

Mitochondrial transfer RNA mutations and male infertility.

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Abstract

Background: Mitochondrial DNA (MtDNA) mutations affecting flagellar movement are often causes of sperm dysmotility. Among these mutations, mitochondrial transfer RNA (mt-tRNA) is the hot spot. However, the role of mt-tRNA mutations in male infertility remains heated debates.

Purpose: To evaluate the roles of mt-tRNA mutations in male infertility.

Methods: To address this problem, we reassessed 5 recent reported mutations: *tRNAHis* C12187A, *tRNAThr* G15928A, *tRNAArg* T10463C, *tRNALys* A8312G and *tRNAAsp* T7572C, in clinical expression of asthenozoospermia. We first performed database searches for these mutations; in addition, the phylogenetic approach was performed to determine the Conservation Index (CI) of these mutations between different species. We also used the bioinformatics tool to predict the secondary structure of mt-tRNAs with and without these mutations. In addition, the pathogenicity scoring system was used to assess the status of each mt-tRNA mutation.

Results: We found that only *tRNALys* A8312G mutation could be regarded as “possibly pathogenic”, whereas other mutations should be classified as “neutral polymorphisms”.

Conclusions: This is the first report dealing with the association between mt-tRNA mutations and male infertility, thus, our study provided novel insight into the molecular basis underlying mt-tRNA mutations in male infertility.

Keywords: mt-tRNA, Mutations, Male infertility, Pathogenic.

Accepted on March 27, 2017

Introduction

Infertility affects about 10-15% of all couples attempting pregnancy, and male infertility accounts for half of them. The suspected causes of male infertility include sperm motility disturbances, Y-chromosome microdeletions, chromosomal abnormalities, and hormonal or receptor disorders [1]. Mutation and depletion of mitochondrial DNA (mtDNA) are associated with poor motility and diminished fertility of human sperm [2]. MtDNA is linked to electron transport system and is thus vulnerable to free radical damage and some of its components mutate 100 times more rapidly than nuclear DNA [3].

Human mtDNA is a 16,569 bp double stranded circular DNA molecule coding for 2 rRNAs, 22 tRNAs and 13 polypeptides [4], mtDNA mutations identified in several studies have been associated with poor semen quality, more recently, absence or abnormal mitochondrion has been reported in asthenozoospermic men [5]. Among these mutations, mitochondrial *tRNA* (*mt-tRNA*) gene is the hot spot for

pathogenic mutations associated with asthenozoospermic men. However, a poor genotype to phenotype correlation is common, as in the case of *tRNALeu* (UUR) T3291C mutation [6]. Distinguishing the polymorphisms and mutations is important, since fail to do so will inevitably lead to poor diagnosis and genetic advice.

In this study, we reassessed 5 reported mutations: *tRNAHis* C12187A, *tRNAThr* G15928A, *tRNAArg* T10463C, *tRNALys* A8312G and *tRNAAsp* T7572C, in clinical expression of asthenozoospermia. We first carried out the database searches for these mutations; moreover, a phylogenetic analysis was used to determine the Conservation Index (CI) of these mutations between different species. We also performed RNA Fold Web Server to predict the Minimum Free Energy (MFE) of these tRNAs with and without these mutations. We finally applied the classic pathogenicity scoring system to discern whether these mutations contributed to the genetic susceptibility of asthenozoospermia.

Materials and Methods

Data extraction

To begin our study, we took the clinical, genetic and molecular datasets from recent publications concerning the association between mt-tRNA mutations and male infertility. Moreover, we searched in the PubMed Central and other public domains with the combination of the following keywords “mitochondrial tRNA mutations, male infertility” or “mt-tRNA variants, male infertility” or “mt-tRNA mutations, asthenozoospermia” or “mt-tRNA variants, oligoasthenozoospermia” to identify the case-control studies examining the link between mt-tRNA mutations and male infertility.

Evolutionary conservation analysis

With the purpose of understanding the molecular basis of mt-tRNA mutations in male infertility, we performed the phylogenetic analysis for the CI of mt-tRNA mutations. Briefly, 17 vertebrate mtDNA sequences were used for the inter-specific analysis. The CI was then calculated by comparing the human nucleotide variants with other 16 vertebrates. Notably, the $CI \geq 70\%$ was considered as having the functional potential.

Haplogroup classification

As specific mtDNA polymorphisms create groups of related mtDNA haplotypes or haplogroups, to find haplotype/haplogroup bearing mt-tRNA mutations, we applied the Phylotree to determine the haplotypes/haplogroups [7].

Prediction the MFE structure of mt-tRNA

We further utilized the RNA Fold Web Server program to predict the MFE structure of these mt-tRNAs with and without these mutations (<http://rna.tbi.univie.ac.at/cgi-bin/RNAfold.cgi>). In addition to MFE folding, RNA Fold results provided the equilibrium base-pairing probabilities that were calculated via John McCaskill's partition function algorithm [8,9].

Determining the pathogenicity

McFarland's team first provided a weighting program for assigning a pathogenicity score to mt-tRNA mutations, their scoring system was revised in 2011 [10,11]. According to that standard, a mt-tRNA mutation will be classified as “neutral polymorphism” if its score is less than 6 points, while the score of a candidate mutation is 7-10, it belongs to “possibly pathogenic”, whereas a mutation with the score more than 11 points, it belongs to “definitely pathogenic”.

Results

Study characteristics

Two potential relevant records were identified by using the keywords mentioned in the previous section. After full-text review, we noticed that one of them was concerning the association between mt-tRNA mutation and asthenozoospermic infertile men from Tunisian population [12]. Whereas another paper which met our inclusion criteria was about the mtDNA polymorphisms in patients with infertility [13]. In total, 5 mt-tRNA mutations have been reported, as shown in Figure 1, the T10463C mutation is localized at the acceptor arm of *tRNAArg* (position 67), the T7572C and C12187A mutations occur at the T arm of *tRNAAsp* and *tRNAHis* genes (position 60 and 54), respectively, the G15928A mutation located at the anticodon stem of the *tRNAThr* (position 43), while the A8312G mutation is located at the D arm of *tRNALys* gene (position 23), the characterization of tRNA mutations was listed in Table 1.

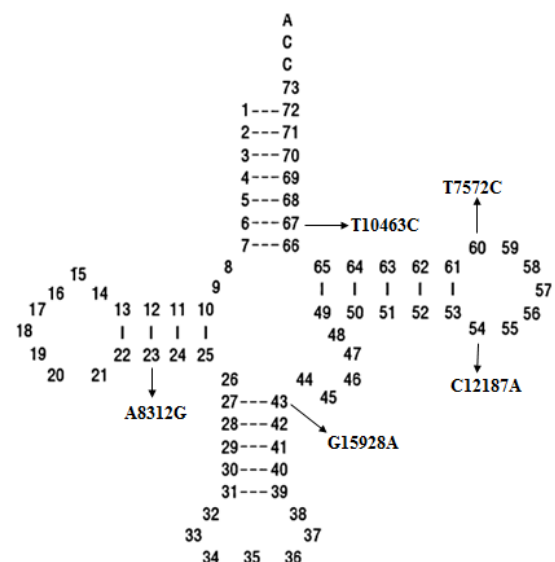


Figure 1. Mitochondrial tRNA secondary structure with standard nucleotide numbering, arrow indicated five mt-tRNA mutations analysed in this paper.

Conservation assessment of mt-tRNA mutations

Conservation assessment of the bases in the *mt-tRNA* gene suggests their evolutionary importance; therefore, it is anticipated that mutations may have pathological consequences. In this study, we performed the evolutionary conservation analysis of these mt-tRNA mutations, the CIs of the C12187A, G15928A, T10463C, A8312G and T7572C mutations are 73%, 76.9%, 94.2%, 96.1% and 19.2% respectively. It seems that the T7572C mutation had a very low level of conservation, which ruled out its involvement in the pathogenesis of male infertility.

Haplogroup analysis

By using the Phylotree (<http://www.phylotree.org/tree/main.htm>), we found that the C12187A and A8312G were non-haplogroup specific mutations, while both of G15928A and T10463C were the haplogroup T specific variants; moreover, the T7572C belonged to haplogroup T1a1, apparently, the G15928A, T10463C and T7572C were neutral polymorphisms.

Prediction the MFE structure of mt-tRNAs

To see whether nucleotide alternations affect the structure of mt-tRNA, the sequences of the wide type and mutant tRNAs harboring the C12187A, G15928A, T10463C, A8312G and T7572C were predicted using the RNA Fold Web Server. As shown in Figure 2, except for the A8312G mutation, other mutations did not affect the secondary structure of the corresponding tRNAs, indicating that the A8312G mutation may play an active role in male infertility.

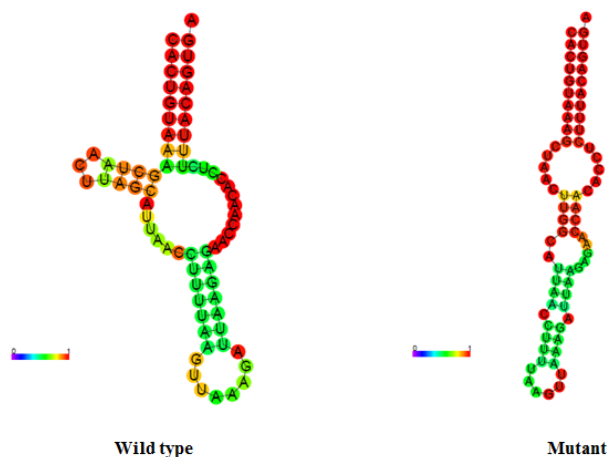


Figure 2. Prediction the MFE structure of *tRNALys* with and without the A8312G mutation.

Pathogenicity scoring system for mt-tRNA mutations

According to the revised pathogenicity scoring system [14], we classified the A8312G mutation as “definitely pathogenic” with a total score of 11 points (Table 2), whereas other mutations, C12187A (3 points), G15928A (5 points), T10463C (4 points) and T7572C (2 points) (data not shown). This indicates that they belong to the “neutral polymorphisms” category.

Discussion

In mammals, mitochondria are important components of spermatozoa and supply energy for their motility. Indeed,

mitochondria are localized in the mid piece of spermatozoa, and deliver ATP, which is mainly used to give flagellar propulsion. To date, more than 150 different mtDNA point mutations as well as a large number of mtDNA deletions are known to be associated with various human mitochondrial diseases [15,16]. However, some of them are single nucleotide polymorphisms and may not play active roles in male infertility, such as the mt-*tRNA*Leu (UUR) T3291C mutation, which is not very conserved from different species, and does not cause the alternation in secondary structure of tRNA Leu (UUR) [6].

For this purpose, we investigated the possible role of mt-tRNA mutations in male infertility. We found that there were 5 mt-tRNA mutations that had been reported: *tRNA*His C12187A, *tRNA*Thr G15928A, *tRNA*Arg T10463C, *tRNA*Lys A8312G and *tRNA*Asp T7572C. Among these, the homoplasmic C12187A and T7572C mutations were localized at the T arm of *tRNA* genes. While the T10463C and G15928A mutations disrupted the base-pairings (6C-67T, 27A-43G) in the corresponding tRNAs. However, RNA Fold Web Server Program showed that none of them caused the alternation of the secondary structure change of the *tRNA* genes. On the other hand, the A8312G mutation occurred at the D arm of *tRNALys*, *mt-tRNALys* gene was the one of the most mutable genes in mitochondrial genome [17] to date, 16 different pathogenic mutations and 8 polymorphisms in this gene have been identified as causes of various clinical symptoms (<http://mamit-tRNA.u-strasbg.fr>) [18]. Phylogenetic analysis suggested that the A8312G mutation was not haplogroup specific variant; in addition, this mutation had a very high CI (96.1%). Bioinformatics tool prediction for the secondary structure of *tRNALys* indicated that it caused a significant change of entropy between the wild type and the mutant, moreover, the pathogenicity scoring system predicted that it had a high score and belonged to a definitely pathogenic mutation, strongly indicated that it may be involved in the pathogenesis of male infertility through the alternation of its secondary structural change, which in turn resulted a failure in tRNA metabolism. Defects in tRNA metabolism will subsequently cause the impairment in mitochondrial protein synthesis.

In summary, this is the first report that reassessing the role of tRNA mutations in male infertility, and our results showed that only the A8312G mutation may be regarded as pathogenic. One of the limitations of the current study was that we did not perform the functional analysis, such as using the cybrid cell containing the A8312G mutation, by examining the steady state level, as well as the aminoacylation ability, to confirm its potential pathogenicity.

Table 1. Molecular characterization of mt-tRNA variants for male infertility.

tRNA	Nucleotide changes	Number of nucleotide in tRNA	Position	Homoplasmy or Heteroplasmy	CI (%)	G kcal/mol (wild-type)	G kcal/mol (mutant)	Haplogroup
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<i>tRNAHis</i>	C12187A	54	T arm	Homoplasmy	73	-11.16	-11.43	None-haplogroup specific mutation
<i>tRNAThr</i>	G15928A	43	Anticodon stem	Homoplasmy	76.9	-15.09	-15.07	T
<i>tRNAArg</i>	T10463C	67	Acceptor arm	Homoplasmy	94.2	-10.58	-10.86	T
<i>tRNALys</i>	A8312G	23	D arm	Homoplasmy	96.1	-16.3	-18.45	None-haplogroup specific mutation
<i>tRNAAsp</i>	T7572C	60	T arm	Homoplasmy	19.2	-10.59	-10.79	T1

Table 2. The pathogenicity scoring system for the A8312G mutation.

Scoring criteria	A8312 mutation	Score/20	Classification
More than one independent report	Yes	2	
Evolutionary conservation of the base pair	No changes	2	
Variant heteroplasmy	Yes	2	
Segregation of the mutation with disease	Yes	2	≤ 6 points: neutral polymorphisms;
Histochemical evidence of mitochondrial disease	No evidence	0	7~10 points: possibly pathogenic;
Biochemical defect in complex I, III or IV	No	0	≥ 11 points: definitely pathogenic
Evidence of mutation segregation with biochemical defect from single-fiber studies	No	0	
Mutant mt-tRNA steady-state level or evidence of pathogenicity in trans-mitochondrial cybrid studies	Weak evidence	3	
Maximum score		11	Definitely pathogenic

Conflict of Interest

None

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