars) with 5 μ M 3-MC for 24 h. CYP1A1 mRNA evels (B) were measured by real time PCR after 8 h of normoxia black) or hypoxia (grey) with 5 μ M 3-MC and normalized to unreated, normoxic controls. Values are the mean and standard error for n=3: $^*p<0.05$; $^{**}p<0.01$; $^{***}p<0.001$.

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AhR-retinoid receptors crosstalk

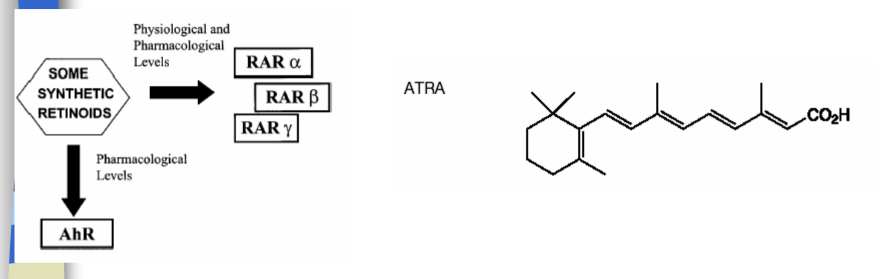


FIGURE 2 Schematic representation of the AhR/Arnt signaling pathway indicating the five steps (see text for descriptions) that have been shown to be modulated by specific retinoids.

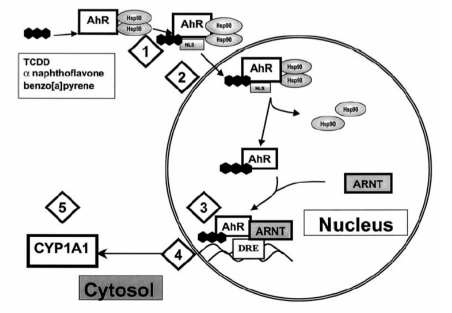


TABLE 2
Effects of Ah Receptor Ligands on Enzyme Activities Involved in Retinoid Metabolism¹

Activity	Effect	Tissue	Reference
Retinoic acid glucuronidation	1	liver, kidney	Bank et al. 1989
	1	liver	Sass et al. 1994
Retinoic acid oxidation	1	liver	Spear et al. 1988
	1	liver	Fiorella et al. 1995
	±0	liver	Andreola et al. 1997
Retinol esterification	1	hepatic stellate cells	Nilsson et al. 1996
	1	kidney	Nilsson et al. 2000
Retinyl ester hydrolysis	±0	liver	Nilsson et al. 2000

¹ TCDD was used in all studies except Sass et al. 1994 (3-methylcholanthrene) and Spear et al. 1998 (3,3',4,4',5,5'-hexabromobiphenyl). All studies were on rats except Andreola et al. 1997 (mice).

Fig. 9. Schematic depiction of the activation of MMP-1 mRNA levels by TCDD and atRA in NHKs. The data presented in this report suggest that TCDD is having an impact on MMP-1 expression in NHKs through at least two mechanisms: 1) by inducing the binding of Fos and Jun proteins to the AP-1 elements in its promoter and thereby activating transcription; and 2) by altering the expression of RAR γ and RXR α expression, which leads to an enhancement of MMP-1 mRNA stability following exposure to atRA.

