Kenyon College Digital Kenyon: Research, Scholarship, and Creative Exchange

Kenyon Summer Science Scholars Program

Summer Student Research Scholarship

Summer 2017

Characterizing the Aryl Hydrocarbon Receptor (AHR) within the order Anura

Reed Crocker *Kenyon College*

Hannah Hertz Kenyon College

Alula Hunsen Kenyon College

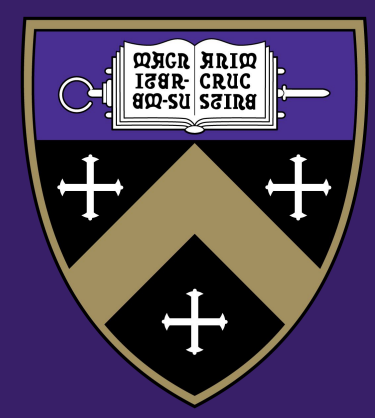
Wade Powell *Kenyon College*

Follow this and additional works at: https://digital.kenyon.edu/summerscienceprogram

Recommended Citation

Crocker, Reed; Hertz, Hannah; Hunsen, Alula; and Powell, Wade, "Characterizing the Aryl Hydrocarbon Receptor (AHR) within the order Anura" (2017). *Kenyon Summer Science Scholars Program.* Paper 64. https://digital.kenyon.edu/summerscienceprogram/64

This Article is brought to you for free and open access by the Summer Student Research Scholarship at Digital Kenyon: Research, Scholarship, and Creative Exchange. It has been accepted for inclusion in Kenyon Summer Science Scholars Program by an authorized administrator of Digital Kenyon: Research, Scholarship, and Creative Exchange. For more information, please contact noltj@kenyon.edu.



Characterizing the aryl hydrocarbon receptor (AHR) within the order Anura

Reed Crocker, Hannah Hertz, Alula Hunsen, Wade Powell Department of Biology, Kenyon College, Gambier, Ohio 43022 USA

Abstract

The aryl hydrocarbon receptor (AHR) is a ligand activated transcription factor that mediates the toxic effects of dioxin-like compounds, such as 2,3,7,8tetrachlorodibenzo-p-dioxin (TCDD). Once bound by TCDD, the AHR complex regulates the transcription of a battery of genes that ultimately impart toxicity. Species specific genotypes of AHR determine structure and affinity for TCDD. These differences in AHR ligand affinity can often explain the degrees of TCDD sensitivity between different vertebrate clades. Previous AHR characterizations in the frog Xenopus laevis, the salamander Ambystoma mexicanum, and the caecilian *Gymnopis multiplicata* predicted low affinity binding and insensitivity to TCDD in all three amphibian orders 1,2,3 . While these data do suggest that low affinity binding and insensitivity is common to all three amphibian orders, they do not confirm this trend is consistent within each clade. In studying a wider group of amphibians from within the frog (Anura) order, we seek to confirm this trend is conserved among related amphibians. To determine this, we chose to characterize two frogs: *Xenopus borealis*, which is phylogenetically similar to the previously mentioned X. laevis; and Bufo marinus, a more distantly related frog. The cloned cDNAs of each frog revealed that the *B. marinus* AHR shared 83-84% sequence identity to both *X. laevis* paralogs, while the two *X. borealis* paralogs shared 93% and 91% identity.

Objective

To characterize the AHR gene(s) from X. borealis and B. marinus and compare them to the well studied model organism, X. laevis. Analysis of AHR sequences and structure will clarify the extent to which the AHR amino acid sequence differs between frogs.

Results

Figure 1		
	<u>RT-PCR</u>	

Figure 2

X. borealis X. laevis B. marinus X. borealis AHR1β cDNA 1 cDNA 2 cDNA

Conclusions

• The degenerate primer design of the experiment is a viable method to find AHR orthologs in non-model organisms. • The AHR amino acid sequences of closely related species are more distantly related ones.



Degenerate Primers	A2 (Forward)	5'-CGGGATCCGAYTAYCTIG GITTYCAR-3'
	B2 (Reverse)	5'-GCTCTAGAGCTCIRCYTCI GTRTAICC-3'
Cycling Conditions		94°C/ 5 min; (94°C/ 15 sec; 50°C/ 30 sec; 68°C/ 1 min)x43; 4°C/ hold

Figure 1. RT-PCR Primer design. The degenerate primers were designed from conserved regions within vertebrate AHRs that have been previously shown to successfully amplify cDNA from many vertebrate species⁵.

<i>Χ. laevis</i> AHR1α	92% (161/174)	95% (164/172)	88% (158/175)	84% (147/176)
<i>Χ. laevis</i> AHR1β		89% (156/174)	93% (164/175)	83% (146/176)
<i>X. borealis</i> cDNA 1			91% (161/175)	86% (151/176)
<i>X. borealis</i> cDNA 2				85% (150/176)
<i>B. marinus</i> cDNA				

Figure 2. Amino Acid identity analysis. X. borealis was found to have two paralogs of AHR which shared high sequence identity with the single B.marinus AHR shared considerably less identity with either Xenopus species. Numbers in parenthesis show shared amino acids divided by total amino acids aligned.

- Perform RACE-PCR to obtain the full open reading frame for both *X.borealis* and *B.marinus*.
- Obtain AHR cDNA and ORF from additional Anura species.
- Begin cataloging the AHR sequences of the order Caudata (salamanders).
- Use the collection of these AHRs to gain predictive power about the TCDD sensitivity of unsequenced or untreated amphibians.

Acknowledgements

- Kenyon College Summer Science Scholars Cascade Program
- Professor Wade Powell for his guidance throughout this project
- The amazing members of the molecular toxicology lab at Kenyon College that aided me in conducting this research

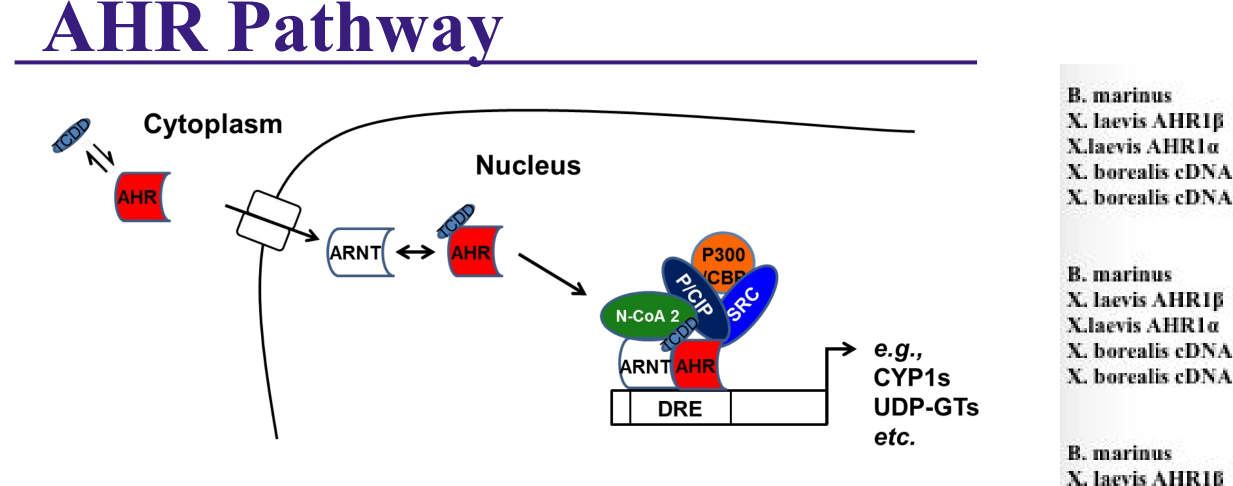


Figure	3	
0		

X.laevis AHR1a

X. borealis cDNA 1

X. borealis cDNA 2 120

- 5	¹ GSDYLGFQQSDVIHQSVFELIHTEDRPEFQRQLHWALDPVQPSD-TKRSSEGNEFTMI ¹ IQDYLGFQQSDVIHQSVYELIHTEDRLEFQRQLHWAFDPAHPSSSVQRSP-DEEFSAL	N -
- 3	1 IODYLGFQQSDVIHQSVYELIHTEDRIEFQRQLHWAFDPAHPSSSLQRSP · · · DDTAL	Τ-
- 3	J GSDYLGFQQSDVIHQSVYELIHTEDRIEFQRQLHWAFDPAHPSSSLQRSP DDTAL	Τ-
- 3	1 GSDYLGFQQSDVIHQSVYELIHTEDRLEFQRQLHWAFDPAHPSSSVQRSPGDKEFSAL	Τ-

us	60	CYNPDQLPPENSSFMERNFVCRLRCLLDNSSGFLAMNFQGRLKFLHGQNKKGKDGSVIPP	H9
AHR16	59	CYKPEQLPPENSSFMERNFVCRLRCLLDNSSGFLAMNFQGRLKFLHGQNKKGKDGS <u>I</u> LPP	II8
AHR1a	57	C F K P E Q L P P E N S S FME R N F V C R L R C L L D N S S G F L AMN F Q G R L K F L H G Q N K K G K D G S T L P P	116
lis cDNA 1	57	CYEPEQLPPENSSFMERNFVCRLRCLLDNSSGFLAMNFQGRLKFLHGQNKKGKDGSMLPP	H6
lis cDNA 2		CYKPEQLPPENSSFMERNFVCRLRCLLDNSSGFLAMNFQGRLKFLHGQNKKVKDGSILPP	

Figure 3. cDNA alignment. The partial cDNA amino acid sequences for each species were aligned using MacVector 14.5.3 Assembler.

F

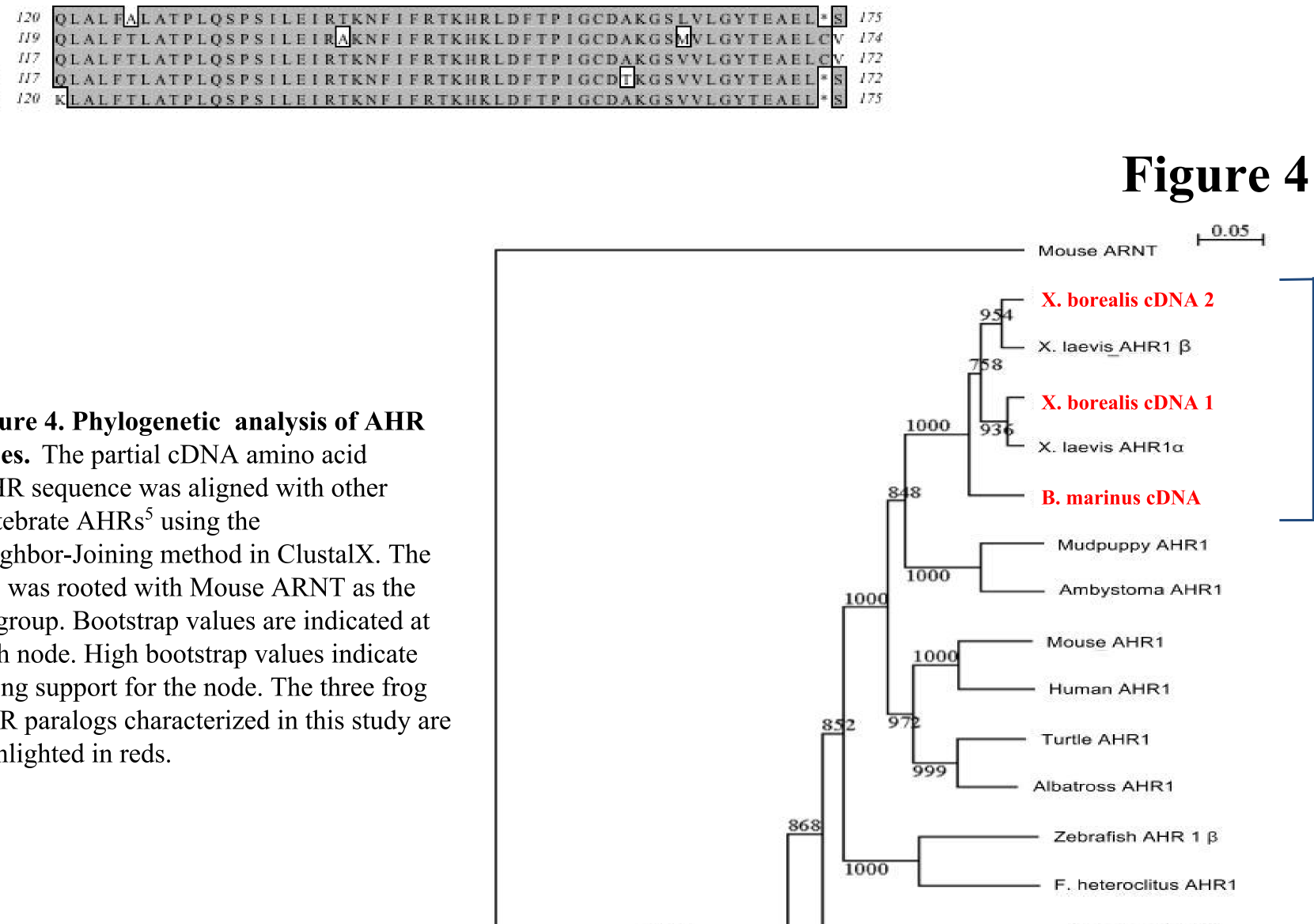
(sbo.

TCDD and other ligands bind AHR in the cytoplasm, inducing translocation to the nucleus and dimerization with ARNT. AHR:ARNT is transcriptionally active and binds specific regulatory elements designated dioxin response elements (DREs), resulting in the recruitment of cofactors and induction of a battery of target genes.

Xenopus Evolution

Studies into the phylogenetic history of *Xenopus* have postulated that 34 million years ago a divergence formed the L & S subgenomic species of ancestral *Xenopus*, one with AHR1α and the other with AHR1 β (T2). The two distinct species then hybridized approx. 17 million years ago to duplicate their genomes (T3). Modern *Xenopus* species like *X*. *laevis* and *X*. *borealis* are now tetraploids with two paralogs of AHR⁴.

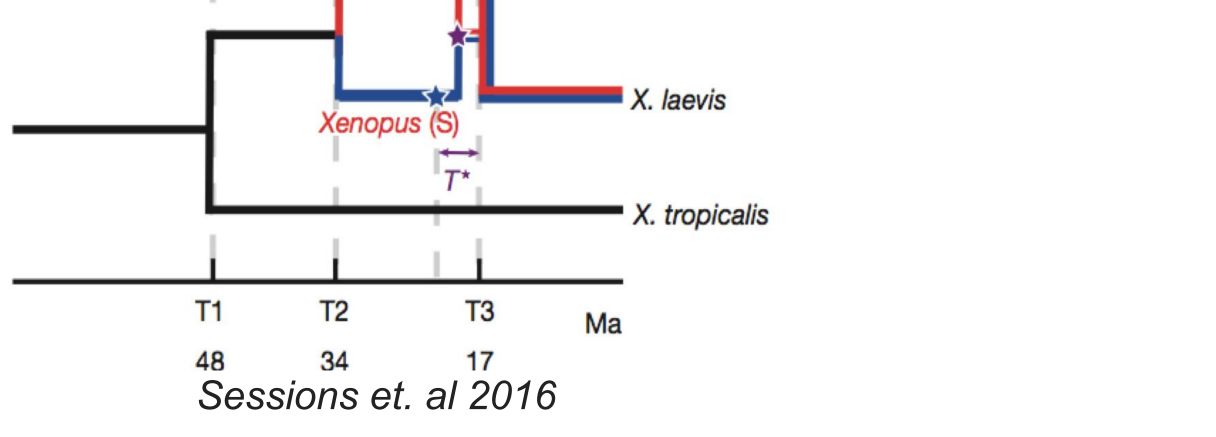
Figure 4. Phylogenetic analysis of AHR genes. The partial cDNA amino acid AHR sequence was aligned with other vertebrate AHRs⁵ using the Neighbor-Joining method in ClustalX. The tree was rooted with Mouse ARNT as the outgroup. Bootstrap values are indicated at each node. High bootstrap values indicate strong support for the node. The three frog AHR paralogs characterized in this study are highlighted in reds.

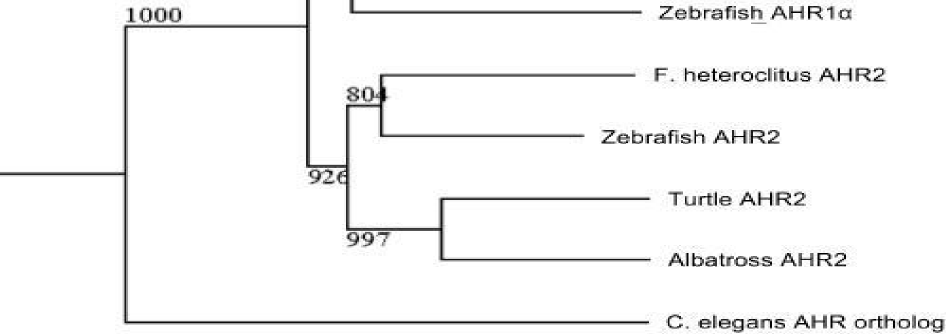


References

- Odio, C., Holzman, S. A., Denison, M. S., Fraccalvieri, D., Bonati, L., Franks, D. G., Hahn, M. E., and Powell, W. H. (2013). Specific Ligand Binding Domain Residues Confer Low Dioxin Responsiveness to AHR1beta of *Xenopus laevis*. *Biochemistry*. 52 (10): 10.1021/bi301722k.
- 2 Shoots, J., Fraccalvieri D., Franks D.G., Denison, M.S., Hahn M.E., Bonati, L., and Powell, W.H. (2015). An Aryl Hydrocarbon Receptor from the Salamander Ambystoma mexicanum Exhibits Low Sensitivity to 2,3,7,8-tetrachlorodibenzo-*p*-dioxin. Environ. Sci. Technol. 49, 6993-7001.
- Kazzaz, S. (2016). An Aryl Hydrocarbon Receptor Sequence from the Caecilian *Gymnopis* multiplicata Suggests Low Dioxin Affinity Is Common to All Amphibian Orders, unpublished poster, Department of Biology, Kenyon College, Gambier OH USA.
- Session, Adam M., Yoshinobu Uno, Taejoon Kwon, Jarrod A. Chapman, Atsushi Toyoda, Shuji Takahashi, Akimasa Fukui, et al. "Genome Evolution in the Allotetraploid Frog *Xenopus* Laevis." Nature 538, no. 7625 (October 20, 2016): 336-43.
- 5 Hahn, M.E., Karchner, S.I., Shapiro, M.A., Perera,

Xenopus (L) X. borealis





S.A. (1997). Molecular Evolution of two vertebrate aryl hydrocarbon (dioxin) receptors AHR1 and AHR2) and the PAS family. *Evolution*. 94: 13743-13748. 6 Hahn, M.E., Karchner, S.I., and Merson, R. R. (2017). Diversity as opportunity: Insights from 600 million years of AHR evolution. *Toxicology*. 2:58-71.