

**Research Paper** 





2016; 4: 29-41. doi: 10.7150/jgen.16885

# De novo Assembly and Analysis of the Chilean Pencil Catfish *Trichomycterus areolatus* Transcriptome

Thomas T. Schulze<sup>1</sup>, Jonathan M. Ali<sup>3</sup>, Maggie L. Bartlett<sup>1</sup>, Madalyn M. McFarland<sup>1</sup>, Emalie J. Clement<sup>1</sup>, Harim I. Won<sup>1</sup>, Austin G. Sanford<sup>1</sup>, Elyssa B. Monzingo<sup>1</sup>, Matthew C. Martens<sup>1</sup>, Ryan M. Hemsley<sup>1</sup>, Sidharta Kumar<sup>1</sup>, Nicolas Gouin<sup>4,5,6</sup>, Alan S. Kolok<sup>1,2</sup>, Paul H. Davis<sup>1</sup><sup>⊠</sup>

- 1. Department of Biology, University of Nebraska at Omaha, Omaha, Nebraska 68182, USA;
- 2. Center for Environmental Health and Toxicology, University of Nebraska Medical Center, Omaha, Nebraska 68198, USA;
- 3. Department of Environmental, Agricultural and Occupational Health, University of Nebraska Medical Center, Omaha, NE,
  - 68198-6805, United States;
- 4. Departamento de Biología, Universidad de La Serena, La Serena, Chile;
- 5. Centro de Estudios Avanzados en Zonas Aridas, La Serena, Chile;
- 6. Instituto de Investigación Multidisciplinar en Ciencia y Tecnología, Universidad de La Serena, La Serena, Chile.

🖂 Corresponding author: Dr. Paul H. Davis. pdavis@unomaha.edu

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Published: 2016.09.06

#### Abstract

*Trichomycterus areolatus* is an endemic species of pencil catfish that inhabits the riffles and rapids of many freshwater ecosystems of Chile. Despite its unique adaptation to Chile's high gradient watersheds and therefore potential application in the investigation of ecosystem integrity and environmental contamination, relatively little is known regarding the molecular biology of this environmental sentinel. Here, we detail the assembly of the *Trichomycterus areolatus* transcriptome, a molecular resource for the study of this organism and its molecular response to the environment. RNA-Seq reads were obtained by next-generation sequencing with an Illumina® platform and processed using PRINSEQ. The transcriptome assembly was performed using TRINITY assembler. Transcriptome validation was performed by functional characterization with KOG, KEGG, and GO analyses. Additionally, differential expression analysis highlights sex-specific expression patterns, and a list of endocrine and oxidative stress related transcripts are included.

Key words: de novo transcriptome, assembly, catfish, Trichomycterus areolatus

## Introduction

*Trichomycterus areolatus* (described by Valenciennes, 1846) is a threatened pencil catfish found in Chile [1, 2]. This freshwater stream fish has a broad distribution ranging from the Coquimbo to the Los Lago regions of Chile (30° *S and* 43° *S respectively*) [1, 3]. The genus *Trichomycterus* is found throughout freshwater ecosystems in both Central and South America and includes over 120 species [4]. Individual species are commonly restricted to specific river areas either by geographical restriction or habitat preference [5]. As a result, *T. areolatus* displays an

intricate integration with its environment; yet the species is fairly distributed throughout the region, providing an opportunity for its use as a model sentinel organism to study environmental stress in Chile's freshwater streams [6,7]. As a benthic fish, *T. areolatus* displays interactions with riverine sediments and substrates [8]. This species tends toward a generalist eating strategy, eating aquatic insects [9] as well as the surface organisms present on stream, plant, and rock surfaces [10]; consequently, its food preferences are known to vary by season [4].

The Trichomycterus fish generally range between 50 - 150 mm [11]. The mature fish in this species will reach average sizes of 56 and 51mm for males and females respectively [12]. They display poor secondary sex characteristics, making the identification of sex by external means difficult [13]. The spawning season takes place during October and November, based on observation of the female gonads rapidly increasing in size just before the season begins [13]. The females are capable of releasing eggs several times throughout the spawning season. The males also exhibit numerous fertilization events throughout the spawning season; however, the male fish in this species do not participate in progeny care [14]. The T. areolatus genome is described as diploid, with the typical individual displaying 2n = 54chromosomes; however, intra-individual variation has been reported [15]. This chromosome number appears common among the Trichomycterus genus and suggests conservation.

Genomes and/or transcriptomes have only been developed for fish that have either economic or scientific value including: *Salmo salar* (Atlantic salmon), *Cyprinus carpio* (common carp), *Pimephales promelas* (fathead minnow), and *Danio rerio* (zebrafish). There are limited transcriptomic resources available for fish native to South America, and none available of the Trichomycterids. Current molecular research with this organism is limited, and includes a karyotype [15], seasonal variation in biomarkers [8], local industrial discharge impacts [6], agricultural disturbances [16], microsatellite loci for conservation resources [17], and population genetics [18].

In this paper, we detail the assembly and analysis of the transcriptome for *T. areolatus*. Fish samples obtained from Chile were sequenced by next-generation sequencing and later assembled into a *de novo* transcriptome using Trinity. Transcriptome validation analyses are included and discussed. Transcriptomic data is deposited at NCBI SRA under accession SRP077018 and the completed assembly is freely available.

# **Results and Discussion**

## **Transcriptome Characteristics**

The constructed transcriptome assemblies are available at http://www.davislab.net/ trichomycterus/. Assemblies available include: a complete transcriptome nucleotide assembly, a representative assembly (see Methods) consisting of 64,385 unique transcripts, and a translated protein assembly of the representative transcripts. De novo assembly was accomplished using TRINITY. Two samples (whole male and whole female adult fish) provided RNA for the assembled transcriptome, which was assembled from Illumina® paired-end reads. A resulting 41.8Gb of output (Table 1) was utilized to create the assembly. General statistics of the assembled representative transcriptome are included in Table 2.

Table 1. Transcriptome Tissue Sequencing Details. RNAsamples were from 4.4 and 5.4ug of RNA for male and female fish,respectively. Sequencing was performed on an Illumina® Hi-Seq2500. BP: base pair, GC: G-C nucleotide ratio.

Tissue	Total Reads	Total Output (bp)	GC Content (%)
Whole Female	328,721,780	32,872,178,000	48%
Whole Male	88,794,542	8,879,454,200	47%

Table2.TrichomycterusareolatusRepresentativeTranscriptomeCharacteristics.TherepresentativeTrichomycterus areolatustranscriptome assemblywas analyzed forgeneralcharacteristicslistedabove.PutativeproteincodingtranscriptswereincludedtranscriptswereincludedandidentifiedbyTransDecoder.Redundanttranscriptswereremoved byCD-HITwhichcollapsesredundantandhighly similar sequencesinto consensus sequences.

Total Transcripts	Mean Length (bp)	Median Length (bp)	N50	GC Content
64889	1484.85	857	2671	47.5%

## **Organism Phylogenetics**

Assembled *T. areolatus* sequences were selected and aligned to publically available sequences from various fish to examine phylogenetic relatedness by identifying orthologs (Table 3). Of the fish compared, *Ictalurus punctatus* (Channel catfish) showed the highest relationship (87.6%) in a concatenated set of conserved sequences, followed closely by *Cyprinus carpio* (Common carp) at 86.5%. Intra-genus phylogenetic comparison of *Trichomycterus* was not possible due to a lack of published sequences.

## Assessing Full-length Transcript Coverage

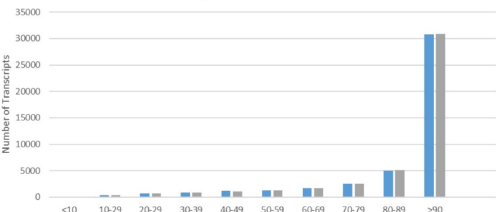
In an effort to deduce the full-length nature of the transcripts within the representative assembled transcriptome, coverage histograms arising from alignments with non-redundant protein sequences of two model organisms—*Salmo salar* and *Danio rerio*—were produced. Model organisms were selected over more closely related organisms (Table 3) due to their more complete publically available transcriptomes. Figure 3 illustrates a histogram for the distribution of transcript length when the representative *T. areolatus* transcriptome was compared to related species. Length coverage exceeding 90% of the other organism's transcript length was found in 64.7% of *T. areolatus* protein sequences upon alignment with *Salmo salar* proteins and 69.1% of *Danio rerio* proteins (Figure 3). Accounting for genetic differences between these organisms, these data suggest that the produced representative transcriptome has a high degree of full-length transcripts.

## Functional Analyses – Putative Transcript Functional Characterization

Multiple analyses exist that allow transcripts to be annotated and grouped by function. This includes Gene Ontology (GO), Eukaryotic Orthologous Groups (KOG), and Kyoto Encyclopedia of Genes and Genomes (KEGG) analyses. These annotation and grouping algorithms were applied to examine the putative function of transcripts, and as a quality control technique to evaluate transcriptome completeness when compared to well-developed transcriptomes from *Danio rerio, Salmo salar,* and *Cyprinus carpio.* Figure 4 details the **GO** analysis, which is capable of classifying predicted gene products by cellular component, molecular function, and biological process [19]. GO groupings between organisms suggest that the selected fish all share a pattern of functional distribution of gene products.

Table 3. Phylogenetic Comparison to Other Fish. Transcripts produced in this study were concatenated and aligned to published sequences of fish species and a percent identity matrix was computed. This analysis utilized *Trichomycterus areolatus* transcripts: Ta\_155828, Ta\_53325, Ta\_192266, Ta\_56196, and Ta\_56194 for cytochrome c oxidase subunit III, HSP70, NADH dehydrogenase subunit 5, estrogen receptor, and glutathione s-transferase kappa 1, respectively.

Species	Percent Identity
Ictalurus punctatus (Channel catfish)	87.6
Cyprinus carpio (Common carp)	86.5
Pimephales promelas (Fathead minnow)	84.5
Oncorhynchus mykiss (Rainbow trout)	82.5
Salmo salar (Salmon)	82.5
Carassius auratus (Gold fish)	76.4
Danio rerio (Zebra fish)	67.1



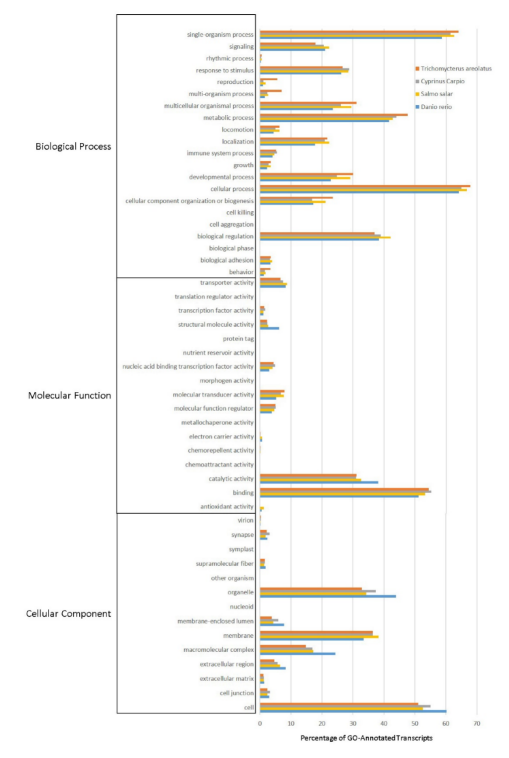
Chilean Catfish Coverage of Salmo salar and Danio rerio Proteins

**Figure 3. Transcript Coverage of Two Model Organisms.** Coverage of Salmo salar and Danio rerio predicted proteins by Trichomycterus areolatus predicted proteins. Predicted polypeptide sequences produced in this study were BLASTed against publically available non-redundant Salmo salar proteins (count = 112,089) and Danio rerio proteins (count = 81,931). The length of the local alignment region reported by the BLASTp algorithm was subsequently divided by the length of the query sequence. Compilation of these results indicated that a vast majority of *Trichomycterus areolatus* predicted protein sequences exhibited greater than 90% coverage of both Danio rerio (64.7%) and Salmo salar (68.9%) protein sequences, suggesting that the assembly produced a high degree of full-length transcripts.

Number of Proteins Covering Salmo salar

Percent Coverage of Salmo salar or Danio rerio Proteins

Number of Proteins Covering Danio rerio



**Figure 4. Gene Ontology (GO) Analysis of the Trichomycterus areolatus Transcriptome.** GO functional analysis was performed on assigned proteins in order to evaluate transcript function and the overall completeness of the isolated transcriptome. GO terms were given for each of the *T. areolatus* predicted proteins as well as the proteomes of *Salmo salar, Cyprinus carpio,* and *Danio rerio* (retrieved from NCBI). The distribution of protein functions closely match one another, suggesting the assembled transcriptome is complete.

Similarly, Figure 5 details the **KOG** transcriptomic analysis; this database allows the user to compare against a collection of seven eukaryote genomes of a diverse set (e.g. human, fly, parasite,

plant, worm, fungi) of model organisms [20]. These annotated proteins are organized by function into clusters of eukaryotic orthologous groups. Orthologs typically have similar functions among different organisms and serve as an effective means of identifying putative functions of gene products [20]. The KOG analysis of *T. areolatus* resulted in KOG groups classifying 81% of transcripts. Finally, **KEGG** analysis was performed to identify biologically relevant pathways of function for predicted protein products from the representative transcriptome [21]. *T. areolatus* was compared to *Danio rerio, Salmo salar* and *Cyprinus carpio* as is shown in Figure 6. Distribution of group numbers also appears consistent as related fish were evaluated. Overall, each annotation and grouping algorithm showed close association with *T. areolatus* and the related interrogated species, providing evidence of the completeness and consistency of the transcriptome.

#### **Differential Expression Analysis**

Table 4 details unique transcripts which were differentially regulated between male and female (sex determined by necropsy) *T. areolatus* samples. Non-inherited transcripts that are unique to individual organisms (e.g. MHC molecules via rearrangement) were excluded.

Among the most highly differentially regulated transcripts, the gene product of A2-macroglobulin (A2M) functions as a protease inhibitor and binds growth factors [22]. A2M, increased in the female by 96-fold, has been shown to increase endogenous production of estradiol resulting in increased follicular cell proliferation and oocyte maturation [22].

Vitellogenin is an egg yolk lipoprotein precursor that serves as an essential material for oocyte development [23, 24, 25]. Vitellogenin production is under estrogenic control, and was upregulated in the female *T. areolatus* 42 fold or more when compared to the male. The female liver (the primary vitellogenin production site) secretes the lipoprotein, which is then transported from the systemic circulation into oocytes within the developing ovary. Because vitellogenins are large lipoproteins, mobilization requires the microsomal triglyceride transfer protein for delivery to the oocyte. Microsomal triglyceride transfer protein is crucial in final yolk lipoprotein assembly [26], and is seen upregulated in the female by 38,604-fold.

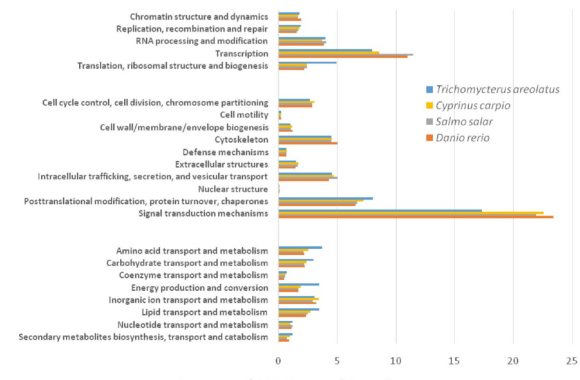




Figure 5. Eukaryotic Orthologous Groups (KOG) Characterization of Trichomycterus areolatus Transcripts. Putative transcript functions were assessed and transcriptome completeness was evaluated using KOG analysis. The Trichomycterus areolatus transcriptome and mRNA nucleotide entries from NCBI of Cyprinus carpio, Salmo salar, and Danio rerio were assigned KOG terms. The three transcriptomes have similar distributions, supporting the completeness of the Trichomycterus areolatus transcriptome.

**Table 4. Differentially Expressed Transcripts.** The resultant male and female *Trichomycterus areolatus* sequence files were interrogated to assess relative differential transcript expression. Unique transcripts, demonstrating changes greater than or equal to 10-fold, are identified by homology (if available) to known proteins, and only the most differentially expressed isoform is presented. Non-inherited transcripts that are unique to individual organisms (e.g. MHC molecules via rearrangement or similar immune transcripts) were excluded. Notably, many transcriptional differences are related to sex-specific expression.

Transcriptomic ID	Fold Change	Name	
Male differentially upregulated transcripts			
TRICH01_163992	15	parvalbumin beta-1	
TRICH01_133698	12	hemoglobin subunit beta-2	
TRICH01_83415	11	sperm acrosome membrane-associated protein 4	
TRICH01_57563	10	endonuclease domain-containing 1 protein	
Female differentially upregulated transcripts			
TRICH01_101761	38604	microsomal triglyceride transfer protein large subunit	
TRICH01_222696	96	complement C4	
TRICH01_180471	96	alpha-2-macroglobulin	
TRICH01_143497	56	3-hydroxyacyl-CoA dehydrogenase type-2	
TRICH01_54871	42	vitellogenin 4	
TRICH01_100604	38	pyruvate dehydrogenase phosphatase regulatory subunit	
TRICH01_142598	35	CD59	
TRICH01_51308	29	ribonucleoside-diphosphate reductase subunit M2	
TRICH01_208172	22	coiled-coil domain-containing protein 36	
TRICH01_51310	21	Jouberin	
TRICH01_100599	19	PEX5	
TRICH01_100601	17	histone-lysine N-methyltransferase ASH1L	
TRICH01_100600	16	A-kinase anchor protein	

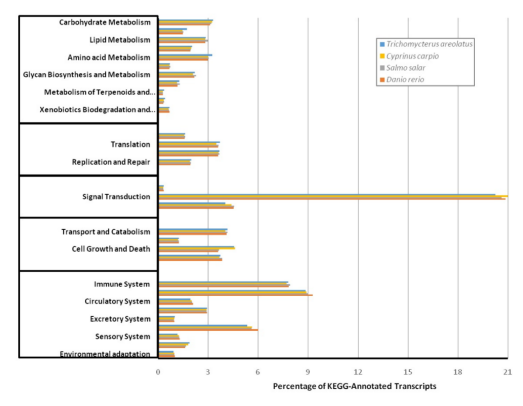


Figure 6. Kyoto Encyclopedia of Genes and Genomes (KEGG) Transcriptomic Analysis. KEGG analysis was performed to functionally describe transcript functions and evaluate transcriptome completeness. To serve as comparisons mRNA sequences for *Danio rerio*, *Salmo salar*, and *Cyprinus carpio* were retrieved from NCBI and characterized into KEGG pathways. The percent distribution shows a similar proportion among compared species indicating a complete transcriptome for *Trichomycterus areolatus*.

Throughout the process of vitellogenesis, the female organism undergoes metabolic changes, shifting energy usage, in order to support the growth of newly forming oocytes. Moreover, lipid metabolism is essential during ovarian development, including lipid storage, oxidation, and as previously mentioned, mobilization. Under some conditions such as starvation, the female may utilize stored lipids for transfer to the growing oocyte, or subsequent oxidation to meet dietary needs when intake is low [27]. increased expression (56-fold) The of 3-hydoxyacyl-CoA dehydrogenase suggests increased lipid breakdown and usage in the female most likely stimulated by increased energy requirements during vitellogenesis.

Similarly, the transcriptomic profile of the female demonstrates increased glycolytic activity, likely also to support the energy-intensive vitellogenesis. The pyruvate dehydrogenase complex is a highly regulated system that connects glycolysis with the TCA cycle. Activation of the complex is enhanced by pyruvate dehydrogenase phosphatase (PDP) [28]. The observed 38-fold increased expression of pyruvate dehydrogenase phosphatase would promote pyruvate progression through the TCA cycle and yield additional cellular energy in the form of ATP.

The complement system in fish is a primary component of their innate immune system and is suggested to be more beneficial to organism defense than that of mammalian systems [29]. Complement proteins are produced by the liver in an inactive form, where they are activated by proteolysis, ultimately leading to either opsonization, phagocytosis, or lysis of the pathogen. Vertebrate fish have multiple isoforms and isotypes of complement proteins, enabling the organism's immune system to recognize a wide range of pathogens, which is crucial due to slow lymphocyte proliferation, and limited antibody production and affinity in the fish immune system [30,31]. Notably, catfish as well as other teleost species, practice external fertilization which may expose the developing embryo and developing fish to waterborne threats, including pathogens. Maternal complement proteins are transferred from the female to the egg and serve to protect the embryos until the immune system and lymphoid organs are competent enough to protect the developing fish [32]. Thus, it is expected that maternal complement proteins, especially CD59 which acts as a stabilizer to prevent premature complement activation, would be Accordingly, the CD59 gene is upregulated. upregulated by 35-fold in the female. However, seasonal temperature and variations cause complement proteins to vary in expression,

accounting for some elevated complement proteins in the male [33].

Relative increases in male-associated transcripts are also presented in Table 4. Parvalbumins are calcium binding proteins found in white, fast twitch skeletal muscle of most fish species. In muscular tissues it acts to sequester calcium, accelerating muscle relaxation. High expression of parvalbumins can promote quicker muscle relaxation, however overexpression leads to smaller mitochondrial densities in slow twitch muscles [34]. More recently, the beta-1 parvalbumin isoform in carp seminal plasma has been characterized [35], and while the specific details of its function remain undefined, it is thought to play a role in sperm motility. Studies in carp have shown that sperm are not mobile without calcium, and initiation of motility is seen with an increase in intracellular calcium levels. The influx of calcium has been suggested to be the initiating factor of sperm motility, and therefore, the presence of parvalbumin as a calcium binding protein may play an essential role in fish sperm function. Additionally, studies in mammalian organisms have suggested parvalbumin regulation in calcium-mediated spermatogenesis and testosterone production [36]. Accordingly, the male transcriptome demonstrates a 15-fold increase in beta-1 parvalbumin expression.

Though many studies do not identify significant sex differences of hemoglobin levels in fish [37], Falahatkar et al. [38] discovered decreased hemoglobin concentrations and hematological changes in juvenile Acipenser stellatus that were treated with estradiol for 7-9 months. They predict estradiol may have an inhibitory effect on erythropoiesis (the production of red blood cells). Hemoglobin transcription levels in the male were found to be 12-fold upregulated compared to the female.

During mammalian fertilization, the acrosome surrounding the sperm head releases acrosomal hydrolases that enables the sperm and egg to combine [39]. The acrosomal-egg interaction, and release of hydrolytic enzymes, are thought to be stabilized by the outer acrosomal membrane matrix interactions, and the specificity of the binding interactions are essential in release of hydrolases [39]. The sperm acrosome membrane associated protein 4 is retained following the egg-sperm binding, localizing to the inner acrosomal membrane in human sperm, and may play a role in fertilization. Of note, teleost organisms differ in fertilization strategies: their sperm do not possess outer acrosomal membranes and therefore, do not undergo the same acrosomal reaction seen in mammals. Interestingly, recent catfish transcriptome

studies have identified a homolog of sperm acrosome membrane-associated protein 4 [40], though the specific function in catfish has not been elucidated [41]. A homolog to acrosome membrane associated protein 4 was observed in this male, expressed at 11-fold higher compared to the female.

#### **Consideration as a Sentinel Organism**

This transcriptome provides a resource to utilize T. areolatus as a sentinel organism or a "canary in the coal mine" for biological effects that may be experienced by local wildlife and nearby human populations. In North America, gene expression biomarkers have been readily applied to environmental monitoring for anthropogenic pollutants [42,43,44,45]. For example, fathead minnows are commonly used environmental sentinel in studies on agricultural runoff [46,47,48], waste water treatment plant effluent [49,50] and industrial waste effluent [51]. The growing reliance on transcriptomic tools in the field of environmental toxicology has been due to their increasing availability for non-model organisms as well as the mechanistic insight they provide for prediction of adverse outcomes at the whole organismal and possibly population level [52,53].

Chile is experiencing significant economic growth driven by agricultural and industrial development which puts considerable pressure on the freshwater resources across much of the country [54,55]. However, scientifically documented adverse impacts on water quality are largely absent, particularly with respect to biologic impacts on sentinel species. Thus, a need exists to develop environmental sentinel organisms suitable for study across the country. T. areolatus is an ideal candidate, as it can be found across much of the range of many rivers in Chile from the mountain watersheds to the low elevation, coastal regions of the country. Furthermore, the fish can be found associated with the benthos in streams where the water is merely a few centimeters in depth. Finally, the previously mentioned benthic behavior of the fish causes it to be intimately associated with sediments, therefore it is susceptible to contaminants that are affiliated with the water and sediment [56]. To provide a resource for the study of environmental systems associated with this organism, specific genes were identified for convenience (Table 5). These selected genes of interest will serve as markers of reproductive dysfunction and/or oxidative stress, in an upcoming manuscript currently in progress.

The availability of a complete transcriptome for *T. areolatus* provides a valuable resource relative to the development of this species as a sentinel organism. The unique niche that *T. areolatus* fills in Chile will allow it to be useful when considering water and sediment contamination.

**Table 5.** *Trichomycterus areolatus* **Environmental Sentinel Biomarkers.** Genes linked to endocrine disruption and/or oxidative stress were identified within the transcriptome assembly for convenience in developing *Trichomycterus areolatus* as an environmental sentinel organism. *Danio rerio* sequences were used as queries to BLAST the full assembly to identify putative homologs. Protein isoforms were differentiated based on query sequence annotation and bitscore.

Gene Name	Gene Symbol	Transcriptomic ID	Transcript Length (bp)	Query ID	Bit Score
Androgen Receptor	AR	TRICH01_58265	4455	NP_001076592.1	788
Aromatase	CYP19a1	TRICH01_14384	284	AAB65788.1	759
Aryl Hydrocarbon Receptor	AHR	TRICH01_166983	3366	NP_001019987.1	593
Aryl Hydrocarbon Receptor 2	AHR2	TRICH01_225654	2827	NP_571339.1	905
Cytochrome P450 1A1	CYP1a1	TRICH01_117246	2028	NP_571954.1	820
Estrogen Receptor Alpha	ESRa	TRICH01_56196	4380	AAK16740.1	729
Estrogen Receptor Beta 1	ESRb1	TRICH01_211240	4700	CAC93848.1	673
Estrogen Receptor Beta 2	ESRb2	TRICH01_95151	3578	CAC93849.1	796
Follicle Stimulating Receptor	FSHR	TRICH01_111037	3596	AAP33512.1	996
Forkhead Box L2	FOXL2	TRICH01_143206	1866	AAI16586.1	370
Heat Shock Protein 70	HSP70	TRICH01_53325	2625	AAF70445.1	1216
Heat Shock Protein 90 Alpha 1	HSP90a1	TRICH01_121146	2863	NP_571403.1	1292
Heat Shock Protein 90 Alpha 2	HSP90a2	TRICH01_121144	2926	AAI63166.1	1278
Metallothionein	MT	TRICH01_130427	554	AAS00513.1	53
Superoxide Dismutase	SOD	TRICH01_28552	2500	NP_571369.1	261
Thyroid Receptor Alpha	THRa	TRICH01_196629	2500	AAA99811.1	760
Thyroid Receptor Beta	THRb	TRICH01_20904	2406	AF109732_1	732
Vitellogenin 1	VTG1	TRICH01_101739	2851	AF406784_1	1224

## Methods

*Tissue Collection and RNA Preparation:* Whole fish were collected from the Choapa River basin in the Coquimbo region (region VII) of Chile in July 2015 under fishing authorization #2017 by the Chilean Subsecretary of Fisheries and Aquaculture. Specifically, a whole male and female with clear sexual differentiation were sampled from a downstream and upstream site respectively. The river sampling site coordinates are shown in Figure 2. This watershed is used intensively for agricultural production.

Fish samples were prepared and immediately submerged in RNA*later*<sup>®</sup> (Ambion) according to specific manufacturer recommendations to preserve RNA integrity. The samples were mechanically homogenized in the presence of Qiagen Lysis Buffer RLT; immediately following, a Qiagen RNeasy Mini Plus isolation kit was used to isolate and purify organism total RNA. The resultant RNA was quantified by Thermo Scientific<sup>™</sup> NanoDrop 2000c and verified for integrity with a bleach denaturing agarose electrophoresis gel [57]. Prior to sequencing, the purified RNA was stored at -80° Celsius with minimal handling and freeze-thawing cycles.

*High-throughput sequencing:* Prior to sequencing, a TruSeq® RNA Sample Preparation Kit was used to prepare the library for sequencing. An Illumina®

HiSeq2500 next generation sequencer was used to generate paired-end 101bp reads. Sequencing was performed at the University of Nebraska Medical Center sequencing core. The whole female and male tissue reads were used exclusively in transcriptome construction.

Data Processing and Assembly: The resulting sequence reads were first processed with FastQC [58] to evaluate sequence quality. Next, PRINSEQ was used to both trim reads by quality score, and apply a dusting technique [59]. Transcriptome assembly was performed de novo with TRINITY assembler [60, 61, 62]. The resultant initial transcriptome assembly was searched for predicted coding sequences at least 50 amino acids long using TransDecoder [61], BLASTed against NCBI Refseq metazoan protein sequences from March 2016 (retained if the bit score was 50 or higher), and filtered for microbial sequence contamination using the same BLAST [63] method. Noncoding RNA was removed based on homology to Rfam sequences of Danio rerio, and highly similar sequences were collapsed using CD-HIT to form the "representative transcriptome". This Transcriptome Shotgun Assembly project has been deposited at DDBJ/ENA/GenBank under the accession GEVC00000000. The version described in this paper is the first version, GEVC01000000.



Figure 1. Representative Photo of Organism-Trichomycterus areolatus organism photos of unspecified sex taken from two different locations in Chile, the Maule River (top) and Pangue River (bottom). Trichomycterus areolatus displays intimate contact with both sediment and the water column. Morphology and behavior is typical of fish who dwell in fast-moving streams. Photos were provided by courtesy of Pablo Reyes, Fundación Ictiológica (Chile) [70].

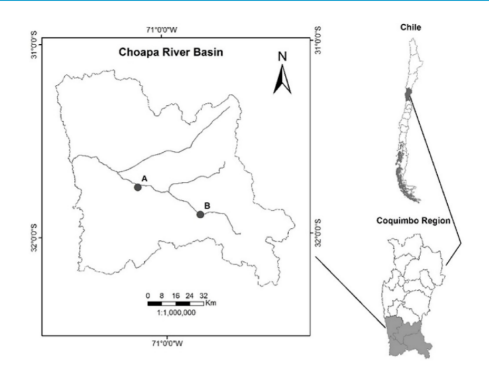


Figure 2. Choapa River Basin Tissue Sampling Sites-The male and female fish samples were collected from the downstream site "A" (altitude 243m, Lat; Lon: -31.749639; -71.160722) and upstream site "B" (altitude 792m, Lat; Lon: -31.89675; -70.783056) respectively. This river basin is proximate to heavy agricultural practice and downstream of heavy metals mining (e.g. copper). The stream itself is predominately supplied by glacial melt from the Andean mountains. Historically, this system drained into the Pacific Ocean but is becoming an isolated system due to the deleterious effects of climate change.

*Percent Identity Matrix:* Sequences acquired from the current assembly were aligned with sequences of previously published sequences from different species to evaluate phylogenic relationships within fish. Alignments were trimmed with Gblocks [64, 65], concatenated manually into a single ordered sequence, and aligned with ClustalW [66] to produce a percent identity matrix.

Transcript Coverage against Model Organisms: A FASTA file was compiled containing all protein sequences derived from the Τ. areolatus transcriptome; the sequences were BLASTed against all non-redundant Salmo salar and Danio rerio protein sequences retrieved from NCBI. The BLASTp algorithm was used to establish local alignments with E-values smaller than 1e-5. The sequences demonstrating the highest-scored alignments were kept. The length of each high-scoring alignment was subsequently compared to the overall length of the reference sequence to obtain the coverage, after which a count of the unique sequences present in a particular range of coverage was obtained through the Linux command line. The results were then compiled into Figure 3.

*Gene Ontology:* BLASTp was used to assign the top hit for the TransDecoded *T. areolatus* proteins and the proteomes of *Salmo salar, Cyprinus carpio,* and

Danio rerio (retrieved from NCBI January 29th, 2016) against the NCBI non-redundant database (GI list: Arabidopsis thaliana, Caenorhabditis elegans, Danio rerio, Dictyostelium discoideum, Drosophila melanogaster, Escherichia coli, Gallus gallus, Homo sapiens, Mus musculus, Rattus norvegicus, Saccharomyces cerevisiae, and Schizosaccharomyces pombe retrieved June 10th, 2016). The resulting file was scanned by BLAST2GO (version 2.8.0) [67] with the b2g\_feb16 GO database; once terms were assigned, level 2 GO ID terms were utilized for a comparison with all groups.

*KOG:* Analysis included translating publically available transcripts from chosen fish species using TransDecoder version 3.0 [61]. These results were aligned using RPS-BLAST (E-value of  $\leq$  1e-5) to the NCBI KOG database (version 3.14).

*KEGG:* The KEGG analysis was conducted by uploading organism transcriptomic sequences to the KEGG Automatic Annotation Server (KAAS) where they are processed by BLAST and GHOST comparisons against a database of KEGG genes. Ortholog assignments were returned and graphed to illustrate a transcriptome pathway comparison.

*Differential Expression Analysis:* Differential transcript expression analysis of *T. areolatus* transcript files were aligned to the assembled reference sequences. Reads were mapped with Bowtie [68], and

RSEM [69] was used to quantify differential expression values. Transcripts Per Kilobase Million (TPM) values were calculated to produce fold changes. Transcripts were then annotated by homology to NCBI Refseq proteins.

*Genes of Interest:* To identify genes related to endocrine and/or oxidative stress, query protein sequences from published sequences available on NCBI were obtained. Sequences from *Danio rerio* were used with BLASTp to find top hits in the putative coding sequence transcriptome (*T. areolatus*). Top hits were reviewed for the highest bit score (in combination with E-value) and included in Table 5. Protein isoforms were differentiated based off of chosen model organism annotation and could vary between query species (e.g. estrogen receptor alpha, beta).

## Acknowledgements

This work was supported by NIH grant P20GM103427 (P.H.D.). The University of Nebraska Medical Center Genome Sequencing Center provided RNA sequencing, and is additionally supported by NIH grant 1P30GM110768 and The Fred & Pamela Buffett Support Cancer Center Grant Additional financial support was P30CA036727. provided by the Fund for Undergraduate Scholarly Experiences (FUSE) and the Biomedical Research Training Program at the University of Nebraska. Jonathan Ali was partially supported by Robert G. Daugherty Water for Food Institute Graduate Assistantship, University of Nebraska, and the Center for Global Health and Development 2015 Summer Program, Research Fellowship University of Nebraska Medical Center. This publication's contents are the sole responsibility of the authors and do not necessarily represent the official views of the aforementioned funding agencies.

# **Competing Interests**

All investigators have declared that no competing interests existed.

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