Integrated Toxicology Program and Nicholas School of the Environment and Earth Sciences, Duke University, Durham, NC

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## 2,3,7,8-Tetrachlorodibenzo-p-Dioxin (TCDD) Induces Organ-specific Differential Gene Expression in Male Japanese Medaka (Oryzias latipes) research findings conclusions

# overview

2,3,7,8-Tetrachlorodibenzo-p-dioxin (TCDD or dioxin) is a widely studied polychlorinated, tricyclic aromatic compound known to induce adverse affects in humans and wildlife, including cancer, reproductive and developmental effects. immunotoxicity, and cardiovascular disease. In recent years, unbiased genomewide gene expression analysis has been exploited using in vitro- and in vivobased mammalian models to uncover additional AhR-dependent or -independent TCDD-responsive genes. However, the majority of these studies have centered on gene expression analysis in TCDD-exposed hepatoma cells or liver. To this end, we utilized suppression subtractive hybridization (SSH) as an impartial screening tool to initially evaluate gualitative gene expression changes in male Japanese medaka (Oryzias latipes) organs (brain, liver, and testis) following intraperitoneal TCDD injection (10 µg-TCDD/kg-body weight) and exposure for 48 h. SSH analysis provides non-biased evaluation of mRNA-level differences between control and toxicant-exposed animal tissues, and relies on hybridization-dependent subtraction of equally abundant transcripts and selective PCR-amplification and enrichment of differentially expressed genes.

In this study, after identification of suspected differentially expressed transcripts based on SSH, expression of genes hypothesized to be strongly responsive to TCDD exposure was semi-quantified using organ-specific replicate nylon membrane cDNA arrays. Moreover, gualitative histopathologic evaluation was used to associate organ histopathology with gene expression patterns in male medaka brain, liver, and testis. Overall, we demonstrate that TCDD induces organ-specific qualitative and semi-quantitative gene expression differences in male medaka, and that these differences are associated with adverse histopathological changes. Based on these data, brain-, liver- and testis-specific mRNA-level targets in male medaka were identified as promising biomarkers of TCDD-induced toxicity for future investigations.



### 2,3,7,8-Tetrachlorodibenzo-p-dioxin

Sources include incomplete combustion, industrial discharge, and biological processes

. In the U.S. (1995), air emissions account for the majority of environ TCDD release, where municipal solid waste incineration, backvard refuse barrel burning, and medical waste incineration contribute >70% of total lioxin releases (NCEA, ORD, U.S. EPA)

Insoluble in water (19.3 ng/L at 25°C) and relatively immobile and persistent on soil ( $t_{_{1/2}}$  = 1-3 yr) and sediment ( $t_{_{1/2}}$  > 1.5 yr) surfaces.

- Highly lipophilic (log  $K_{\rm yw}$  = 6.80) and significant bioaccumulation and biomagnification potential (log BCF = 3.2-3.9 in fish) in ecological food chains, resulting in elevated levels in wild animals and commercial food



Figure 1. Upon entry into cellular cytosol, TCDD is highly specific for the aryl hydrocarbon (Ah) receptor (AhR). Following ligand binding, activated AhR enters the nucleus, dissociates from the XAP2-HSP0 complex, and forms a heterodimer with AhR nuclear transporter (ARNT); this transcription factor complex binds to complex binds dioxin response elements (DREs) in promoter regions of target genes, driving activation of the "Ah gene battery" and regulation of numerous yet unidentified dioxin-inducible genes. In general, these TCDD-induced signaling pathways elicit down-stream adverse effects such as oxidative stress, cell cycle arrest, and apoptosis. Pathway image from Biocarta (biocarta.com).

			Medaka Organ							(6-8-μm) (below) of a control	Contraction of the second	THOMAS STATES
	-	Brain	Liver	Testis	_	Total				adult male medaka ip-exposed to vehicle (DMSO) for 48 hr.		
Colonies screened		768	768	768		2304		_		egg-laying freshwater fish	and the second of the	
True positives sequenced		184 (23.9% <sup>a</sup> ) 335 (43.6%)		261 (33.9	9%)	780 (33.9%)			native to Japan, Korea, and eastern China. Adult males (6-	-		
Redundant genes		98 (12.8%)	258 (33.6%)	.6%) 214 (27.9%)		570 (24.7%)			8-months-old) typically have an	Adult male med	laka (3-4 cm l	
No significant homology		36 (11.2%)	77 (10.0%)	77 (10.0%) 47 (6.1%)		210 (9.1%)				average body weight of ~350-		
Non-redundant genes		58 (7.6%)	112 (14.6%)	165 (21.5	165 (21.5%)		335 (14.5%)			400 mg.		0.25 000
enes spotted on cDNA	array 8	3 (1.0%)	22 (2.9%)	12 (1.6%	)	42 (1.8	3%)	_		80	See Company Stations	0.25 cm
All percentage values a	re relative to	total coloni	es screened from eac	h organ.						Cr-		
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58 7 11	2	1	2% Metabo	am.		$\mathbf{N}$	12%					the second second
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		1										
165					and instantion		United			Control		10 µg/kg
			19% 20%		17%		105					And the second second
Testis		Tes	stis	Figure	3. Top si	x Gene (	Ontology	™ (GO)		A	B	Al Share
Figure 2 Numerical distribution of		Cell mobility 10%			term distribution of differentially expressed						a the start	2.24
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edaka brain, liver, and testis	following a 48-	h		m TCDD/	kg-body	weight e	xposure	GO		State of the second		12 8 83
) µg-TCDD/kg-body weight e	xposure. Of 33	5		terms	were de	termined	by Lo	cusLink	.⊆	A STATE AND A STATE AND A STATE		1
tal genes identified, only 2	2-3% of TCDL	)- Metabo 121		(NCBI)	, and w	ere bas	ed on	general	<u>n</u>	12月1日 - 12月1日 - 12月1日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日日		and the second
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igh degree of organ specificit	y at the level	of Signal 1	ransduction	gene	unctionali	tv. espec	ially wit	hin the			1 S. A. C.	1.1
ene expression in respon	nse to TCD	D	13% Proteolysia 13%	medak	a brain an	d liver libr	aries.					1. 184
posure.									I	1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1. 1		
le 2. Array data summary of gene	expression chan	ges in male meda	ka organs following 48-h expo	sure to 2,3,7,8-to	trachlorodi	benzo-p-di	oxin (TCD	D).				
e Name (in alphabetical order)	0	Drigin * Accession	ID <sup>b</sup> Gene Ontology™	Brain TRT/CTL <sup>c</sup>	- L	CTL P	Testi TRT/CT	s L P		C	D	1
n, beta	1	estis AJ630123	Cell motility	0.951 0.	11 1.116	0.319	0.684	0.035**			2 S.A	
coenzyme A dehydrogenase a.2.HS.obyccomtein	8	krain AAH45911	Fatty acid metabolism	1.049 0.	19 1.15	0.209	0.764	0.079*			第二人工工作	and the second
ase, alpha	1	estis AF416651	Carbohydrate metabolisr	1.032 0.	197 1.18	0.077*	0.683	0.017**	er			NAME OF STREET
poprotein A-1 poprotein B	L L	iver AAP20157 iver AAP97376	Lipid transport Lipid transport	1.029 0. 0.920 0.	1.000	0.995	0.730	0.009***	.≥	and the second second second second		/ Deletion
oprotein C-I	L	iver Q9XSN5 krain AAP20154	Lipid transport Lipid transport	1.279 0.	18 1.244 79 1.143	0.096*	0.990	0.922 0.008***				1 and the second
poprotein E salt-activated linase	i.	iver CAB65356	Lipid transport	2.245 0.	121 1.065	0.557	0.849	0.052*				Call Carlos
oxypeptidase B	1	estis AB099302	Proteolysis	1.023 0.	18 1.160	0.123	0.716	0.073*			17 P ( ) ( )	ACC STREET
c binding protein julation factor XIIIb, precursor	1	estis XP_21651 iver XP_35523	6 Mitosis 5 Blood coagulation	1.052 0. 0.957 0.	1.181 144 1.170	0.093*	0.710	0.013** 0.012**		And the second second second second		S. S. S. S.
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plement C4 chrome P450 1A	E	Iver AB025577 Irain AY297923	Complement activation Drug/toxicant metabolism	1.071 0.	51 1.090 88 1.61	0.267	0.886	0.012** 0.292			1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1 1	18 A 10
chrome P450 1A tase 1. precutsor	1	iver AY297923 estis AB029755	Drug/toxicant metabolish Proteolysis	1.585 0.1	37 1.000	0.069*	1.009	0.958		E	1 - A AR	
tase A, precursor	1	estis AB029756	Proteolysis	1.051 0.	1.12	0.265	0.803	0.103			1	1 Va
nogen, alpha	i.	iver AAH41754	Blood coagulation	1.008 0.	73 1.07	0.318	0.746	0.006***		A STATE OF A STATE OF A STATE		
nogen, gamma eraldehyde 3-phosphate dehydrogena	150 E	Iver AJ310418 Irain BAB62812	Blood coagulation Glycolysis	1.053 0.0	1,114	0.224	0.823	0.150 0.026**				A CONTRACT
cidin precursor choriolytic enzyme 1, precursor	i i	iver AAS66305 estis AAL40376	Immune response Unknown	0.969 0.	91 1.464	0.074*	0.587	0.003***	<u>.</u>	The states		19
sex-determining protein	1	estis AY129241	Sex differentiation	1.224 0.	1.45	0.056*	0.768	0.087*	st		Carlos A	0 9 8 Ve
hondrial DNA	1	estis AP004421	Unknown	0.935 0.	59 0.97	0.831	0.843	0.045**	<u>ه</u>		A AND	and a
inositol oxygenase H dehydrogenase	L	Iver AF401311 Irain NP_73981	5 Fatty acid biosynthesis	1.046 0.	48 1.17 88 1.078	0.119	0.756	0.112 0.006***		A State of the second second	A 4	ARCEN S
eoside diphosphate kinase n A2	E	krain AF266216	Nucleoside metabolism Development	1.128 0.0	62 1.200	0.260	0.867	0.206			3 3 3 3 4	the seal of a bull
erebellin-like protein	L	iver AAF04305	Neurogenesis	0.993 0.	71 1.134	0.659	0.745	0.002***				2000
tyltin-binding protein	1	iver BAB83525	Spermatogenesis Unknown	1.015 0.1	1.10	0.286	0.767	0.009***				TRUE TOWN I WE
sferrin sinogen 1	1	iver D64033 estis AB029750	Iron ion transport Proteolysis	0.949 0.1	79 1.116	0.133	0.725	0.010***		Figure 5. Histopathology of adult male medak	a following a 48-h 10	ua-TCDD/ka-body
lin, alpha	E	Irain AAP89018	Microtubule movement	0.930 0.	21 1.47	0.046**	0.686	0.003***		Serial 6-8-µm whole-body sagital sections we	re mounted on glass	slides and stained
-temperature-acclimation-related-65	kDa-protein L	iver AB075198	Acute phase response	1.022 0.1	14 0.91	0.413	0.729	0.019**		and eosin. All slides were imaged at 40X obje	ective. Scale bar = 10	µm. (A) Brain from
										control medaka: (B) brain from TCDD-expose	d medaka: (C) liver fro	m vehicle (DMSO)
rol (Arabidopsis Cab1)		4/A X56062	NA	0.984 0.	15 1.021	0.466	1.006	0.777		(D) liver from TCDD-exposed medaka: (E) toot	tis from vehicle (DMSO	) control medaka: a

ubtracted cDNA library where gene was identified. I GenBank (<u>http://www.iccl.ntm.mk.gov)</u> and identify sequences with significant homotogy (E-1X10 identify gene expression differences based on least payler means of normalized array density values identify gene expression differences based on least payler.

nd stained with hem A) Brain from vehicle (DMSO)
 icle (DMSO) control medaka;
 icl medaka; and (F) testis from



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## suppression subtractive hybridization

• 780 total cDNAs were sequenced: 210 (27%) sequences had no significant homology and 98, 258, and 214 sequences from the brain, liver, and testis respectively (570 total) sequences had significant homology to known sequences in NCBI databases.

 58, 112, and 165 total distinct genes for the brain, liver, and testis respectively; although the same number of colonies were screened, the frequency of one or more genes within the library affected the library gene diversity and number of non-redundant genes.

· 2-3% of TCDD-responsive genes in all three libraries were shared between any two libraries, and only 4 genes were identified in all three libraries, suggesting a high degree of organ specificity at the level of gene expression in response to TCDD exposure.

 31% (13 genes) and 32% (26 genes) of genes identified in the brain and liver library are involved in metabolism, and 41% (50 genes) of identified genes in the testis library have no known function.

### gene arrays & organ histopathology

· Of 42 transcripts screened. CYP1A mRNA was the only transcript significantly higher in TCDD-exposed brain, whereas 12 transcripts (including CYP1A) were significantly higher in TCDD-exposed liver and 34 transcripts were significantly lower in TCDD-exposed testis.

· Minimal histopathological changes observed in brain; glycogen depletion and mild hepatocyte hypertrophy observed in liver; and disruption of primary spermatocytes and interstitia observed in testis.

· Based on gene expression and histopathological data, male medaka testis is highly sensitive to TCDD exposure, and mechanisms of reproductive toxicity are currently being pursued.

- · Based on these data, promising organ-specific biomarkers of acute TCDD-induced toxicity include:
- Brain: CYP1A (xenobiotic metabolism) Liver: α-Amylase (glycogen metabolism), Hepcidin (immune defense), Tributyltin-binding protein (unknown function) Testis: Male sex-determining protein (sex differentiation), Protamine (spermatogenesis)

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