



Influence of pesticides pollution on mitochondrial 12S rRNA gene in cultured *Mugil capito* from different regions at Kafr El-Shiekh governorate, Egypt

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ABSTRACT

The aim of this study was to evaluate the impacts of pesticides pollution on the sequence of mitochondrial 12S rRNA in liver tissues of *Mugil capito* that obtained from three different regions at Kafr El Sheikh Governorate, Egypt. The results revealed ten SNPs between three sequences of different regions after alignment of these sequences submitted under accession numbers (MF817450.1), (MG210582.1) and (MG210583.1) with sequence with accession number (KU681005.1). Overall, the pesticides pollution has able to happen mutation in the mitochondrial gene.

Keywords: *Mugil capito*, pesticides, SNPs, 12S rRNA gene sequence.

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1. INTRODUCTION

Recently, there are found that human health might be exposed to hazards from fish consumption due to persistent organic pollutant residues in fish tissues like pesticides (Greco et al., 2010). These pesticides are utilized in agricultural applications and finally diffused into natural water affecting aquatic living. Hence, fish was a good biomarker for aquatic environment pollution with pesticides (Lakra and Nagpure, 2009).

Nowadays, with advanced in biological science field, some studies have focused on determining the side effect of pesticides residues on gene level. There had been detected that it had able to interact with genetic material causing either inactivation or

stimulation for nucleic acid replication (Farid and El-Sayed, 2015). Therefore, gene expression can play a crucial role to know the state of fish in the surrounding environment as well as early detection for any changes in the expression of some key genes (Schulte, 2001; Steiner and Anderson, 2000).

Mitochondrial 12S rRNA has proven to be a useful molecular marker for better conservation and management of the endangered species (Siddappa et al., 2013). The rRNA plays a primarily role in protein synthesis. It is, therefore, of particular importance that the structure is taken into account when rRNA genes are aligned, especially when the alignments are intended for phylogenetic analyses (Wang and Lee,

2002). In order to quantify the extent of compensatory mutations that occurs within 12S rRNA sequences and to discuss the phylogenetic implication of the degree of constraint on compensatory mutations as well as, the nucleotide composition and the patterns of nucleotide substitutions.

Therefore, these changes are very important tool for diagnosis of chemical pollution and pesticides in fish. The aim of this study was to assess the impact of pesticides pollution on mitochondrial 12S rRNA gene sequence in cultured *Mugil capito*.

2. Materials and methods

Fish sampling

Total number of 150 *Mugil capito* was collected from the study regions (Al-Hamol, Al Riad and Sidi Salem) fish farms. Liver and dorsal musculature were excised, packed and kept at -80°C.

RNA extraction

Total RNA was extracted from muscle and liver tissues of *Mugil capito* using TRIzol method according to manufacturer's instructions (Chomczynski, 1993). The quantity of RNA was evaluated by using Nano-drop spectrophotometer and purity by OD₂₆₀/OD₂₈₀ nm absorption ratio 1.8:2.0. The obtained total RNA was treated with DNase to remove any contamination with genomic DNA.

PCR reaction and program

The isolated cDNA were amplified using script RT-PCR two-step kit following the manufacturer protocol (Jena Bioscience, Germany). The used primers obtained using primer3 tool <http://primer3.ut.ee/cgi-bin/primer3/>. PCR amplification was carried out by SensoQuest (Labcyler, Germany) using a 50 µl of polymerase chain reaction mixture contained: 2µl of cDNA used as template separately, 5 µl of 10x Hot Start Buffer complete, 1µl of dNTP Mix, 0.25µl of

Hot Start Pol, 1µl of forward primer (0.1-0.5 µM), 1µl of reverse primer (0.1-0.5 µM) and 39.75µl of RNase free Water. The final reaction mixture was placed in a thermal cycler and the PCR program was carried out by initial denaturation at 94 °C for 2 min followed by 40 cycles of 94 °C for 30 sec for DNA denaturation, annealing temperatures as seen in (Table 1) for 30 sec, extension at 72 °C for 1 min and final extension at 72 °C for 10 min then were held at 4 °C. Amplified PCR product was analyzed by electrophoresis in 2 % agarose gel stained with Ethidium bromide using 50 bp DNA ladder (Thermo Scientific, USA), then visualized under UV Trans-illuminator.

Cloning and Gene sequencing

The purified PCR fragment, almost 250bp, was ligated in pGEM®-T Easy Vector Systems (Promega, Germany) according to the manufactures. The competent cells were prepared and transformed according to (Inoue et al., 1990). The white colonies were picked up from LB/Amp/Xgal plates and inoculated on LB/Amp broth media (Sigma-Aldrich, Germany). Then it incubated overnight at 33°C for stabilizing the plasmid inside the transformed cells with shaking. The plasmid was isolated according to alkaline method (Birnboim and Doly, 1979). The purified plasmids were analysed by electrophoresis in 1.5 % agarose gel using 50bp DNA Ladder (Thermo Scientific, USA) to confirm the recombinant plasmids. The recombinant plasmids were sent to MacroGen Company (South Korea) for sequencing by ABI 3730XL DNA sequencer (Applied Biosystem, USA).

Phylogenetic analysis

The obtained sequence for 12S rRNA was analyzed with VecScreen tool for vector contamination (<http://www.ncbi.nlm.nih.gov/tools/vecscreen/>). The obtained sequence from mitochondrion of *Mugil capito* was registered

at NCBI database under accession numbers: MF817450.1, MG210582.1 and MG210583.1 (<http://www.ncbi.nlm.nih.gov>).

3. RESULTS

Identification of 12S rRNA SNPs

This study revealed effect of pesticides pollution on mitochondrial 12S rRNA gene in *Mugil capito* that obtain from three different regions within kafr elshiekh government. After sending this gene for sequencing, the results elucidated presence variations among

the three different sequences. These sequences were submitted into genebank with accession numbers (MF817450.1) (MG210582.1) (MG210583.1). They aligned to highly similar sequence on genebank with accession number (KU681005.1) reported that there were ten single nucleotide polymorphisms (SNPs). Analysis of those sequences revealed SNPs C29G, G34T, A41T, G57T, A65C, G66C, G79A, C82G, G206T, T220C, and A226G as shown as in Fig 1.

Table 1: The used primers sequences in the current study.

Gene of interest	Primer sequences (5' - 3')	Ta (°C)	amplification size (bp)	Reference
12S rRNA	F: CCCACTATGCTCAGCCCTAA R: CCTGGCGTTTTGGGTCATAC	53	250	Current study

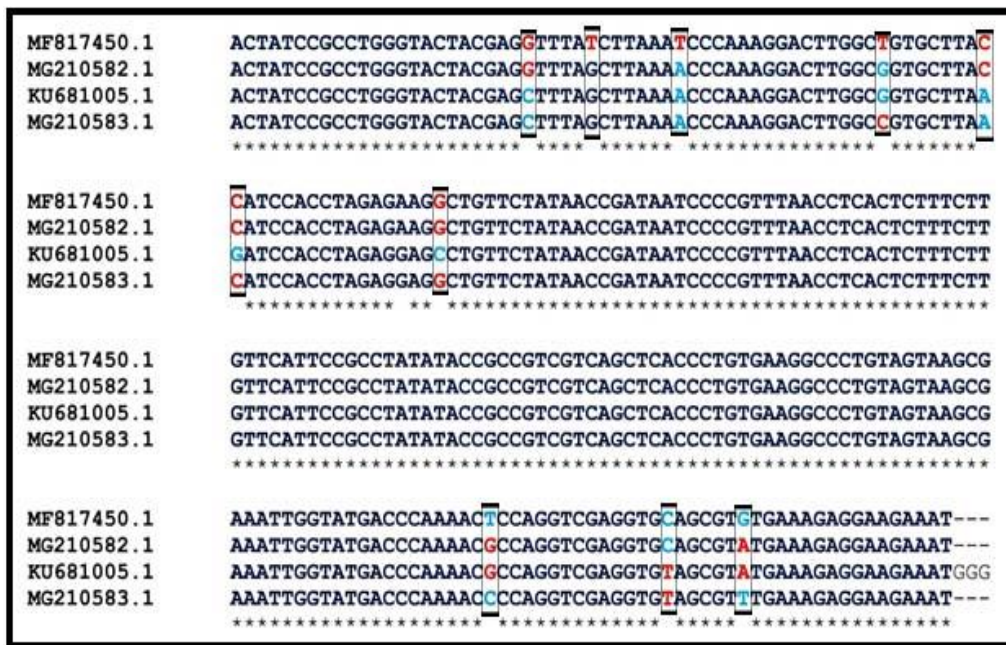


Fig.1. Single nucleotide polymorphism (SNP) image of *Mugil capito* in different study regions (Al Hamol, Al Riad and Sidi-Salem) with accession number (MF817450.1) (MG210582.1) (MG210583.1) respectively current study produced by Jalview, version 2.10.1 software.

4. DISCUSSION

The present study is an extension to a previous work of Abd El Megid *et al.*, (2018) (under publishing) where the authors did survey on pesticide residue in the studied areas (Al Hamol, Al Riad and Sidi-Salem) for the fish farm water and tissue levels of cultured *Mugil capito* in the same fish farms belonging to the study areas. The obtained results revealed that Delta-BHC compound was the dominant organochlorine with concentration ranging from 150 to 450ng/L. In addition to other pyrethroids residues were detected in the ponds water. It was also found that the highest pesticides residue was detected at Al Hamol and Al Riad fish farms and the lowest at Sidi-Salem fish farms. But fortunately; the pesticides residues in fish tissues were below the permissible limits.

Mitochondrial genome is extremely necessary for a life of almost eukaryotic that encoded into 13 proteins, two ribosomal rRNAs and 22 tRNAs that are required for mitochondria protein synthesis (Yang *et al.*, 2014). The results of the nucleotide sequence organization alignment are similar to fish species. In consistence, we have a tendency to found high similarity (95% identity) in nucleotide sequences of 12S rRNA between *Mugil capito* in current study with that submitted under accession number (KU681005.1). To our data, this can be the primary study to spot ten loci of mitochondrial 12S rRNA in *Mugil capito* in Egypt. The sequences of those loci were submitted into GenBank databases with accession number (MF817450.1), (MG210582.1) and (MG210583.1). Analysis of those sequences revealed SNPs C29G, G34T, A41T, G57T, A65C, G66C, G79A, C82G, G206T, T220C, and A226G. These

SNPs may be attributable to presence of pesticides within the study regions led to some mutations in the sequences of 12S rRNA. Surprisingly, variations of SNPs among the various study regions were correlated with the degree of pollution in every region. It could be concluded that, IGF1 and CYP1A genes can be used as sensitive biomarker for assessment contamination in all stages of fish. In addition to the new marker represented by, 12SrRNA sequence analysis.

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