The Structural Annotations of The Mir-122 Non-Coding RNA from The Tilapia Fish (Oreochromis niloticus)

Arli Aditya Parikesit1*, Imron Imron2*, Rizky Nurdiansyah1, David Agustriawan1

1Department of Bioinformatics, School of Life Sciences, Indonesia International Institute for Life Sciences, Jakarta, Indonesia
2Research Institute for Fish Breeding, Subang, Indonesia

ARTICLE INFO

Article history:
Received April 21, 2021
Received in revised form September 27, 2021
Accepted October 11, 2021

KEYWORDS: tilapia, Oreochromis niloticus, miRNA, transcriptomics, RNAcentral, molecular modeling,

ABSTRACT

Tilapia (Oreochromis niloticus) is an important fisheries commodity. Scientific efforts have been done to increase its quality. One of them is staging a premium diet such as a fat-enriched diet. The transcriptomics approach is able to provide the signatures of the diet outcomes by observing the micro(mi)RNA signature in transcriptional regulation. Hence, it was found that the availability of mir-122 is essential in the regulation of a high-fat diet in tilapia. However, this transcriptomics signature is lacking structural annotations and the complete interaction annotations with its silencing(si)RNA. RNAcentral website was navigated for the latest annotation of mir-122 from tilapia and other species as a comparison. MEGA X was employed to comprehend the miRNA evolutionary repertoire. The RNA secondary structure prediction tools from the Vienna RNA package and the RNA tertiary structure prediction tools from simRNA and modeRNA are secured with default parameters. The HNADOCK tools were leveraged to observe the interaction between mir-122 and its siRNA. The post-processing was conducted with the Chimera visualization tool. The secondary and tertiary structure of the mir-122 and its siRNA could be elucidated, docked, and visualized. In this end, further effort to develop a comprehensive molecular breeding tool could be secured with the structural annotation information.

1. Introduction

Tilapia fish, or Oreochromis niloticus, is considered an important aquaculture product worldwide that originated from Africa and spread into more than 90 countries (De Silva et al. 2004). As it could live in a temperature range of 8 to 42°C in freshwater and hot spring, tropical countries such as Southeast Asia could serve as their habitat (FAO 2019a). It was projected that Tilapia fish production already reaching 6.3 million tonnes, and south-east Asian countries will play an important part there (FAO 2019b). The breeding techniques of Tilapia fish are not considered complicated. They may be carried out in the aquaponic system as well as in pond-based or hapa-based breeding systems (Bhujel et al. 2001; Fessehaye et al. 2006; Storey 2017). As male Tilapia significantly make more profit due to its bigger size and less time or energy for breeding. Various techniques such as methyltestosterone and YY super male were generated to breed male-only offspring (Towers 2013). In this respect, with the advance of molecular biology, more sophisticated techniques were introduced as breeding molecular markers based on genomics and proteomics technology. There are currently molecular markers categorized into cytoplasmic, dominant, and codominant types, and the most common molecular markers in use are based on mitochondrial (mt)DNA and simple sequence repeat (SSR) signatures (Amoussou et al. 2019). Moreover, sex-specific DNA markers have already been developed for facilitating the production of the genetically male Tilapia with different determination loci (Chen et al. 2018). A more specific molecular marker, such as Prolactin I microsatellite, has been developed with satisfactory performance to assist breeding (Chi 2014).

However, it is already known that the transcriptomics approach could eventually detect the activities of the genes that are beyond the coverage of the genomics and proteomics-based
technology (Dong and Chen 2013). In this regard, transcriptomics-based markers are already widely utilized for molecular medicine and agriculture research (Parikesit et al. 2017; Anurogo et al. 2019). Moreover, aquatic-based biomarkers have been developed for fish breeding as well (Collí-Dulà et al. 2016). Bioinformatics has played important role in developing the transcriptomics markers with specialized machine-learning-based tools (Parikesit 2018b). Molecular modeling and simulation-based approaches such as RNA structure elucidation and molecular docking methods are widely utilized to annotate the transcriptomics biomarkers (Parikesit 2018a). Chou in Chou (2004) stated that the current approach in biomarkers development always considers bioinformatics tools in assisting the wet laboratory efforts. The important biomarker that is currently becoming a focal point in transcriptomics development is the non-coding(nc)RNA (Mattick 2005). ncRNA is gaining importance because ncRNA could become a viable alternative toward the standard proteomics one (Amaral and Mattick 2008). One of the important types of small ncRNA is micro(mi)RNA, which is approximately 22 base pair long and explicitly inhibit gene function (Kozomara and Griffiths-Jones 2011). miRNA plays an important role in various metabolic and physiological functions of the organism such as the immune system, digestive pathway, and neurological function (Griffiths-Jones 2006). Another type of small ncRNA that is very similar to miRNA is the silencing(si)RNA or the interference RNA (RNAi). The difference between both types is the biogenesis of miRNA involves the unwinding process, while the siRNA involves the cleavage process. Moreover, both of them have a distinct secondary structure (Ui-Tei 2016). The interesting part is the availability of siRNA to inhibit miRNA activities could trigger developmental defects in various organisms (Chapman et al. 2004). Moreover, artificial miRNA has already been utilized as siRNA expression vector for human disease, and this is a strong argument that actually there is a feasible interaction between miRNA and siRNA (Boudreau et al. 2009). In this end, the utilization of siRNA is crucial because it is already proven be able to knock down genes and transcripts that are responsible for human and animal disease.

In this regard, in order to provide more biomarkers alternatives for fish breeding, a specific transcriptomics approach should be devised for Tilapia fish markers (Li et al. 2015). Moreover, as one of the problems of Tilapia’s breeding is their inclination towards a high-fat diet, a specific biomarker should be leveraged. Hence, it is found that high-fat diets in Tilapia will induce the upregulation of mir-122 transcriptomics-based biomarkers that are highly correlated with lipid metabolism and fat deposition (Qiang et al. 2018). Moreover, mir-122 also plays an important role in regulating stress response (Qiang et al. 2017). Although the metabolic pathway of the mir-122 has already been elucidated, its structural annotations are still not determined yet due to the inherent difficulty in RNA crystallography experiment (Holbrook and Kim 1997; KE 2004; Reyes et al. 2009). The absence of structural information may hamper the development of more fine-grained biomarkers in the future. The information on the interaction between mir-122 of Tilapia with other biomarkers is also absent because the database for ncRNA in animals is currently unavailable (Bonnici et al. 2018). Thus, in this end, the objective of this research is to determine the in silico structural annotations of the mir-122 Tilapia biomarker in order to shed light on its molecular interactions. It is expected that the elucidated structure will pave the way to find more information about the fish breeding transcriptomics-based pathway, especially their molecular interactions.

2. Materials and Methods

The pipeline for this experiment was developed based upon previous venues that have been improved significantly (Parikesit and Anurogo 2018; Parikesit et al. 2018). The computer used is a MacBook Pro® Laptop (13-inch, Late 2011) with 16 GB RAM, Intel HD Graphics 3000®, and 500 GB SATA disk for securing good performance in molecular computation (Cense 1989; Baro and Hughes 1991; Smith 1995; Oliver et al. 2019). The pipeline is mainly comprised of two parts, namely the sequence and structural analyses.

2.1. Sequence Analysis

The first step would be retrieving mir-122 sequences from RNAcentral database with respect to representative species of Mus musculus, Bos Taurus, Danio rerio, Oreochromis niloticus (Tilapia fish), Xenopus tropicalis, Homo sapiens, and Gallus gallus that inspired from UCSC genome browser annotated genomes(Speir etal. 2015; TheRNAcentralConsortium 2017). Moreover, in order to determine the possible
common ancestry of the mir-122 sequence in those species, MEGA X software was employed to generate the phylogenetic tree. The evolutionary history was inferred by using the Maximum Likelihood method and Jukes-Cantor model with Gamma distribution. All positions with less than 95% site coverage were eliminated, i.e., fewer than 5% alignment gaps, missing data, and ambiguous bases were allowed at any position (partial deletion option). The tree was replicated with 1,000 bootstraps.

2.2. Structural Analysis

2.2.1. 2D Structural Analysis

The earliest step is to determine the conserved 2-D structure for the mir-122 from the representative species with RNAalifold and determine the secondary structure of mir-122 from Tilapia fish with RNAfold (Gruber et al. 2008; Lorenz et al. 2011). The resulted ‘dot-bracket notation’ data of secondary structure will be utilized for predicting the transition state of the mir-122 stability with Barrier server (Flamm et al. 2002; Wolfinger et al. 2004). Thus, the important step in this part is the elucidation of the siRNA for mir-122 Tilapia with RNAxs (Tafer et al. 2008).

2.2.2. 3D Structural Analysis

Then, the three-dimensional structure prediction of both mir-122 and its siRNA for Tilapia was conducted with modeRNA and simRNA, both of them for generating predictions with homology modeling and ab initio methods respectively (Magnus et al. 2016; Rother et al. 2011; Magdalena Rother et al. 2011). In this end, after obtaining the protein databank (PDB) file of the models, the HNADock software was utilized to predict the docking possibility of mir-122 Tilapia with its respective siRNA (He et al. 2019). All software was employed by using the default parameters, and the HNADock software has utilized a specific scoring function for RNA-RNA interaction.

3. Results

3.1. Sequence Analysis

Based upon the retrieved sequences from the RNACENTRAL database, the representative of each species from mammalian, aves, amphibia, and actinopterygii classes is shown in the Table 1.

However, in order to obtain information on the common ancestor of those genes and their clustering, the phylogenetics tree method was devised in Figure 1. It is expected that the correct tree will eventually provide the correct structural annotations.

The tree shows that, unsurprisingly, the mir-122 of Homo sapiens, Bos Taurus, and Mus musculus are clustered together as the mammalian class. While Xenopus tropicalis and Gallus gallus purposely are in one distinct cluster. It is inferred that the tree is biologically correct because Danio rerio and Tilapia fish are clustered together as part of Actinopterygii class. In this end, as the tree is biologically correct, the sequence annotation could be leveraged to search for the conserved structure of the ncRNA.

3.2. Structural Analysis

3.2.1. 2D Structural Analysis

As the phylogenetic tree already validated the biological inference, it is the next logical step to progress further with the structural annotation pipeline. Hence, the following 2D structural conservation alignment was executed in order to comprehend the structure diversity among different organisms. As shown in Figure 2A, the base pair probability is very high, and it shows the plausibility that the structure of all sampled organisms mir-122 actually derived from a single entity. Hence, the Tilapia mir-122 structure in Figure 2B closely resembles the conserved structure with only a subtle variation. The calculation of the predicted free energy of the conserved structure is -54.10 kcal/mol, while the predicted structure of mir-122 Tilapia is -33.30 kcal/mol.

The trajectory of the RNA structures could be elucidated with fine resolution with the barrier server application. The Tilapia mir-122’s transition state could be observed in Figure 3. As a standard feature, exactly 10,070,026 structures were predicted by the RNAsubopt applets in the server that formed a way in an energy range of 35.4 kcal/mol above the minimum free energy.

<table>
<thead>
<tr>
<th>Table 1. The retrieved mir-122 sequences from the representatives species. These are the model species for genome annotation research</th>
</tr>
</thead>
<tbody>
<tr>
<td>Rncentral ID</td>
</tr>
<tr>
<td>----------------------------------</td>
</tr>
<tr>
<td>URS000006F971C</td>
</tr>
<tr>
<td>URS000015B1DD</td>
</tr>
<tr>
<td>URS000067AD20</td>
</tr>
<tr>
<td>URS000071B92F</td>
</tr>
<tr>
<td>URS0000914C02</td>
</tr>
<tr>
<td>URS000065381A</td>
</tr>
<tr>
<td>URS0000646248</td>
</tr>
</tbody>
</table>
Figure 1. Molecular phylogenetic analysis of mir-122 sequences by maximum likelihood method (JC + G model; 1,000 bootstrap). The number depicts the evolutionary distance between one node and the others. The identifiers of nodes are RNAcentral ID, species name, and RNA identifier.

Figure 2. (A) RNAalifold, (B) RNAfold Tilapia. The scale of base-pair probability is using a color legend that starts from probability 0 (dark purple), inferring that the structure is not feasible, and probability 1 (dark red) is the most feasible one. The total domination of the red color in the structures means that the structures are plausible, as the scale inferring to the probability of the structure existence.
specific molecular simulation method is necessary to examine the interactions of those structures. The utilized standard for this experiment is the docking between chains A and B of the r(GGUCACAGCCC)2 crystal structure (PDB ID: 1KD5) shown in Figure 6. Based on the computational validation result, the structure in Figure 6A is predicted as the best one due to its existence in the 1st rank among 100 annotated structures in PDB format. Moreover, the 1st rank structure also gives the ligand RMSD (Å) of 0.60. Figure 6B shows the top 10 structures superimposed in one Figure. The designated validation method of this research made it clear that the prediction pipeline is actually worked to observe the molecular interaction based on real wet laboratory experiment settings.

Figure 4 shows the RNAxs application output of the siRNA that downregulates the mir-122 of Tilapia fish. The taken siRNA is the most favorable design compared with others based on the thermodynamics approach. Thus, the elucidated 2D ncRNA designs are prepared for the next step of the 3D design pipeline.

3.2.2. 3D Structural Analysis

Hence, as shown in Figure 5, the molecular visualization of both the mir-122 and its siRNA could be done in a fine-grained manner. More importantly, the visualization shows that the hydrogen bonds of the RNA molecules could be elucidated. It is an important indicator that the internal structure of the functional groups could sustain the availability of the bonding and proves that there is a high probability if the structures indeed exist. In this end, a more specific molecular simulation method is necessary to examine the interactions of those structures.
Figure 4. The RNAxs output of the mir-122 siRNA. The black plot refers to the accessibility of the target sequence and the red plot refers to the accessibility of siRNA sequence. If approaching zero or touching the base of X-axis, the probability of accessibility for the sequence is zero. If it is farther from zero, the probability will be higher. BLAST iteration could be executed in the low-right of the window to check the homology of the query in the NCBI genbank.

Figure 5. Visualization by UCSF Chimera for the de novo models of (A) mir-122 and (B) siRNA mir-122. The hydrogen bonds that sealed the double-helix structures were depicted in blue planar rectangles. The pentagonal shapes represent the bases.
In this regard, the visualization of the molecular docking result between mir-122 and its respective siRNA could be observed in Figure 7. The docking experiment result will be compared with the validated standard as stated beforehand.

Based on the computational result, the structure in Figure 6A is predicted as the best one due to its existence in the 1st rank among 100 annotated structures in PDB format. Moreover, the 1st rank structure also gives the ligand RMSD (Å) value

Figure 6. The molecular docking of the crystal structures of r(GGUCACAGCCC)2 (PDB ID: 1KD5) that have already been validated by the wet experiment. The brown-colored RNA is the 1KD5 chain A, while the other colors are the 1KD5 chain B in different conformations. (A) The best model docking, (B) the 10 best models docking in the cumulative structure

Figure 7. Molecular docking of mir-122 with its respective siRNA. The brown-colored RNA is the mir-122 Tilapia, while the other colors are the siRNA in different conformations (A) the best model docking, (B) the 10 best models docking in the cumulative structures
of 50.49. Figure 6B shows the top 10 structures superimposed in one Figure. As the ligand RMSD is way above the accepted cut off the 5 Å, it is predicted that the binding between the siRNA and mir-122 will not be permanent and prone to be unbound in a very short period of time. However, the docking study implies that the interaction between the mir-122 and siRNA is indeed happening. Further studies are needed to observe the implication of the siRNA unbinding in this docking system.

4. Discussion

The phylogenetic tree as depicted in Figure 1 is employed because it is necessary to observe whether the biological clustering information of the sequences is in accordance with the established tree in the NCBI taxonomy database (Federhen 2012). The spontaneous values of both structures in the Figure 2 are the strong argument for the feasibility of those structures in nature. Hence, the structural crystallography will eventually be complicated to do (Anderson-Lee et al. 2016).

Hence, as wet laboratory experiment for determining the RNA structure transition is inherently feasible with kinetic isotope effects and single-molecule fluorescence spectroscopy that is not within reach in many laboratories, a more feasible approach is developed for this regard (Chen et al. 2000; Russell et al. 2002; Bartley et al. 2003). Thus, the computational environment is the fittest approach in order to observe this phenomenon. The total amount of structure in the Figure 3 is almost similar to the one that was detected in other experiments with other species and biomarkers (Parikesit 2018c). Hence, it could be inferred that the typical miRNA structures pool could always be detected in a certain amount.

As a plausible approach, silencing mir-122 could be a way to downregulating that transcriptomics biomarker and regulating the fat metabolism (Esau et al. 2006). In this regard, utilization of silencing (si)RNA to silence the mir-122 is a viable option for the down-regulation process (Girard et al. 2008). Based on the homology search in the template of the modeRNA database, the Homology models of mir-122 and its siRNA of Tilapia are not found. It suggests that the RNA structure of the mir-122 Tilapia could be considered novel and distinct as it is not similar to any 3D structure in the homology model database (Figure 4). In this regard, the only method that could be relied on predicting the 3D structure is with the de novo prediction that works by constructing the structure based upon the thermodynamics and kinetics parameters, as depicted in Figure 5 (Das and Baker 2007).

The most viable method to observe the interaction between the RNA molecules is by utilizing molecular docking tools. The docking method is currently very common to observe the interaction between protein-protein, and protein-ligand (Morris and Lim-Wilby 2008). However, until recently, the utilization of this method to comprehend RNA-RNA interaction is not doable due to the unavailability of the scoring function. Hence, the current protocol already provides RNA-RNA scoring function within the HNADOCK application, and the validation of this method is already leveraged by running the structural samples from the wet experiments (He et al. 2019). Based upon the accepted consensus, the RMSD cut-off below 5 Å for results in both Figure 6 and 7 is considered a good binding mode (He et al. 2019). The utilized standard for this experiment of the r(GGUCACAGCCC)2 in the Figure 6 was taken from the wet laboratory experiment. However, the experiment only provided the sequences, and the structure is predicted with our own in-house pipeline.

The application of structural bioinformatics in RNA modeling was previously focused mainly to human biomedical studies. Examples are in the domain of Cancer and infectious disease studies, for devising biomarkers and therapeutics (Agustriawan et al. 2021; Ivan et al. 2021; Parikesit and Nurdiansyah 2020). Comprehensive molecular simulation pipeline for this effort has already been established, albeit with an extensive demand for computational power and specific parameters and scoring functions (Ivan et al. 2020). Up to now, similar studies for the molecular aspect of the fisheries sciences are considered very limited and barely exist in the structural bioinformatics community. As the focal point of structural bioinformatics application in humans are to elicit biomedical interests, most of the designs are catered for diagnostics, drugs, and vaccines. Similar demand is very scarce in fisheries studies, except for diagnostics. It is an exception because DNA barcoding in fish is already an established method and is widely deployed in this field (Fitrian and Madduppa 2020; Ayu et al. 2021). However, as the field of agriculture
Acknowledgements

The author would like to thank the Research and Community Service Department (LPPM) of the Indonesia International Institute for Life Sciences (i3L) and Research Institute for Fish Breeding (BRPI) for their heartfelt support and providing the facilities for this research. Thanks also goes to Prof. Rosalba Giugno from the Department of Computer Science, University of Verona, Italy, for providing information on how to leverage the arena-ldb database, and Andamar Pradipita, M.A. from the i3L Community Language Center for his excellent proofreading of this manuscript.

References


