

Identification of single nucleotide polymorphisms associated
with growth traits in rainbow trout by RNA Sequencing

by
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ABSTRACT

The rainbow trout (*Oncorhynchus mykiss*) are a species of great importance for aquaculture. The efficiency and sustainability of commercial farms depends on strains with traits such as high muscle yield. These characteristics could be improved by selective breeding based on genetic differences between subgroups of *O. mykiss*. Single nucleotide polymorphisms (SNPs) account for the majority of these variations and can be studied via whole transcriptome sequencing. The characteristics of muscle mass, fat content, and flesh whiteness were measured in about a 100 families from two generations produced at the USDA/NCCCWA rainbow trout breeding program. RNA extracted from the samples was sorted by its correlation with the traits, sequenced by RNA-Seq, and mapped to a reference transcriptome. SNPs with a very high likelihood of trait influence were confirmed by Sequenom genotyping in a small population. Seven SNPs were confirmed as genetic markers for traits of interest.

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This bar graph shows the mapping statistics of the reads of high and low-ranked families. Reads were mapped to the previously generated reference transcriptome. An average of 8,600,000 reads were mapped per group.

FIGURE 3:

Table of Biological Significance of SNP Genes 20

This table shows the analytical results of the association between 41 validated nuclear SNPs and three traits (percent muscle mass, percent fat content, and percent color absorbance). The analysis was completed with PLINK genetic toolset in a population of 871 fish. These seven markers were each associated with at least one trait. Low false discovery rates indicate probable associations and are shaded deep red in this table. For each polymorphism, the major and minor alleles are designated and assigned color for contrast: yellow for adenine, green for guanine, blue for cytosine, and red for thymine. Each minor allele frequency [MAF] is also shown, with darker shading corresponding to greater frequency. Genotyping data for each SNP is given (number of individuals homozygous for major allele / number of heterozygous individuals / number of individuals homozygous for minor allele).

INTRODUCTION

RAINBOW TROUT

The rainbow trout (*Oncorhynchus mykiss*) are an iconic species which have greatly impacted humans. Native to the eastern coast of Russia and the western coast of North America, this valuable food animal was transported eastward and has been introduced to all continents besides Antarctica. They have become the most widely cultivated cold freshwater fish in the world (Palti *et al.*, 2011).

Traits such as large size and adaptability to artificial rearing and large size make rainbow trout ideal for such various purposes as food production, sport fishing, and research (Ma *et al.*, 2012, Pulcini *et al.*, 2012, Thorgaard *et al.*, 2002). They are larger than most other model fish species such as zebrafish and pufferfish, so greater quantities of specific organ tissue can be obtained from an individual (Ali *et al.*, 2014).

O. mykiss have been used as a model organism for such fields of research as ecology (Davidson, 2012), evolutionary biology (Taylor *et al.*, 2011), immunology (Nya and Austin, 2011), nutrition (Wong *et al.*, 2013), oncology (Tilton *et al.*, 2005), toxicology (Köllner *et al.*, 2002), and physiology (Palti *et al.*, 2011). More is known about the physiology and biology of rainbow trout than any other fish species (Salem *et al.*, 2015).

AQUACULTURE

Aquaculture remains the most sustainable solution to rising global demand for seafood products. Aquaculture is the fastest growing animal production sector and has expanded to accommodate demand that fisheries alone cannot meet (Lazzarotto *et al.*, 2015). The quota for aquaculture production worldwide increased from 38.5% to 48% from 2005 to 2011. Currently, aquaculture is experiencing a growth rate of 8%, the most for any global food production industry (Safari and Mehraban Sang Atash, 2015).

The success of rainbow trout aquaculture is due to innovations in management, better understanding of nutrition, advanced disease control, and improved strain genetics. Fast and efficient growth highly influence the profitability of food animal production. Through improvements in feed quality and uptake, growth rate and feed conversion ratio must be maximized and mortality must be minimized in order for aquaculture to remain profitable (Ramzani *et al.*, 2014). Furthermore, weight gain can also be influenced by pro-nutrients that may affect utilization of nutrients (Ramzani *et al.*, 2014). There is also interest in using terrestrial plant resources to replace or supplement aquatically sourced components in this salmonid's carnivorous diet, both to reduce cost and to minimize dependence on marine fishery resources (Lazzarotto *et al.*, 2015).

MARKER-ASSISTED SELECTION

Modifications to growth characteristics of commercial strains may help trout farmers bolster their production. While growth rate and adult size of *O. mykiss* can be increased by selection based on adult phenotype, this process leads to slow, modest gains

over several generations (Fishback, 2002, Salem *et al.*, 2015). The efficiency of fish farms cannot improve, through traditional means alone, by the rate at which need increases (NCCCWA, 2005). Muscle traits are complex, polygenic and difficult to improve by conventional selection. As an additional impediment to traditional selection, some traits such as fatty acid content cannot easily be determined in living individuals; for such traits, once an individual's desirability for breeding is ascertained, it is no longer able to breed.

The information obtained by an analysis of genetic differences can lead to quicker, more cost-effective improvement in commercial populations. If a characteristic in an individual's genome serves as a marker for improved adult phenotype, the individual can be chosen for breeding at any stage of development. Genetic marker assisted selection for desired phenotypes can be accomplished in fewer generations with better results than traditional selection, allowing for quicker improvement in commercial populations (Salem *et al.*, 2015).

GENOME AND TRANSCRIPTOME

Genetic research of *O. mykiss* is complicated by the size and complexity of its genome. An ancestor of the ray-finned fishes underwent an autotetraploid event, which effectively doubled its genome. An additional autotetraploid event occurred in the ancestor of the salmonids (Genet *et al.*, 2011). Due to crossovers between homologous chromosomes, modern rainbow trout still possess duplicate copies of most genes (Sakamoto and Danzmann, 2000, Palti *et al.*, 2011, Christensen *et al.*, 2013). In addition,

over half of all rainbow trout genes contains interspersed repeating sequences (Genet *et al.*, 2011; Salem *et al.*, 2015). The presence of many similar sequences hinders the development of a standardized genome reference sequence.

Alternatively, genes transcribed from different regions of the genome can be deduced from a transcriptome, the collection of RNA molecules synthesized in a population of cells. The muscle cell transcriptome of rainbow trout can reveal which variants in the DNA are correlated with important aquacultural traits by comparison with phenotypic data from adult individuals.

SINGLE NUCLEOTIDE POLYMORPHISMS

A single nucleotide polymorphism (SNP) is a variation in one particular base of the DNA in an individual in comparison either to a paired chromosome of the same individual or to a homologous chromosome in another individual from the same population or species. Though any of the 4 nucleotides may be present in a position, SNPs are typically bi-allelic. SNPs by common definition must be significant in a population, with the minor variation having a frequency of at least 1% (Vignal *et al.*, 2002). In rainbow trout, SNPs can potentially exist as variation between alleles in a single chromosome or between two paralogous chromosomes (Christensen *et al.*, 2013).

SNPs are a significant portion of the information that can be determined from comparison of individual genomes; potentially, hundreds of thousands of SNPs can be found in a population (Platzer 2013). Because they represent a large proportion of total genetic differences (as much as 90% among humans), they are ideal for genetic-assisted

selection of desired fish muscle traits (Pertea 2012). SNPs are also ubiquitous and readily scored, providing a relatively simple and inexpensive means to generate a vast amount of genotyping data (Vignal et al., 2002). SNPs which are proximal to or located within coding sequence are more likely to alter a completed protein and are the primary interest of this study.

QUALITY TRAITS

Traits that are beneficial for aquaculture include fast and efficient growth of individuals as well as pleasing appearance of fillets. The traits of percent muscle mass, percent fat mass, and flesh whiteness were selected for an operational definition of fish quality in this study.

As stated above, growth traits such as muscle yield and fat content are highly complex, as the adult size of an individual is determined by many genes, overall nutrition, and exposure and response to antigens. Diverse genes with many functions may potentially affect the adult phenotype.

Color is perceived as an indicator of overall quality and is a highly influential aspect of consumer approval (Safari and Mehraban Sang Atash, 2015). The characteristic pinkish coloration of rainbow trout fillets depends primarily on carotenoid pigments supplied in feed, as salmonids are unable to synthesize these pigments *de novo*. If carotenoids are not supplied in feed, the fillets will be a darker red (Hardy and Barrows, 2002). Carotenoids are absorbed enzymatically as lipophilic pigments and transmitted to the circulatory systems via carotenoid-carrying lipoprotein (Storebakken and No, 1992).

Carotenoids are deposited into the fillets, and only a negligible portion is remobilized in the blood of adult fish (Guillou and Choubert, 1992). However, color loss can occur in processed fillets during storage (Safari and Mehraban Sang Atash, 2015). Any gene that contributes to absorption, circulation, and retention of carotenoids could potentially affect this trait. Flesh whiteness, without the influence of dietary carotenoids, is heritable and was selected as a trait of interest for this study.

MATERIALS AND METHODS

ETHICS STATEMENT

Institutional Animal Care and Use Committee of The United States Department of Agriculture, National Center for Cool and Cold Water Aquaculture (Leetown, WV) specifically reviewed and approved all husbandry practices used in this study (IACUC approval #056).

TISSUE SAMPLES

Our samples were obtained from rainbow trout that were raised in Leetown, West Virginia at a USDA facility, the National Center for Cool and Cold Water Aquaculture. Extensive phenotypic data, including several size measurements and muscle and organ masses, were recorded for approximately 500 fish. Rainbow trout with phenotypic variations in traits were selected for this experiment from a hundred families and 2 generations (hatch years 2010 and 2012). Tissue samples were shipped on ice to MTSU and stored at -80°C. TRIzol[®] from Invitrogen (Carlsbad, CA) was utilized for total RNA isolation from each sample.

PREPARATION OF LIBRARIES AND SEQUENCING

Preparation of libraries and sequencing were performed at the University of Illinois. The TruSeq RNA Preparation Kit from Illumina (San Diego, CA) was used for construction of RNA-seq libraries. High-quality RNA (1 µg) was used as the source of mRNA. Strands were synthesized and 3'-end A-tailed double-stranded DNA was ligated

to indexed adaptors and amplified by PCR for 10 cycles. Libraries were Qubit-quantitated and pooled, followed by qPCR quantification to produce the maximum number of clusters in the flow cell. The libraries were sequenced from one end of the molecules on an Illumina HiSeq2000 to give reads with a length of 100 nucleotides.

Reads were mapped to the reference transcriptome assembled from 13 tissues of a single double-haploid fish using the default settings. Putative SNPs were called using the default parameters of CLC. Very strict parameters for SNPs detection were used (depth of coverage > 10; $P > 97\%$, quality score > 30) and followed by Fisher's exact test ($P < 0.05$).

ISOLATION OF DNA FOR GENOTYPING

To isolate DNA, 500 μL extraction buffer, 5 μL DTT (1M) and 6 μL Pro-K (10 mg/mL) were added to each fin clip in a 1.5 mL microtube. The tubes were stored at 37°C overnight on a rotary shaker, then spun for 2 min at 14,000rpm to separate the pigments. The supernatant was transferred to a new 1.5mL microtube and mixed with 200 μL 5M NaCl. These tubes were spun for 10 min at 14,000 rpm, and the supernatant was transferred to a new 2 mL screw-cap tube containing cold 100% ethanol (1 mL). The tubes were cooled on ice for 10 min or in -20°C freezer then spun for 10 min at 14,000 rpm and 4°C. The resulting pellets were aspirated. For a total of 2 times, the pellet was rinsed in 1 mL of 70% EtOH, spun for 5 min at 14,000 rpm and 4°C, and aspirated. Afterwards, the pellets were allowed to air dry for 20 min. Finally, 100 μL Low TE

(10mM Tris, pH8.0; 0.1mM EDTA) was added, and each tube was left overnight at 37°C. DNA extracts were analyzed for concentration and purity by spectrometer.

SNP MARKER VALIDATION AND GENOTYPING

Samples were sent to the University of Arizona Genetics Core for genotyping to validate SNPs. We expected the SNPs in the resulting genome to match many of the ones we had developed computationally from data obtained by RNA-Seq.

SNPs markers identified by RNA-Seq were validated by Sequenom genotyping at University of Arizona Genetics Core. Associations of markers with muscle mass, fat content, and flesh whiteness were assessed by genotyping 871 additional fish. SNPs associated with each phenotype were determined using PLINK, a whole genome association toolset. The detailed workflow for our pipeline to identify SNPs associated with muscle yield, fat content, and color is shown in Figure 1.

FIGURE 1

[Page 16 is saved as a separate document for color printing.]

RNA-SEQ ANALYSIS FOR SNP DETECTION

SNPs discovery was achieved using tissues collected from high-ranked families versus low-ranked families. For muscle mass and fat content, cDNA libraries from 8 different full-sibling families of high fat content (5 families) versus low fat content (3 families) and high muscle yield (4 families) versus low muscle yield (4 families) were used for RNA-Seq, with two families shared between muscle yield and fat content. For flesh whiteness, cDNA libraries from 9 different full-sib families of high color (5 families) versus low color (4 families) were used for RNA-Seq, with two families shared between fat content and flesh whiteness. Bar-coded libraries were prepared using single-end multiplexed sequencing protocol. A total of 21 RNA-Seq libraries were produced for all traits. Libraries were pooled and sequenced on a single Illumina HiSeq2000 flow cell.

In this study, important putative SNPs in cDNAs from high versus low ranked families were identified using the reads from RNA-Seq high rank and low rank families (each read 100 base pairs in length, an average of 123,000,000 reads per group). Reads were mapped to the previously generated reference transcriptome (Salem *et al.*, 2015). An average of 8,600,000 reads were mapped per group. Mapping statistics for the traits of muscle yield, fat content, and color are shown in Figure 2.

Putative SNPs identified by RNA-Seq were validated and genotyped by Sequenom genotyping platform. Associations of markers with muscle mass were further estimated by genotyping 871 individuals of the selected line at NCCCWA (hatch years 2010 and 2012).

FIGURE 2

[Page 18 is saved as a separate document for landscape and color printing.]

RESULTS

Probabilistic variant detection yielded 127,401 putative SNPs. Strict SNP filtration parameters (coverage > 20 reads; 4 reads to call variant; $P \geq 0.95$; forward/reverse balance < 0.05; quality score > 30) resulted with 31,419 putative SNPs. Fisher exact test on high rank and low rank families yielded 147 SNP ($P < 0.1$). These were submitted for genotyping and pared down to 41 validated SNPs considered for association study with phenotypes.

We measured muscle yield for individuals in 100 families from the USDA rainbow trout breeding program, and variations were compared with SNP allele frequencies. Out of 41 validated SNPs, the study identified 7 SNPs as potential genetic markers associated with quality traits. These results are given in Figure 3 and examined in the discussion.

FIGURE 3

[Page 20 is saved as a separate document for landscape and color printing.]

DISCUSSION

Each of the 7 SNPs identified in this study were synonymous mutations, meaning that each single base substitution does not directly alter the transcribed protein. Most of the SNPs were associated with housekeeping genes or genes with significant immune function. Housekeeping genes are always expressed and contribute to basic cell function, whereas tissue-specific genes are differentially expressed and contribute to cell specificity (Salem *et al.*, 2015).

VOLTAGE-DEPENDENT ANION-SELECTIVE CHANNEL PROTEIN 2

A guanine-to-thymine SNP with a frequency of 0.19 was correlated with muscle mass, fat content, and flesh whiteness. This SNP is associated with a gene for voltage-dependent anion-selective channel protein 2 (VDAC2).

VDAC2 is a protein found abundantly in the outer membrane of mitochondria as a constituent of mitochondrial porins. These channels mediate a significant portion of all molecular traffic to and from the mitochondria and are involved in regulation of mitochondrial apoptosis (Hoogenboom *et al.*, 2007). This gene could be a potential marker for immune response in rainbow trout (Long *et al.*, 2015).

Since the gene for VDAC2 is important for cell housekeeping and immune response, its associated SNP could conceivably have implications for many diverse traits.

CYTOPLASMIC FMR1-INTERACTING PROTEINS

Two SNPs were found associated genes with cytoplasmic FMR1-interacting proteins (CYFIP). An adenine-to-cytosine SNP with a frequency of 0.22 was correlated with muscle mass. This SNP is associated with a gene for CYFIP1. A thymine-to-cytosine SNP with a frequency of 0.10 was also correlated with muscle mass. This SNP is associated with a gene for CYFIP2.

CYFIP are very highly conserved proteins. CYFIP1 and CYFIP2 in humans are, respectively, 92% and 98% identical to the homologous genes in zebrafish. This suggests structural of functional importance for these genes, though neither protein contains a recognized functional domain or motif. More likely, these proteins function by binding to other proteins (Mayne *et al.*, 2004).

CYFIP is broadly expressed in the CNS (Schenck *et al.*, 2001), where it has implications for axon pathfinding and retinal lamination (Kobayashi *et al.*, 1998). It is also associated with both fibronectin-mediated primary CD4⁺ cell adhesion and Rac-1-mediated T cell adhesion (Mayne *et al.*, 2004). CYFIP2 is highly abundant in CD4⁺ cells in patients with multiple sclerosis, an autoimmune disorder (Pittman *et al.*, 2010).

Since CYFIP has an immune component, it is plausible that these 2 associated SNPs may affect immune response to parasites and other diseases, which in turn affect adult weight.

TAXILIN BETA MUSCLE-DERIVED 77-LIKE

An adenine-to-guanine SNP with a frequency of 0.31 was correlated with muscle mass. This SNP is associated with a gene for taxilin btw muscle-derived 77-like.

The taxilin family interacts with syntaxin proteins in the process of intracellular vesicle traffic, especially calcium-dependent exocytosis. They are expressed ubiquitously as housekeeping proteins (Nogami *et al.*, 2003; Nogami *et al.*, 2004). They are also involved with recycling transferrin receptors which have been endocytosed. (Sakane *et al.*, 2014).

Since this gene for a taxilin protein contributes to exocytosis and iron uptake, it is possible that an associated SNP could have implications for muscle development.

NUCLEOLIN ISOFORM X2

A thymine-to-cytosine SNP with a frequency of 0.50 was correlated with fat content and flesh whiteness. This SNP is associated with a gene for nucleoli isoform X2.

Nucleolin is a eukaryotic nucleolar phosphoprotein, and the most abundant nucleolar protein, excluding ribosomal proteins (Lapeyre *et al.*, 1997). It influences synthesis and maturation of ribosomes, from transcription of rDNA to assembly of ribosomal components (Ginisty *et al.*, 2000). Primarily located in dense fibrillar regions of the nucleolus, it is essential for general cell activity but is expressed in greater quantities in metabolically active cells (Bayne *et al.*, 2006).

For rainbow trout in particular, nucleolin production increases in cold seasons when fish are less metabolically active, serving to down-regulate production of both rDNA (Roger *et al.*, 2002) and rRNA (Alvarez *et al.*, 2003).

Since this gene for a nucleoli protein is involved in metabolic regulation, it appears likely that this associated SNP could affect fat storage in an adult rainbow trout. The association with color could be due to the altered composition of muscle cells as a result of metabolic down-regulation.

EUKARYOTIC TRANSLATION INITIATION FACTOR 3, SUBUNIT I

Two SNPs were found associated genes with Eukaryotic translation initiation factor 3, subunit I (EIF3I). An adenine-to-guanine SNP with a frequency of 0.27 was correlated with fat content. A adenine-to-cytosine SNP with a frequency of 0.30 was correlated with fat content and flesh whiteness.

EIF3I is one of 13 subunits of eukaryotic initiation factor (EIF) 3 (Graf *et al.*, 2014). EIFs support the initiation of translation in eukaryotes (Rezende *et al.*, 2014). EIF subunits may be overproduced in a carcinoma cell, resulting in larger size, increased proliferation, and enhanced cyclic progression (Cuchalová *et al.*, 2010).

Since EIFs have implications for many cellular processes, it is conceivable that this associated SNP could affect fat content and flesh whiteness.

CONCLUSION

RNA Sequencing was used in this current study to identify single nucleotide polymorphisms (SNPs) to serve as markers associated with the traits of muscle mass, fat content, and flesh whiteness in rainbow trout. We validated many of these presumptive SNPs. We attempted to identify candidate genes that could be responsible for phenotypic variations. Additionally, we investigated why each SNP may elicit its effect and identified genes that could be responsible for variations occurring among phenotypes.

This study included SNP identification in only the muscle tissue of adult rainbow trout. In future studies, we intend to examine other tissues and life stages and tissues for SNPs associated with desirable traits for aquaculture in order to improve the productivity and yield of commercially farmed strains.

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APPENDIX 1

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Scholars Week Abstract 2013

RNA-SEQ IDENTIFIES SNP GENETIC MARKERS AND DIFFERENTIAL GENE EXPRESSION ASSOCIATED WITH INCREASED MUSCLE YIELD IN RAINBOW TROUT

Ashlin Harris, Undergraduate, (Honors College); Rhett Layman, Graduate student; Mohamed Salem, Faculty; Mohamed Salem (Faculty Sponsor),

Background: Rainbow trout is an important food animal, making high muscle mass a desired trait. Muscle traits are complex, polygenic and difficult to improve by conventional selection. In addition, genetic marker assisted selection for desired phenotypes can be accomplished in fewer generations than traditional selection, allowing for quicker improvement in commercial populations. Single nucleotide polymorphisms (SNPs), which account for 90% of genetic differences among individuals, are ideal for marker assisted selection of desired phenotypic traits. The objective of this project is utilization of state-of-the-art RNA-Seq (whole-transcriptome sequencing) analysis to identify SNP genetic markers and characterize genes controlling muscle yield.

Results: Phenotypic variations in muscle yield were measured in 100 families from the USDA rainbow trout breeding program. Variations in muscle yield were correlated to transcriptome-wide SNP allele frequencies and to global patterns of gene expression in families showing extreme phenotypes (4 high-muscle yield families [50.9% of BW \pm 1.8] versus 4 low-muscle yield families [43.2% of BW \pm 2.1]). We identified 127,401 presumptive SNPs; from these we identified 143 SNPs in the “high” group and 96 SNPs in the “low” group, all with a false discovery rate (FDR) \leq 0.1. Currently, these putative SNP markers are being genotyped and evaluated for association with muscle yield in a ~500-fish panel. Differential gene expression between groups “high” and “low” identified 60 genes with a fold change \geq \pm 2 and a FDR \leq 0.1. Gene pathway analysis identified 239 SNPs and 60 differentially expressed genes responsible for amino acid and sugar metabolism as well as various metabolic pathways.

Conclusion: The study identified SNP markers and differentially expressed genes and gene pathways predictive of increased muscle yield in rainbow trout. These SNPs and differentially expressed genes can be used by aquaculture institutions, such as the USDA, to improve the efficiency of trait selection in rainbow trout.

Scholars Week Abstract 2014

**RNA SEQ IDENTIFIES SINGLE NUCLEOTIDE POLYMORPHISM (SNP)
ASSOCIATED WITH MUSCLE YIELD AND SHEAR FORCE OF FLESH IN
RAINBOW TROUT**

Bam Paneru, Graduate student, Biology; Ashlin Harris, Undergraduate, Biology; Rhett Layman, Graduate student, Biology; Timothy D. Leeds, NCCCWA; Jianbo Yao, UWV; Brett Kenney, UWV; Mohamed Salem (Faculty Sponsor), Biology

Rainbow trout (*Oncorhynchus mykiss*) is one of the top five sport fish species in North America and the second most widely cultivated fish species for food. Muscle content and shear force are two important characteristics of fish flesh which determine consumer satisfaction. Single nucleotide polymorphism (SNP) determines 90% of genetic difference between individuals and is utilized in genetic evaluation and selective breeding. Our previous studies using the RNA-Seq technique were successful in finding SNPs markers associated with growth traits in rainbow trout. In this study, RNA-Seq was used to identify SNPs associated with fish muscle yield and shear force. RNA was isolated from fish belonging to 100 different families. These families were classified into two groups: high muscle yield families (50.9% of BW \pm 1.8) and low muscle yield families (43.2% of BW \pm 2.1). RNA sample from all individuals belonging to each muscle yield group was pooled and sequenced by illumina platform. Sequencing reads from high muscle and low muscle families were mapped to a transcriptome reference to predict the SNPs associated with muscle content and shear force. Phenotypic variations in muscle yield were measured in these 100 families from the USDA rainbow trout breeding program. Predicted SNPs were validated by the sequenom genotyping technology. Out of validated SNPs, 42 SNPs were evaluated for association with muscle traits in 1000 fish from two generations. The study identified SNP (potential genetic markers) associated with shear force.

Scholars Week Abstract 2015

IDENTIFICATION OF SINGLE NUCLEOTIDE POLYMORPHISM (SNP) MARKERS FOR MUSCLE YIELD AND QUALITY IN RAINBOW TROUT USING RNA SEQ

Ali Ali, Graduate student, Biology; Bam Paneru, Graduate student, Biology; Rafet Al Tobasei, Graduate student, Biology; Mohamed Salem, Staff, Biology; Mohamed Salem (Faculty sponsor), Biology

Background: Rainbow trout (*Oncorhynchus mykiss*) is one of the favorite sport fish species in North America and the second most important fish species for aquaculture in the U.S. Muscle yield and quality traits are important factors of profitability for the food fish aquaculture industry. Most of the genetic variations are caused by Single nucleotide polymorphism (SNP). Our previous studies were successful in finding SNP markers associated with different traits in rainbow trout.

Methods: In the current study, whole-transcriptome sequencing (RNA-Seq) was used to detect SNPs associated with fish muscle yield, fat content, flesh whiteness and softness (shear force). Variations in these muscle quality traits were measured in about a 100 families from two generations produced at the USDA/NCCCWA rainbow trout breeding program. RNAs isolated from four full-sib families (5 fish per family), sorted as high or low for each of these traits, were separately sequenced using an illumine RNA-Seq technique. Sequences from different families were mapped to the reference genome/transcriptome to predict SNPs that are associated with each quality traits. Predicted SNPs were confirmed on a small set of fish using the Sequenom genotyping technology. Then, the validated SNPs were assessed for association with the aforementioned traits in approximately 1000 fish from two generations. Results: The study identified seven SNPs (potential genetic markers) associated with these quality traits. In addition, digital gene expression profiling identified a small number of differentially regulated protein coding genes and long non-coding RNAs (LncRNAs) that are associated with each quality attribute, perhaps, indicating that selective breeding have had a relatively low impact on modifying gene expression.