Genetic variation and identification of single nucleotide polymorphism of insulin-like growth factor-1 gene in *Tilapia guineensis*

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**Abstract.** Ukenye E, Megbowon I, Oguntade O, Oketoki T, Amusa O, Usman A, Sokenu B, Adeleke R, Joseph B, Omatah C. 2020. Genetic variation and identification of single nucleotide polymorphism of insulin-like growth factor-1 gene in *Tilapia guineensis*  
Biodiversitas 21: 5317-5321. *Tilapia guineensis* is an important cichlid species of West African coastal waters with good nutritional, economic, and aquaculture relevance. The knowledge of the genetic basis of variation in growth traits in Tilapia fish is of great importance to support genetic improvement in the context of aquaculture. In this study, regions of the *Tilapia guineensis* IGF-1 gene were sequenced, aligned and compared across populations to identify single nucleotide polymorphism and genetic diversity among four populations of *T. guineensis* in South-west Nigerian coastal waters. A total of thirty-four SNPs were identified across the studied populations and were detected in the forward reaction with twenty-two transversions and twelve transitions. Badagry population showed the highest genetic diversity with the highest molecular diversity indices; number of polymorphic sites, pairwise differences, number of segregating sites and nucleotide diversity while the least diverse population was Pepe. Analysis of molecular variance (AMOVA) revealed that genetic variation was mostly within populations. This finding provides more information regarding variation in insulin growth factor I of *T. guineensis* and will encourage association study for production traits that will inform useful selection marker for breeding program.

**Keywords:** Igf-1, genetic diversity, SNPs, *Tilapia guineensis*

**INTRODUCTION**

*Tilapia guineensis* is among the most exploited fish species of cichlids usually found in brackish water regions of West Africa and has good aquaculture potential. It has been successfully raised in ponds, enclosures, cages, and tanks and accepts the high density necessary for economical culture (Penna-Mendoza 2005). *T. guineensis* has continued to contribute significantly to the nutrition value and economic growth of many developing countries. In spite of these potentials, *T. guineensis* like other Tilapias is still faced with the problem of stunted growth due to its prolific nature. Information on the molecular basis of growth traits for growth and productivity enhancement of *T. guineensis* will increase aquaculture production. Consequently, genetic variation and single nucleotide polymorphism identification from candidate genes associated with growth as a molecular technique to improve its growth is crucial.

In view of the success of genetic improvement programs focused on the selection of faster-growing traits, identification of the genetic variation in candidate genes related to growth and other traits of economic importance is an essential step (Lupchinski Jr et al. 2011). According to Zhang et al. (2018), growth traits of animals are always of primary concern during breeding for its determinant economical value. Fish growth is an important productive trait in aquaculture and identification of single nucleotide polymorphism (SNP) for growth performance is considered as a key step during breeding. Growth traits of fishes and other farmed animals could possibly be improved by selective breeding under specific conditions (Zhang et al. 2018). The contribution of genetic variation as an essential molecular tool in selective breeding for growth enhancement and conservation programs in aquaculture and fisheries cannot be overemphasized. Therefore, the improvement of relatively slow growth rates in Tilapia as reported by many workers is considered necessary with a view to improve aquaculture production. Insulin-like growth factor I (IGF-1) is one of the candidate genes associated with growth in different species of fish and is considered a highly conserved and crucial growth-regulating gene in Tilapia species (De-Santis and Jerry 2007). It plays important role in fish growth, development and metabolism. It is also a mediator of the endocrine action of hormonal growth and is able to stimulate the synthesis of DNA and subsequent cell division in many cell types. Microsatellites and single nucleotide polymorphism are mainly molecular markers of choice for genetic studies in fishes (Sundaray et al. 2016). Due to the large numbers of SNPs in almost all populations of organisms for genome-wide association studies, SNPs are becoming the most abundant type of DNA sequence polymorphism with important role in genetic studies.
(Danzamann et al. 2016). According to Yáñez et al. (2015), SNP has been successfully used for the identification of candidate genes of traits and strains useful in aquaculture. A 7K SNP-chip has been used to study genetic differences between farmed and wild Atlantic salmon in Norway (Karlsson et al. 2011); SNPs were also used to assess the genetic impact of gene flow on wild populations of Atlantic salmon in the Gulf of Finland, Baltic Sea (Ozerov et al. 2016). Cuevas-Rodriguez et al. (2016) successfully identified genetic variation and single nucleotide polymorphism in candidate genes for growth in Oreochromis niloticus.

Although, major aquaculture efforts have been focused on Oreochromis niloticus particularly in the area of molecular mechanisms of some candidate genes associated with growth and selective breeding to improve its growth and overall productivity, but not much attention has been given to T. guineensis species commonly found around us in this research area. Therefore, identifying and validating genetic variation and single nucleotide polymorphisms (SNP) in IGF1 gene related to growth in T. guineensis necessitated this study.

MATERIAL AND METHODS

Fish samples collection and DNA extraction
Forty (40) broodstocks of T. guineensis belonging to four different populations (10 per location) namely; Pepe and Ugbonla from Ondo state, Badagry and Epe from Lagos state were used for the present study. The coordinates of the sampling locations are presented in Table 1. Phenol-chloroform protocol Sambrook and Russel (2001) was used to extract genomic DNA from the caudal fin tissue of the experimental fish. The purity check of extracted DNA was done using a Nano-drop spectrophotometer (Shimadzu corporation Japan, MODEL UV-1800, 2000 series) at absorbance of 260/280nm. After checking the quality and quantity of genomic DNA, it was diluted to a final concentration of 50-100 ng/μl in highly purified water for amplification.

Polymerase chain reaction amplification
To identify SNPs within the IGF-1 gene, the specific primer (ALT-IGF-1 forward-CTTGGACGAGTFAAAG GCAATG and ALT-IGF-1 reverse- GAATAAAC AAGCAGTGATAAGA) of 399bp was designed to amplify regions of the IGF-1gene. The Tilapia specific primer (ALT-IGF-1) was developed from the sequence available from public database and synthesized. DNA amplification was carried out by polymerase chain reaction (PCR) in a thermocycler (Eppendorf Master Cycler) at Inqaba biotechnology laboratory, South Africa. 50 ng DNA, 1.25U Taq DNA polymerase, 0.4 mM dNTPs and MgCl2 at the concentration 12.5mM and the annealing temperatures 55 °C were used for the primer-pair. PCR products were analyzed by gel electrophoresis on 1% agarose gel with ethidium bromide. This was followed by PCR product purification. The primer sequences, annealing temperature and ampiclon size of the amplified fragments are shown in Table 2.

DNA sequencing and SNP identification
Purified PCR products were bidirectional-sequenced in an automatic sequencer (ABI 3500XL, Genetic analyzer), at Inqaba biotechnology laboratory, South Africa. Immediately nucleotide sequences were obtained, the sequence of each fragment from each fish sample was aligned and SNP identification was done using Arlequin v. 3.5.2 software.

Sequence analyses
The aligned sequences were utilized to detect genetic diversity indices, single nucleotide polymorphisms, nucleotide diversity, polymorphic sites and mean number of pairwise differences using Arlequin software. The molecular diversity indices measured include number of haplotypes (H), number of segregating sites, average number of nucleotide differences (K), singleton variable site and parsimony-informative site for each population of T. guineensis. Percentage of variation among and within-population was also estimated using AMOVA.

RESULTS AND DISCUSSION

Polymorphisms detected
ALT-IGF-1 gene was used to investigate and validate genetic variation and single nucleotide polymorphism associated with growth traits in T. guineensis. Overall, 34 polymorphic sites with 34 SNPs were identified in four populations. Out of 34 SNPs, 22 were transversions and 12 were transitions; 33 were revealed in Badagry, 1 in Epe, non in Ugbonla and Pepe populations respectively. The highest number of polymorphism was detected in samples from Badagry with Ugbonla and Pepe samples showing no polymorphism. On the basis of the number of SNP loci in this gene fragment among the four T. guineensis populations, the most diversity was observed in the Badagry population while the least diversity was observed in Ugbonla and Pepe (Table 3). A total of 8 indel polymorphisms were detected across the four populations.

The value of expected heterozygosity ranged from 0.002 to 0.021 with the lowest value in Pepe population of T. guineensis (Table 3). The polymorphisms and the different population carrying it is shown in table 4. It is observed that Badagry population had 33 SNPs out of the 34 SNPs identified in these populations. However, Pepe and Ugbonla did not show any polymorphism. In table 5, Badagry population has the highest nucleotide diversity, mean number of pairwise differences and number of polymorphic sites. Overall, the populations recorded low nucleotide diversity. Badagry population showed higher

| Location | Latitude | Longitude | State |
|----------|----------|-----------|-------|
| Pepe     | N06°10'01.3" | E02°52.988" | Ondo |
| Ugbonla  | N06°08'31.1" | E004°47'39.8" | Ondo |
| Badagry  | N04°25.012" | E02°52.988" | Lagos |
| Epe      | N04°27.200" | E007°19.618" | Lagos |

Table 1. Geographical location of the sampling stations

| Location | Location | State |
|----------|----------|-------|
| Pepe     | N06°10'01.3" | Ondo |
| Ugbonla  | N06°08'31.1" | Ondo |
| Badagry  | N04°25.012" | Lagos |
| Epe      | N04°27.200" | Lagos |

Table 2. Primer sequences, annealing temperature and ampliclon size of the amplified fragments.
molecular diversity indices Theta S, haplotypes (H), average number of nucleotide differences (K) and the highest nucleotide percentage of cytosine and thymine compared to other populations studied. The AMOVA analysis showed that genetic variation (98%) was mostly within population (Table 6).

Table 2. Primer sequence, annealing temperature and amplicon size, used in amplification of IGF-1 gene.

| Primer  | Sequence                        | Annealing temperature (°C) | Amplicon size (bp) |
|---------|---------------------------------|---------------------------|-------------------|
| ALT-IGF-1 | F-CTTAGGASGAGTAGGAAGGCAATG   | 55                        | 399               |
|         | R-GTGAATACAAGCAAGGATAAG         |                           |                   |

Table 3. Genetic diversity and SNPs of IGF-1 gene in four Tilapia guineensis populations

| Population | A: average number of alleles, He: expected heterozygosity |
|------------|----------------------------------------------------------|
| Badagry    | 1.101±0.334, 0.021±0.072 | Transitions: 12, Transversions: 22, Indels: 3 |
| Ugonla     | 1.005±0.071, 0.001±0.020 | Transitions: 0, Transversions: 0, Indels: 2 |
| Epe        | 1.005±0.020, 0.001±0.020 | Transitions: 0, Transversions: 1, Indels: 1 |
| Pepe       | 1.005±0.071, 0.002±0.033 | Transitions: 0, Transversions: 0, Indels: 2 |

Table 4. Single nucleotide polymorphism of IGF-1 gene in Tilapia guineensis populations

| No | Position | SNP | Mutation | Population |
|----|----------|-----|----------|------------|
| 1  | 2        | T>G | Transversion | Badagry   |
| 2  | 14       | T>A | Transversion | Badagry   |
| 3  | 19       | A>C | Transversion | Badagry   |
| 4  | 21       | A>C | Transversion | Badagry   |
| 5  | 30       | A>T | Transversion | Badagry   |
| 6  | 34       | A>C | Transitions  | Badagry   |
| 7  | 35       | C>A | Transversion | Badagry   |
| 8  | 36       | T>C | Transitions  | Badagry   |
| 9  | 38       | T>C | Transitions  | Badagry   |
| 10 | 43       | A>T | Transversion | Badagry   |
| 11 | 56       | T>A | Transversion | Badagry   |
| 12 | 58       | G>C | Transversion | Badagry   |
| 13 | 62       | A>T | Transversion | Badagry   |
| 14 | 64       | T>C | Transitions  | Badagry   |
| 15 | 72       | C>A | Transitions  | Badagry   |
| 16 | 78       | A>T | Transversion | Badagry   |
| 17 | 89       | G>T | Transversion | Badagry   |
| 18 | 93       | T>A | Transversion | Badagry   |
| 19 | 115      | G>C | Transversion | Badagry   |
| 20 | 144      | T>A | Transversion | Badagry   |
| 21 | 154      | T>A | Transversion | Badagry   |
| 22 | 160      | T>A | Transversion | Badagry   |
| 23 | 199      | T>G | Transversion | Badagry   |
| 24 | 254      | T>A | Transversion | Badagry   |
| 25 | 310      | T>G | Transversion | Badagry   |
| 26 | 335      | C>T | Transitions  | Badagry   |
| 27 | 339      | T>C | Transitions  | Badagry   |
| 28 | 343      | A>C | Transversion | Badagry   |
| 29 | 356      | G>A | Transitions  | Badagry   |
| 30 | 1        | T>G | Transitions  | Badagry   |
| 31 | 4        | C>A | Transitions  | Badagry   |
| 32 | 20       | A>T | Transitions  | Badagry   |
| 33 | 3        | T>A | Transitions  | Badagry   |
| 34 | 4        | C>A | Transversion | Epe       |
The current study, sequence alignment ofresults
suggested a significant difference in body and Caudal Peduncle Depth of *T. guineensis* species from Badagry lagoon reported by Kuton and Adeniyi (2014). These results indicate that the values of each SNP site vary among the four *T. guineensis* populations. The different SNP sites might differentially affect *T. guineensis* growth traits and may be used as breeding selection markers for *T. guineensis*. It is noteworthy that association between single nucleotide polymorphisms detected and phenotypic traits was not investigated in the present study. Therefore, further study could be on the association between single nucleotide polymorphism and size of the specimen. The complete coding DNA sequence of the ALT-IGF-1 gene of the *T. guineensis* species shared 100% identity with *Oreochromis aureus* (GeneBank accession No.: XM031746329.1), *Maylandia zebra* (GeneBank accession No.: XM012918326.3), and *Pundamilia nyererei* (GeneBank accession No.: XM013914390.1). The results suggest that conservation of the ALT-IGF-1 gene is very high across different species. Anurodh et al. (2014) made a similar observation in sequence characterization of IGF1 gene in Indian goats. Similarly, Ugbonla and Pepe did not have any polymorphism, which might suggest that the region is relatively conserved in the populations, as this confirms little to no recombination or mutation within the gene region. This might affect the morphology in terms of body size and growth of the *T. guineensis* species in these populations. This finding is similar to that of Babatunde et al. (2016) who identified no polymorphism in IGF-1 gene of the chicken strains studied.

The low nucleotide diversity observed in all the studied populations of *T. guineensis* suggested low mutation rate in this region and indicated a relatively low level of genetic diversity, which may be a result of genetic drift and might have contributed to low growth. Hence, there is urgent need for conservation. This finding is in line with the report of Agbebi et al. (2016) who observed low nucleotide diversity in all the four Tilapia species studied. Babatunde et al. (2016) also reported a low level of nucleotide diversity in Nigerian locally adapted chickens. The SNPs polymorphism revealed a total of 22 transversions and 12 transitions; transversions are higher than transitions. This result suggests that transversions are more easily

### Table 5. Summary of molecular diversity Indices of *Tilapia guineensis* populations studied

| Indices               | Badagry | Ugbonla | Epe     | Pepe     |
|-----------------------|---------|---------|---------|----------|
| Nucleotide diversity  | 0.021±0.012 | 0.001±0.001 | 0.001±0.001 | 0.002±0.002 |
| No. Polymorphic sites | 33      | 0       | 1       | 0        |
| Mean no. pairwise difference | 8.37±4.240 | 0.56±0.494 | 0.56±0.494 | 7.67±3.907 |
| Theta (S)             | 11.665  | 0.000   | 0.353   | 0.000    |
| Theta (H)             | 2.403   | 0.542   | 0.781   | 2.122    |
| Theta (K)             | 3.302   | 1.052   | 1.052   | 1.956    |

### Table 6. Analysis of molecular variance (AMOVA)

| Source of variation | df | Sum of squares | Variance component | Percentage of variation |
|---------------------|----|----------------|--------------------|------------------------|
| Among population    | 3  | 4.4            | 0.017              | 1.24                   |
| Within population   | 36 | 46.9           | 1.303              | 98.76                  |
| Total               | 39 | 51.3           | 1.32               |                        |

### Discussion

In the current study, sequence alignment of *Tilapia guineensis* ALT-IGF-1 gene was used to investigate and validate genetic variation and single nucleotide polymorphism associated with growth traits in tilapia. Previous studies have confirmed this candidate gene approach as a strong technique for the identification of polymorphisms in genes that associate with economic traits (Hammer-Hansen et al. 2011). The present study revealed the existence of 34 SNPs detected in its coding and noncoding region across the four populations which is an indication of genetic polymorphism in *T. guineensis* populations. This finding is in accordance with the earlier reports of Cuevas-Rodriguez et al. (2016) on the novel single nucleotide polymorphisms in candidate genes for growth in tilapia (*Oreochromis niloticus*). They identified single nucleotide polymorphisms that will allow their association with growth traits in *O. niloticus*. Salem et al. (2012) also identified 22 SNPs associated with growth in rainbow trout. Roman (2018) stated that if an adequate number of SNPs is identified and used, mutations in genes associated with a trait can be detected and used in marker-assisted selection. The identified SNPs facilitate the finding of genotypes useful in aquaculture. Badagry population displayed the most diversity among the four populations followed by Epe population as demonstrated by the number of SNP loci and polymorphic sites. This suggests that there is diversity in IGF-1 gene in the Badagry and Epe populations of *T. guineensis*. A similar observation was made by Ukenye et al. (2016) on genetic diversity assessment of *T. guineensis* using microsatellite markers where Badagry population exhibited high genetic diversity. The observed polymorphisms in Badagry samples conform to a significant difference in body and Caudal Peduncle Depth of *T. guineensis* species from Badagry lagoon reported by Kuton and Adeniyi (2014). These results indicate that the values of each SNP site vary among the four *T. guineensis* populations. The different SNP sites might differentially affect *T. guineensis* growth traits and may be used as breeding selection markers for *T. guineensis*. It is noteworthy that association between single nucleotide polymorphisms detected and phenotypic traits was not investigated in the present study. Therefore, further study could be on the association between single nucleotide polymorphism and size of the specimen. The complete coding DNA sequence of the ALT-IGF-1 gene of the *T. guineensis* species shared 100% identity with *Oreochromis aureus* (GeneBank accession No.: XM031746329.1), *Maylandia zebra* (GeneBank accession No.: XM012918326.3), and *Pundamilia nyererei* (GeneBank accession No.: XM013914390.1). The results suggest that conservation of the ALT-IGF-1 gene is very high across different species. Anurodh et al. (2014) made a similar observation in sequence characterization of IGF1 gene in Indian goats. Similarly, Ugbonla and Pepe did not have any polymorphism, which might suggest that the region is relatively conserved in the populations, as this confirms little to no recombination or mutation within the gene region. This might affect the morphology in terms of body size and growth of the *T. guineensis* species in these populations. This finding is similar to that of Babatunde et al. (2016) who identified no polymorphism in IGF-1 gene of the chicken strains studied.

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substituted than transitions. This finding is contrary to other previous research (Vera et al. 2011).

According to Tian et al. (2014), genetic heterozygosity indicates the proportion of the group with site heterozygotes at some loci. The average heterozygosity of loci reflects the level of variation in the genetic structure, and variability is directly correlated with heterozygosity and the ability to adapt to the environment. Among the four populations, the average expected heterozygosity and molecular diversity indices of the Badagry population were higher than others reflecting greater diversity for Badagry population. Thus, Badagry population represents the best population of *T. guineensis* associated with growth traits in the present study. The least diversity that was observed in Ugbonla and Pepe populations based on number of SNPs may be due to inbreeding or founder effect. Analysis of molecular variance (AMOVA) revealed that genetic variation was mostly within populations suggesting that bulk of the variation seems to be accounted for by differences within populations.

In conclusion, Badagry population that showed greater diversity may represent the best population of *T. guineensis* connected with growth traits among the populations studied. The polymorphisms identified in the current study indicated some level of genetic variability hence; SNPs identified in the *T. guineensis* ALT-IGF-1 gene contributed to the genetic variation in *T. guineensis* populations and should be further investigated through association study to detect their production potential which may inform useful selection marker for breeding program.

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