Comparative transcriptome analysis of Nile tilapia (*Oreochromis niloticus*) in response to alkalinity stress

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**ABSTRACT.** To identify genes of potential importance to alkalinity tolerance, RNA sequencing (RNA-Seq) was performed to survey gill transcriptome profiles from freshwater (FW) and alkaline water (AW) exposed Nile tilapia (*Oreochromis niloticus*). A total of 22,724,036 (AW)/16,461,040 (FW) single-end reads were generated in which 20,304,348 (AW)/14,681,290 (FW) reads (90.0/89.72%) were aligned to the reference genome. Differential expression analysis revealed 302 up-regulated and 193 down-regulated genes between AW- and FW-exposed fish. These differentially expressed genes were enriched in several Gene Ontology (GO) terms related to "stress response", "heme binding", and "carbonate dehydratase activity". Meanwhile, significant KEGG pathways were enriched in energy metabolism, including nitrogen and sulfur metabolism. These results demonstrate the response of Nile tilapia exposed to alkaline-water and might provide valuable information to further understand the molecular mechanisms of adaptation of fish to alkaline environments.

**Key words:** Alkalinity stress; Nile tilapia; Transcriptome; Carbonate
INTRODUCTION

Alkaline water is usually characterized by relatively high pH and bicarbonate ($\text{HCO}_3^-$) and carbonate ($\text{CO}_3^{2-}$) concentrations. High-carbonate-alkalinity is considered a major stressor on fishes in alkaline water. High pH may inhibit ammonia excretion and increase $\text{CO}_2$ excretion in fishes living in alkaline water (Parra and Baldisserotto, 2007). Several fish species are able to survive in such harsh environments (Wilkie and Wood, 1996). For example, *Alcolapia grahami*, a well-studied alkaline-resistant fish, inhabits Lake Magadi (pH 10) Kenya (Seegers and Tichy, 1999), Lahontan cutthroat trout (*Oncorhynchus clarki henshawi*) inhabits Pyramid Lake (pH 9.4), Nevada (Coleman and Johnson, 1988), and Amur ide (*Leuciscus waleckii*) inhabits Dali Nor Lake (pH 8.25-9.6), China (Xu et al., 2013a,b). These fishes have evolved special strategies to overcome problems associated with ammonia excretion and acid-base regulation in alkaline environments (Wilkie and Wood, 1996; Bergman et al., 2003; Wang et al., 2003).

In contrast, alkalinity causes several abnormal behavioral, physiological, and biochemical reactions and even quick death in freshwater teleosts exposed to alkaline water (Yao et al., 2010). Freshwater fishes can endure alkalinity up to a certain extent (pH close to 8) depending on the species or individuals (Lei et al., 1985). Under such conditions, fishes must maintain their homeostasis by activating complex regulatory mechanisms including those for acid-base and osmotic control, nitrogen and respiratory metabolism, and immune system. Few studies have focused on molecular genetic responses stimulated by carbonate-alkalinity stress in fishes (Perry and Gilmour, 2006). Metabolic alkalosis induced by intra-arterial infusion of NaHCO$_3$ was reported to be accompanied by a marked increase in the Cl/HCO$_3^-$ exchanger mRNA hybridization signal in the rainbow trout gills (Sullivan et al., 1996). No significant changes were noticed in the expression of three lungfish genes, Na$^+$/HCO$_3^-$ cotransporter (NBC), V-type H$^+$-ATPase, and carbonic anhydrase, in gill and kidney tissue 4 and 8 h following a 1-h infusion of base (NaHCO$_3$) (Gilmour et al., 2007). DNA microarray was used for expression profiling in Japanese medaka (*Oryzias latipes*; Yao et al., 2012). After exposure to high carbonate-alkalinity solution containing NaHCO$_3$ and Na$_2$CO$_3$ (carbonate alkalinity = 30.4 ± 0.4 mM, pH = 9.2 ± 0.0) for 96 h, 512 genes were up-regulated and 501 genes were down-regulated in the gills. Most of the genes were associated with stress response, energy regulation pathways, immune system, iron, and osmotic, or acid-base acclimation; these genes included carbonic anhydrase, Na$^+$, K$^+$-ATPase, and V-H-ATPase. However, the previous studies were fragmentary and inconclusive about the complex molecular genetic response to alkalinity stress or about the mechanism of alkaline tolerance in fishes.

With the emergence of high-throughput (next-generation) sequencing, RNA sequencing (RNA-Seq) can be applied to transcriptome studies across the whole genome. Compared to the traditional cDNA microarrays, RNA-Seq is more sensitive in detecting all the expressed genes (Hale et al., 2011). Since gene expression generally changes very significant under abiotic stress compared to that under control conditions, RNA-Seq has been widely used to profile differential gene expression and pathways under certain environmental stress (Dugas et al., 2011; Teets et al., 2012). The Nile tilapia (*Oreochromis niloticus*), which inhabits freshwater, is one of the most important commercially cultured species. It is also a good model organism for studies on ionic and osmotic acclimation in euryhaline teleosts (Wang et al., 2009). Recently, selective breeding program on Nile tilapia has been initiated in China to improve their growth and alkaline tolerance. The strains with better performance could be important for the rural regions having large saline-alkaline open waters.
In the present study, we obtained the first high throughput data for the alkalinity stress responses in Nile tilapia using an Illumina sequencing platform. Transcriptome sequences from gills were assembled into contigs, function annotation and gene ontology (GO) analyses were performed, and candidate genes involved in alkalinity stress response were identified. These results provide a valuable resource for unveiling the mechanisms for alkaline tolerance, as well as for facilitating the genetic improvement of tilapias.

MATERIAL AND METHODS

Ethics statement

All experiments were performed according to the Guide for the Care and Use of Laboratory Animals of China. This study was approved by the Committee on the Ethics of Animal Experiments of Shanghai Ocean University.

Fish and carbonate-alkalinity exposure

Nile tilapias were obtained from fishery farm of the Shanghai Ocean University. They were maintained in a water circulation system at 26° ± 2°C under a 12-h light:12-h dark photoperiod. Carbonate-alkalinity solution, prepared using NaHCO₃ and de-chlorinated filtered tap water, was maintained in a plastic tank and aerated for 24 h before use. The carbonate alkalinity [expressed in millimolar (mM) concentration] was determined every day throughout the exposure period by acidimetric titrations. Exposure tests were conducted in 200-L tanks. Ninety percent solution in the test tanks was changed daily. The fish were exposed to freshwater (FW; carbonate alkalinity = 1.1 ± 0.1 mM, pH = 7.5 ± 0.2) and alkaline water (AW; carbonate alkalinity = 30.4 ± 0.4 mM, pH = 9.1 ± 0.2) for 96 h; each treatment had three tanks, each with 10 fishes. This alkalinity concentration was used because our previous study indicated that growth of tilapia was greatly affected at this concentration (Zhao et al. 2013). The fishes were not fed during the stress experiment.

Tissue sampling and RNA extraction

Based on a previous report (Lavado et al., 2013) about their importance under hyperosmotic and hypersaline conditions, gills of 12 fishes (6 from FW and 6 from AW) were collected and stored in liquid nitrogen for RNA isolation.

Total RNA was extracted from the gills using TRIZOL Kit (Invitrogen, Carlsbad, CA, USA) following manufacturer instructions. RNA samples were digested by DNase1 to remove possible genomic DNA contamination. Equal amounts of the RNA samples from FW and AW were then pooled together for cDNA synthesis and sequencing.

cDNA library construction and sequencing

RNA-Seq library preparation and sequencing was carried out by Oebiotech Lab (Shanghai, China). cDNA libraries were prepared from 2.5 μg total RNA using Illumina TruSeq RNA Sample Preparation Kit (Illumina, San diego, CA, USA) following the manufacturer protocol. The final library had an average 270-bp fragment size and a 400-ng final yield. The library was sequenced on
Illumina Hiseq2500 sequencing system. Adaptor sequences were trimmed and reads with low quality or length less than 10 were removed by the Trim Galore software.

Read mapping and differential expression analysis

The reads were processed and aligned with the *O. niloticus* reference genome (oni_ref_Orenil1.1, ftp: //ftp.ncbi.nlm.nih.gov/genomes/Oreochromis_niloticus/Assembled_chromosomes/seq/) using TopHat v1.0.12 (Trapnell et al., 2009). Alignment was performed with TopHat, which incorporates the Bowtie v0.11.3 algorithm (Langmead et al., 2009). TopHat initially removes a portion of reads based on the quality information accompanying each read and then maps the reads to the reference genome. It allows multiple alignments per read (up to 40 by default) and a maximum of 2 mismatches when mapping reads to the reference. The minimum/maximum intron length was set as 50/50,000 bp.

TopHat builds a database of potential splice junctions and confirms these by comparing the previously unmapped reads against the database of putative junctions. Default parameters for TopHat were used.

The aligned read files were processed by Cufflinks v0.8.0 (Trapnell et al., 2010). Reads were assembled into transcripts, their abundance was estimated, and tests for differential expression and regulation between different samples were performed. Cufflinks uses normalized RNA-Seq fragment counts to measure the relative abundance of transcripts.

The unit of measurement was fragments per kilobase of exon per million fragments mapped (FPKM). Confidence intervals for FPKM estimates were calculated using a Bayesian inference method (Jiang and Wong, 2009).

Functional annotation

GO annotations of *O. niloticus* genome were downloaded from Ensembl BioMart (http://asia.ensembl.org/biomart/martview/5addcfdd189de6499adfd439b3d7c206). The GO terms associated with transcriptome contigs were then obtained to describe the genes in the areas of biological processes, molecular functions, and cellular components.

KEGG pathways were assigned to the assembled contigs using the online KEGG Automatic Annotation Server (KAAS; http://www.genome.jp/kaas-bin/kaas_main; Moriya et al., 2007). The Bi-directional Best Hit (BBH) method was used to obtain KEGG Orthology (KO) assignment.

Quantitative reverse transcription-PCR (qRT-PCR)

To validate RNA-Seq results, 27 genes with high level of significance or important stress-responding functions were selected for qRT-PCR analysis with β-actin as the reference gene. Total RNA was reverse-transcribed with SuperScript III First-Strand Synthesis System for qRT-PCR (Takara, Dalian, China) and amplified with SYBR Green PCR Master Mix (Takara). β-actin was detected as the internal normalization control. The primers used for mRNA detection are listed in Table S1. Briefly, qRT-PCR was performed using MyiQ5 Real-time PCR Detection System (Bio-Rad, Hercules, CA, USA). The PCR conditions were: 50°C for 2 min, 95°C for 10 min, 40 cycles at 95°C for 15 s, and 60°C for 1 min. The PCR data were analyzed with the ABI 7500 SDS software. The comparative CT method (2^-ΔΔCt method) was used to analyze the expression of the target genes.
RESULTS

Transcriptome sequencing, assembly, and alignment

To obtain comprehensive understanding of the differences in expression between AW- and FW-exposed Nile tilapia, we performed deep sequencing of RNA samples from the fish gills. A total of 22,724,036 (AW)/16,461,040 (FW) single-reads were generated. After the removal of ambiguous nucleotides and low-quality sequences (Phred quality scores <20), a total of 22,553,326 (AW)/16,362,920 (FW) cleaned reads (98.6/99.0%) were obtained (Table 1). Of the total 20,304,348 (AW)/14,681,290 (FW) reads (90.0/89.72%) aligned to the reference genome, 19,612,054 (AW)/14,119,220 (FW) reads were unique (Table 2). All the RNA-Seq data generated in this study have been deposited in the NCBI SRA database (Accession Nos. SRR2074560 and SRR2074567).

Identification of differentially expressed genes

Differentially expressed genes (DEGs) were identified by calculating the ratio of FW versus AW FPKM values. Between AW- and FW-exposed fishes, 495 DEGs were identified with P value ≤0.05 and fold-change ≥2 (see Table S2). Among these genes, 302 were up-regulated and 193 were down-regulated.

We further examined 27 typical DEGs related to signal transduction, cellular stress response, acid-base and ion regulation, metabolism, immune response, and reproduction processes. Overall, the expression patterns of these genes were in agreement with the RNA-Seq and qRT-PCR analyses with minor differences observed in their expression levels (Figure 1). Only a single gene did not show consistency of expression in the two assays. Thus, these genes showing similar patterns of mRNA abundance in RNA-Seq analysis and qRT-PCR, validated the genome-wide expression profiling of gills in response to AW stress.

Functional annotation

GO-enrichment analyses were conducted on the DEGs; the top 30 significant GO terms in each group are shown in Figure 2. In response to AW stress, we observed significant gene
enrichment of several GO terms in gills that related to stress response including “cellular response to xenobiotic stimulus (GO: 0071466)” [3 genes]. The expression of more stimulus-related genes in AW-exposed fishes was consistent with the extreme environmental stress. GO-enrichment analysis indicated that the up-regulated genes were enriched in “heme binding (GO: 0020037)” [8 genes], “electron carrier activity (GO: 0009055)” [6 genes], “ATP biosynthetic process (GO: 0006754)” [2 genes], and “oxidoreductase activity (GO: 0016712)” [3 genes], while the down-regulated genes were enriched in “heme binding (GO: 0020037)” [1 gene], and “electron carrier activity (GO: 0009055)” [1 gene]. Genes in the molecular function category of “oxidoreductase activity” and “electron carrier activity” are widely studied and recognized to associate with oxidative stress and in adaptation to environmental stimuli in coupling with mitochondrial functions. It was also suggested that under severe osmotic and alkaline stress more energy was consumed than in the freshwater environment. The increased energy requirement leads to active proteolysis in the gill, which was observed as the DEGs enrichment in “proteolysis (GO: 0006508)”. In the present study, DEGs were also enriched in “carbonate dehydratase activity (GO: 0004089)” and “transport vesicle membrane (GO: 0030658)”, suggesting their important roles in regulating homeostasis of various substrates in response to the AW stress; this was consistent with previous reports on AW stress adaptation (see Table S3).

Figure 1. Differentially expressed genes validated by qRT-PCR. The white columns represent results of RNA-Seq, and the black columns indicate results of qRT-PCR. X-axis represents gene name and y-axis indicates log2 ratio [freshwater (FW)/alkaline water (AW)].
Additionally, KEGG analysis was performed as an alternative approach for functional categorization and annotation. Top 16 significant pathways are shown in Figure 3 (P < 0.05). Fifty-four DEGs were annotated with these KEGG pathways. Briefly, the most significant pathway was of energy metabolism, including nitrogen (3 DEGs) and sulfur metabolism (3 DEGs). Further, 33 DEGs were classified into metabolism, including major sub-groups of metabolic pathways (25 DEGs) and microbial metabolism in diverse environments (8 DEGs). DEGs grouped into endocrine system accounted for 11 pathways, including insulin signaling (7 DEGs) and adipocytokine signaling pathway (4 DEGs). Endocrine and metabolic diseases (type II diabetes mellitus), metabolism of other amino acids (selenocompound metabolism), excretory (aldosterone-regulated sodium reabsorption), and immune system (NOD-like receptor signaling pathway) groups contained 4, 2, 3, and 3 KEGG-annotated DEGs, respectively (see Table S4).

Figure 2. Gene ontology (GO) enrichment. Distribution of the GO categories assigned to the Oreochromis niloticus transcriptome. Transcripts were annotated in three categories: molecular functions (blue), cellular components (yellow), and biological processes (red). X-axis represents -log (P value) and y-axis indicates GO terms.
DISCUSSION

RNA-Seq and qRT-PCR data indicated that the expression of mitogen-activated protein kinase (MAPK), eukaryotic translation initiation factor 4E-binding protein 3 (EIF4EBP3), insulin receptor substrate 1-like, and insulin-like growth factor-binding protein 5-like in tilapia gill tissue was up-regulated under high alkalinity stress. The MAPK are key elements of protein phosphorylation cascades that integrate and amplify signals from osmosensors to activate appropriate downstream targets mediating physiological acclimation (Kultz, 2001). EIF4EBP3 belongs to the EIF4E-binding protein family, which regulates EIF4E activity by preventing its assembly into the EIF4F complex. It mediates the regulation of protein translation through the MAPK and mammalian target of rapamycin (mTOR) signal transduction pathways (Poulin et al., 1998). mTOR plays important roles in response to stress, including activation of autophagy (Jung et al., 2010) and modulation of

Figure 3. KEGG pathway analysis. Red columns represent significant enrichment pathways (P ≤ 0.05), and blue columns indicate insignificantly pathways (P > 0.05). X-axis represents KEGG pathway terms and y-axis indicates -log (P value).
protein synthesis (Sengupta et al., 2010). These responses can conserve energy and promote 
survival during prolonged periods of stress (Teets et al., 2012). Insulin-like growth factor I is 
identified in long-term systemic response. The effect of this hormone is fine-tuned at the cellular 
level via adjustment of the expression and membrane insertion of the corresponding hormone 
receptors (Yadav, 2010). The presence of MAPK, EIF4EBP3, and insulin-like growth factor I with 
high level of significance suggested that there should be a complex alkalinity signal transduction 
network in fishes for integration and amplification, and this network might include MAPK, mTOR 
signaling pathways, and systemic response. These signal transduction pathways control effector 
mechanisms responsible for acclimation to changes in the environmental alkalinity.

At circumneutral pH (pH 6-8), ammonia excretion is mainly dependent upon the suitable 
NH₃ partial pressure gradients between the blood and the unstirred boundary layers of the gill 
(Randall and Ip, 2006). In this model, hydration of CO₂ in the unstirred boundary layers of the gills 
provide H⁺ to trap NH₃ as NH₄⁺, as it diffuses across the branchial epithelium (Wright et al., 1989). 
This effectively creates a “sink” that continuously favors NH₃ diffusion. In alkaline water, there is 
less H⁺ available for transforming NH₃ into NH₄⁺. When freshwater fishes are exposed to alkali 
water, there is an immediate reduction in ammonia excretion rate and a corresponding increase in 
plasma ammonia concentration (Wright, 1993). High ammonia concentration may pose potentially 
lethal problems for fishes (Parra and Baldisserotto, 2007). In our study, increases in the expression 
of carbonic anhydrase (CA), ammonium transporter Rh-type C 2-like, and glutamine synthetase 
(GS) were found under high alkalinity stress. CA catalyzes the hydration of CO₂ to provide H⁺ and 
HCO₃⁻ (Fisher et al., 1996; Whitman et al., 2000). Ammonium transporter is electroneutral and 
may regulate transepithelial ammonia secretion (Han et al., 2006). GS plays an essential role in 
the metabolism of nitrogen by catalyzing the condensation of glutamate and ammonia to form 
glutamine, which is considered less or non-toxic (Eisenberg et al., 1987). Thus, we hypothesized 
that increased expression of CA, ammonium transporter, and GS can reduce toxicity of high 
ammonia concentration in fishes under alkalinity stress.

Many highly expressed proteins involved in the regulation of ion transport were also 
obscurred; these included sodium-coupled monocarboxylate transporter 1, voltage-dependent 
anion-selective channel protein 2-like, sodium/potassium-transporting ATPase subunit alpha-
1-like, solute carrier family 25 member 39, sodium/calcium exchanger 1-like, and ATP-sensitive 
inward rectifier potassium channel 1-like. We consider all these as candidate genes to maintain 
acid-base and ion balance directly during the high alkalinity stress. Similar to the previous studies, 
identification of genes encoding for oxidation-reduction such as cytochrome P450 suggests a high 
energy demand for tilapia acclimation to alkalinity stress. Furthermore, glutathione S-transferases, 
serine/threonine-protein kinase, and heat-shock 70-kDa proteins were rapidly up-regulated during 
alcalinity stress, suggesting that they were fundamentally important in the alkalinity stress response 
of fish. Heat-shock proteins target damaged proteins to the proteasome to prevent accumulation 
of dysfunctional proteins and to recycle peptides and amino acids (Goldberg, 2003), suggesting 
possibly high level of autophagy under alkalinity stress.

In conclusion, comparative transcriptome study conducted on Nile tilapia exposed 
to alkaline and fresh water identified a relatively large number of genes that displayed distinct 
differences in their expression in gills. We identified significant DEGs encoding important 
modulators of alkaline stress adaptation and tolerance, including carbonic anhydrases, glutamine 
synthetase, cytochrome P450. Further analysis revealed that several well-known signaling 
pathways and functional categories of genes were significantly enriched, including the pathways of
“mTOR signaling”, “EIF2 signaling”, and “superpathway of cholesterol biosynthesis” and functional categories of “response to stimulus”, “heme binding”, and “transporter activity”. Overall, this study demonstrated transcriptome changes in Nile tilapia in response to alkaline stress. The results set a foundation for further analyses on alkaline-responsive candidate genes, which would help in better understanding of teleost adaptation under extreme environmental stress and ultimately benefit future breeding for alkaline-tolerant fish strains.

Conflicts of interest

The authors declare no conflict of interest.

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Supplementary material

REFERENCES


