MERRF and MELAS: current gene therapy trends and approaches

Ciara Ann Agresti, Penelope Nicole Halkiadakis, Peter Tolias

Center for Healthcare Innovation and Department of Chemistry & Chemical Biology, Stevens Institute of Technology, Hoboken, NJ 07030, USA.

Correspondence to: Dr. Peter Tolias, Center for Healthcare Innovation and Department of Chemistry & Chemical Biology, Stevens Institute of Technology, 1 Castle Point on Hudson, Hoboken, NJ 07030, USA. E-mail: ptolias@stevens.edu


Received: 4 Apr 2018  First Decision: 21 May 2018  Revised: 6 Jun 2018  Accepted: 12 Jun 2018  Published: 3 Jul 2018

Science Editor: Sheng-Ying Qin  Copy Editor: Jun-Yao Li  Production Editor: Cai-Hong Wang

Abstract

The mitochondrion is a unique organelle that predominantly functions to produce useful cellular energy in the form of adenosine triphosphate (ATP). Unlike other non-nuclear eukaryotic organelles (with the exception of chloroplasts), mitochondria have two lipid membranes that enclose their own mitochondrial DNA (mtDNA) and ribosomes for protein production. Similar to nuclear DNA, mtDNA is equally susceptible to mutations that may be classified as either pathogenic or nonpathogenic. Myoclonic Epilepsy with Ragged Red Fibers (MERRF) and Mitochondrial Encephalomyopathy, Lactic Acidosis, and Stroke-like Episodes (MELAS) are mitochondrial diseases originating from pathogenic point mutations located within mtDNA. Currently, there is no cure and patient care primarily focuses on treating each disease's associated symptoms. When considering the multiple barriers existing between the extracellular surface of the plasma membrane and the location of the mtDNA within the mitochondrial matrix, developing a pharmacological therapeutic that can both overcome these barriers and correct an mtDNA causing mitochondrial disease remains difficult at best. Interestingly, the field of gene therapy may provide an opportunity for effective therapeutic intervention by introducing a genetic payload (to a particular cellular gene) to induce the correction. This review primarily focuses on understanding the principles of mitochondrial biology leading to the mtDNA diseases, MERRF and MELAS, while providing a landscape perspective of gene therapy research devoted to curing these diseases.

Keywords: Mitochondria, mitochondrial biology, mitochondrial DNA, mitochondrial diseases, gene therapy, MERRF, A8344G, MELAS, A3243G
INTRODUCTION TO MITOCHONDRIAL BIOLOGY

Mitochondrial structure and function

The scientist, Ivan Wallin, stated that mitochondria had originated as endosymbionts in 1927[1]. Since each component of a mitochondrion is unique onto itself, its ancestral roots are likely to have originated from an “alpha-proteobacteria (gender: Rickettsia) that would have been engulfed by a heterotrophic host cell, probably an archeobacterium, 1.5 billion years ago” as proposed by Lynn Margulis[2]. In Margulis’ book, Origin of Eukaryotic Cells, “symbiosis usually means two or more organisms with distinct evolutionary histories derive benefit from highly specific associations with each other[3]”. While this notion may have been initially controversial, it has since been postulated that symbiosis is a necessary component of evolution. Moreover, scientific evidence has emerged to support the endosymbiotic origin of mitochondria including: free-living form of early mitochondria, mitochondrial DNA and structure, as well as the relegation of mitochondrial metabolic processes to nuclear regulation[3].

The structure of the mitochondrion is quite unique from the remaining membrane-bounded eukaryotic organelles. Its composition consists of an outer mitochondrial membrane, an inter-membrane space, an inner mitochondrial membrane, and a mitochondrial matrix (containing the mtDNA and ribosomes). Due to the four hydrocarbon tailed phospholipid, cardiolipin, the inner mitochondrial membrane is able to form cristae structures that are resultant from membrane in-folding[4]. In 1988, Robin and Wong[5] had determined that mitochondrial statistical information (number of mitochondria/cell and number of mtDNA/cell) varied between the following mammalian cell lines: rabbit lung macrophages, rabbit peritoneal macrophages, mouse LA9 fibroblasts, human lung fibroblasts and rat L-8 skeletal muscle cells. As per their analysis, the following statistical measures were obtained: the amount of mitochondrial DNA per mitochondrion remained the same in all cell varieties, the amount of mitochondrial DNA molecules per cell experienced an eight-fold difference between the different cell types, while the virtual number of mitochondria per cell also varied[5]. Therefore, dependent on the cell type, the morphology of mitochondria can vary greatly, appearing as either distinct entities ranging from 1 to 4 µm in length, or as part of an organized network[6,7]. As an interesting point, the extent of mitochondrial morphology is dependent on the balance between mitochondrial fusion and fission[6]. Furthermore, these events have the potential to be influenced by the mitochondrial membrane mechanical characteristics[8].

Resultant of its unique structure, the mitochondrion serves as a crucial cellular metabolism component. Mitochondria are well-defined as the powerhouse of the eukaryotic cell, but in addition to its famous oxidative phosphorylation role, they participate in: the creation of iron-sulfur (Fe-S) clusters, β-oxidation of fatty acids, synthesis of heme prosthetic groups, steroidogenesis (dependent on the cell variety), the urea cycle, as well as the homeostatic maintenance of calcium[2]. Mitochondria contribute greatly to the generation of free radicals, and have a participatory role in inflammation and innate immunity[4]. Moreover, mitochondria serve as a component for critical cellular signals relating to either its survival or death[2]. A summary of mitochondrial functions, with their associated purpose, appears in Table 1[2,4,9-17].

Mitochondrial DNA

While studying chick embryo mitochondria, mtDNA was discovered in 1963 by Nass and Nass[18]. Located inside of the matrix, mtDNA is a critical component of mitochondrial processes. Seemingly, the circular nature of the mitochondrial genome is reminiscent of its bacterial origins, and interestingly, its composition varies according to the nucleotide densities of its component strands. Therefore, each strand of the circular molecule is denoted as either the heavy (predominantly composed of guanine) or the light (predominantly composed of cytosine) strand[2].

Unlike nuclear DNA, mtDNA follows a non-mendelian transmission pattern and is inherited maternally via the oocyte (despite the postulation of a possible chance of paternal inheritance). In 1981, Anderson et al.[19] had
initially determined the sequence of the human mitochondrial genome, as well as provided an explanation of the sequence in relation to the arrangement and expression of its genes, which was subsequently revised in 1999\textsuperscript{20}. Human mtDNA is organized as a circular double-stranded molecule that is 16,569 base pairs in length and has a unique mitochondrial genetic code (that is different from the nuclear genetic code) due to certain codons that represent alternative amino acids or stop codons\textsuperscript{2}. Similar to nuclear DNA, mitochondrial DNA is capable of undergoing the following processes: DNA replication, RNA transcription, protein translation, DNA repair, and DNA recombination.

The mitochondrial genome [Figure 1] is highly efficient, containing no introns and a small portion of noncoding DNA\textsuperscript{2}. Interestingly, mtDNA's unique noncoding region is defined as the displacement loop (D-loop), which is approximately 1118 base pairs in length and exists approximately from nucleotides 577 to 16,028\textsuperscript{21}. This region contains transcription promoters and a replication origin defined respectively as: the heavy strand promoter (HSP), the light strand promoter (LSP) and the heavy strand replication origin (OH)\textsuperscript{2,21}. An alternative

<table>
<thead>
<tr>
<th>Mitochondrial function</th>
<th>Purpose</th>
</tr>
</thead>
</table>
| Formation of Iron-Sulfur (Fe-S) clusters\textsuperscript{2} | Small prosthetic groups that are inorganic and composed of iron-sulfur clusters\textsuperscript{3}  
Role in electron transfer, binding and activation of substrates, redox catalysis, DNA replication, DNA repair, gene expression regulation, and modification of tRNAs\textsuperscript{37} |
| \(\beta\)-oxidation of fatty acids\textsuperscript{2}                                                         | Functions to drive oxidative phosphorylation and supply acetyl-CoA for ketogenesis\textsuperscript{32}                                    |
| Synthesis of heme prosthetic groups\textsuperscript{2}                                                      | Production of heme through a complex biochemical process occurring in cells\textsuperscript{10}                                             |
| Steroidogenesis\textsuperscript{2}                                                                       | Production of steroid hormones from cholesterol\textsuperscript{2}                                                                    |
| Urea cycle\textsuperscript{2}                                                                          | Production of urea from ammonia\textsuperscript{33}                                                                                   |
| Homeostatic maintenance of calcium\textsuperscript{2}                                                     | Responsible for maintenance of calcium homeostasis required for biological processes\textsuperscript{74}                                |
| Generation of free radicals\textsuperscript{4}                                                           | Production of toxic byproducts resultant from mitochondrial energetic processes (i.e., oxidative phosphorylation)\textsuperscript{76} |
| Inflammation and innate immunity\textsuperscript{4}                                                       | Mitochondria serve as a hub for innate immune signaling\textsuperscript{76}                                                        |
|            | Mitochondrial damage leads to inflammation by releasing mitochondrial alarmins\textsuperscript{76}                                         |
| Cellular signals associated with cell survival and death\textsuperscript{2}                             | Mitochondrial participation in the regulation of cell survival and cell death induced by oxidative stress\textsuperscript{77}              |

![Figure 1. Overview of critical regulation, MERRF, and MELAS features located within the mitochondrial genome](image-url)
replication element, the light strand replication origin (OL), is situated externally to this region\[^2,^21\]. In addition, the D-loop segment is triple-stranded and contains a unique 650 nucleotide strand located between both the heavy and light strands, named 7S DNA, as determined by its sedimentation attributes\[^2,^22\]. Within the D-loop, 7S DNA is bound to the light strand, which displaces the heavy strand\[^23\]. While the entire function of 7S DNA is not completely defined, it has been hypothesized that it participates in mtDNA transcription and replication\[^6\].

DNA sequencing analysis has revealed unique regions on both strands of the human mitochondrial genome coding for 22 transfer RNAs (tRNAs), 2 ribosomal RNAs (rRNAs), and 13 polypeptides that function exclusively for mitochondrial-related processes. The 13 polypeptides corresponding to respiratory chain subunits are: complex I subunits (7), complex III subunit (1), complex IV subunits (3), and complex V subunits (2)\[^2\]. However, not all of the vital mitochondrial proteins are translated inside of the mitochondrion; only about 1% of mitochondrial proteins are synthesized within the mitochondrial matrix\[^24\]. Mitochondrial proteins that are nuclear encoded and subsequently translated in the cytosol must be trafficked and integrated into the mitochondrion. Only precursor proteins containing a specialized signal sequence are granted entry across the mitochondrial membranes using specialized multimeric protein translocator complexes: translocase of the outer membrane (TOM), the translocase of the inner membrane (TIM), and the OXA complex.

Divergent from nuclear DNA, mtDNA is not packaged like chromatin; rather it is organized into structures known as nucleoids. Since mtDNA lacks the attributes embodied in chromatin, the nucleoid structure functions to provide sequential regulation of mtDNA related processes, while simultaneously protecting against degradation and damage\[^9\]. Dependent on the cell line, nucleoids contain between one and eight mtDNA copies\[^5\]. While it remains challenging to fully characterize the composition of a nucleoid, the protein TFAM (transcription factor A of mitochondria) functions as a critical component of packaging that non-selectively binds to mtDNA to create negative supercoiling, thereby significantly compacting the mitochondrial genome\[^2,^22\]. A unique attribute is their close association to the mitochondrial inner membrane, via anchoring, as visualized by high resolution microscopy\[^2\].

Multiple copies of mtDNA can be contained within a single cell and are either identical (homoplasmy) or dissimilar (heteroplasmy)\[^9\]. Some potential reasons mitochondrial genomes may become heteroplasmic are: maternal transmission, location of the mtDNA relative to oxidative phosphorylation, lack of histones, and the truncated efficiency of the mitochondrial DNA repair machinery\[^2\]. While some mtDNA mutations may be considered pathogenic, there also exists a potential for others to have no effect. A pathogenic mutation can be overcome if the amount of non-mutated (wild-type) mtDNA is greater than mutated mtDNA within the same cell. While an advantage lies in this scenario, the opposite circumstance also exists and can induce deleterious effects. When the amount of mutated mtDNA is greater than non-mutated mtDNA, it will likely result in a mitochondrial-related disease. This phenomenon is known as the threshold effect and is a critical component regarding the existence of numerous mitochondrial myopathies\[^9\]. Further, due to these varying ratios of normal to diseased mtDNA, its imbalance will likely cause a spectrum of mitochondrial defects leading to a variation in symptom severity per patient.

**MITOCHONDRIAL DISEASES**

**Overview**

Mitochondrial diseases are a collection of illnesses arising from defects relative to the respiratory chain and oxidative phosphorylation\[^25\]. As of current, the United Mitochondrial Disease Foundation (UMDF) has compiled a comprehensive list of varying mitochondrial diseases [Table 2]\[^26\]. Mitochondrial diseases can either involve affecting a single organ or numerous organ systems, which often phenotypically have a neurologic and myopathic presentation\[^27\]. These diseases have an ability to arise from either nuclear or
mitochondrial DNA gene mutation(s), since mitochondria rely heavily on the interaction between these genomes\(^{[27]}\). As previously mentioned, the thirteen polypeptides mtDNA encodes for are respiratory chain subunits and, in addition, these complexes contain several subunits encoded by the nuclear genome\(^{[25]}\). Aside from their participation as integral members of oxidative phosphorylation, the gene products encoded by the nuclear genome dictate specific mitochondrial functions\(^{[25]}\). Mitochondrial diseases can participate in numerous genetic disorders and partake in the aging process\(^{[27]}\).

Of these mutations, upwards of 250 pathogenic mutations and rearrangements have been identified in a multitude of diseases affecting both central nervous and muscle systems\(^{[26]}\). After collecting and analyzing epidemiological data regarding both childhood and adult mitochondrial diseases, Schaefer et al.\(^{[29]}\) suggests the lowest prevalence of mitochondrial disease is approximately 1 in 5000. However, mitochondrial diseases could be far less obscure; accurate identification is limited by: disease onset related to patient age, the dual role and interplay of nuclear and mitochondrial genomes, symptoms similar to other diseases, as well as a

<table>
<thead>
<tr>
<th>Table 2. Classified mitochondrial diseases according to the United Mitochondrial Disease Foundation(^{[26]})</th>
</tr>
</thead>
<tbody>
<tr>
<td>Mitochondrial diseases</td>
</tr>
<tr>
<td>Progressive infantile poliodystrophy (Alpers disease)</td>
</tr>
<tr>
<td>Autosomal dominant optic atrophy (ADOA)</td>
</tr>
<tr>
<td>Barth syndrome/lethal infantile cardiomyopathy (LIC)</td>
</tr>
<tr>
<td>Beta-oxidation defects</td>
</tr>
<tr>
<td>Carnitine-acyl-carnitine deficiency</td>
</tr>
<tr>
<td>Carnitine deficiency</td>
</tr>
<tr>
<td>Cerebral creatine deficiency syndromes (CCDS)</td>
</tr>
<tr>
<td>Co-enzyme Q10 deficiency</td>
</tr>
<tr>
<td>NADH dehydrogenase (NADH-CoQ reductase) deficiency (complex I deficiency)</td>
</tr>
<tr>
<td>Succinate dehydrogenase deficiency (complex II deficiency)</td>
</tr>
<tr>
<td>Ubiquinone-cytochrome c oxidoreductase deficiency (complex III deficiency)</td>
</tr>
<tr>
<td>Cytochrome c oxidase deficiency (complex IV deficiency/COX deficiency)</td>
</tr>
<tr>
<td>ATP synthase deficiency (complex V deficiency)</td>
</tr>
<tr>
<td>Chronic progressive external ophthalmoplegia syndrome (CPEO)</td>
</tr>
<tr>
<td>CPT I deficiency</td>
</tr>
<tr>
<td>CPT II deficiency</td>
</tr>
<tr>
<td>Kearns-Sayre syndrome (KSS)</td>
</tr>
<tr>
<td>Lactic acidosis</td>
</tr>
<tr>
<td>Leber’s hereditary optic neuropathy (LHON)</td>
</tr>
<tr>
<td>LBSL - leukodystrophy</td>
</tr>
<tr>
<td>Long-chain acyl-CoA dehydrogenase deficiency (LCAD)</td>
</tr>
<tr>
<td>LCHAD</td>
</tr>
<tr>
<td>Subacute necrotizing encephalomyelopathy (Leigh disease or syndrome)</td>
</tr>
<tr>
<td>Luft disease</td>
</tr>
<tr>
<td>Multiple acyl-CoA dehydrogenase deficiency (MAD/glutaric aciduria type II)</td>
</tr>
<tr>
<td>Medium-chain acyl-CoA dehydrogenase deficiency (MCAD)</td>
</tr>
<tr>
<td>Mitochondrial encephalomyopathy lactic acidosis and stroke-like episodes (MELAS)</td>
</tr>
<tr>
<td>Myoclonic epilepsy and ragged-red fiber disease (MERRF)</td>
</tr>
<tr>
<td>Mitochondrial recessive ataxia syndrome (MIRAS)</td>
</tr>
<tr>
<td>Mitochondrial cytopathy</td>
</tr>
<tr>
<td>Mitochondrial DNA depletion</td>
</tr>
<tr>
<td>Mitochondrial encephalopathy</td>
</tr>
<tr>
<td>Myoneurogastrointestinal disorder and encephalopathy (MNGIE)</td>
</tr>
<tr>
<td>Neuropathy, ataxia, and retinitis pigmentosa (NARP)</td>
</tr>
<tr>
<td>Pearson syndrome</td>
</tr>
<tr>
<td>Pyruvate carboxylase deficiency</td>
</tr>
<tr>
<td>Pyruvate dehydrogenase complex deficiency (PDCD/PDH)</td>
</tr>
<tr>
<td>POLG2 mutations</td>
</tr>
<tr>
<td>Short-chain acyl-CoA dehydrogenase deficiency (SCAD)</td>
</tr>
<tr>
<td>Encephalopathy and possibly liver disease or cardiomyopathy (SCHAD)</td>
</tr>
<tr>
<td>Very long-chain acyl-CoA dehydrogenase deficiency (VLCAD)</td>
</tr>
</tbody>
</table>
spectrum of disease characteristics. This begs the question that given the possible prevalence of mtDNA mutations, what differentiates an asymptomatic carrier, oligosymptomatic patient, and an individual presenting a phenotypic clinical manifestation of the disease? As mutant and wild-type mtDNA can exist in tandem, the clinical observance of an mtDNA pathogenic mutation is largely determined by the proportion of mutant to wild-type genomes in varying tissues. Typically, greater percentages of mutant heteroplasmy are linked with a severe clinical presentation and an earlier disease onset. The variability of mitochondrial heteroplasmy can impact the clinical phenotypic manifestations of disease. Generally, the mtDNA deletion phenotypic threshold value is approximately 60%, and for other mtDNA mutations is close to 90%. The critical threshold level seems to vary within different tissues and organs, which is directly due to their energy demand. The causal relationship between heteroplasmy and the threshold effect can explain clinical phenotype variations in one individual, or the same family of individuals, via their percentage of mutated mtDNA. Moreover, on a global cellular level, mitotic segregation can further explain how some patients can possess a clinical phenotype in childhood and alternative phenotype in adulthood.

Mitochondrial diseases have been divided into the following categories: mtDNA mutations, nuclear DNA mutations, and intergenomic signaling defects. The first category, diseases attributed to mtDNA, involves the identification and characterization of pathogenic mutations with a larger number of mtDNA deletions and duplications. Conversely, the second category, nuclear DNA mutations, focuses on how the defects of the respiratory chain due to mendelian genetics comprises a considerable portion of mitochondrial diseases. Lastly, the third category, intergenomic signaling defects, are mutations that arise from the nuclear genome that directly affect the mtDNA copy number and its stability. Mitochondrial disease can present itself as either a distinct clinical syndrome or an myriad of disease phenotypes.

Diagnosing a disease of this variety can remain challenging as its fundamental roots may point to either mitochondrial or nuclear DNA, which can be determined using a two-tiered approach that includes both non-molecular (neuroimaging, muscle biopsy, cardiac and lactate concentration evaluation) and molecular (sequencing either nuclear or mitochondrial DNA for pathogenic mutations from isolated tissue) genetic testing methods. The vast diversity of causes leading to mitochondrial encephalomyopathies correlates to a diverse patient treatment plan that relies on: physiological, biochemical, and genetic approaches. Medicine, surgery, prosthetics, and perhaps dialysis or transfusions are at the forefront of physiological treatment options. Whereas biochemical treatments rely on the supplementation of oxidative phosphorylation components, and the reduction of accompanying toxicity. As a complement to the physiological and biochemical approaches, genetic treatments may depend on either genetic counseling, the alteration of heteroplasmy, and the cellular import of exogenous biological material (i.e. nucleic acids/proteins).

MERRF and MELAS

Of particular interest, MERRF and MELAS are debilitating diseases that are among the multisystemic mitochondrial diseases/syndromes that are clinically defined, alongside Kearns-Sayre syndrome (KSS). MERRF and MELAS, both classified as orphan diseases, are caused by a mtDNA point mutation that is responsible for disrupting the pairing of the codon-anticodon that is necessary for protein synthesis by disturbing the tRNAs tri-dimensional structure, as well as any associated post-transcriptional modifications.

MERRF syndrome, commonly associated with an origination in childhood is described as a multisystem disease that is characterized by spontaneous muscle contractions, generalized epilepsy, loss of control regarding bodily movements, dementia and weakness. Uncommon pathogenic variants within mtDNA encoded tRNA genes (defined as MT-TX) correlated with MERRF are: MT-TF, MT-TL1, MT-TI, and MT-TP. However, the most prominent pathogenic variant (present in approximately 80% patients or greater) is MT-TK (encodes for tRNALys), which can be observed by an A-to-G nucleotide switch at position 8344 (m.8344A>G) [Figure 1]. While the clinically detectable pathogenic variants associated with MERRF can
mostly be found in all tissues, the existence of heteroplasmy can cause mutated mtDNA to be variably distributed in tissues that leads to a changeable disease-triggering threshold level\[36]\.

Similar to MERRF, MELAS is also a multisystem disease, with an origination usually in childhood, that is characterized by stroke-like episodes, dementia, seizures, lactic acidosis, ragged red fibers (visible during muscle biopsy), vomiting and headaches\[37,38]\.

The most prominent pathogenic variant (present in approximately 80% patients) is MT-TL1 (encodes for tRNALeu(UUA/UUG)), which can be observed by an A-to-G nucleotide switch at position 3243 (m.3243A>G) [Figure 1]\[37\]. Other common pathogenic variants, such as MT-ND5, are associated with MELAS\[37\]. Confirming a patient positive for MELAS focuses on correctly diagnosing phenotypic characteristics associated with the disease, as well as employing genetic testing for further validation (extraction of mtDNA from leukocytes acts as a reliable source)\[37\]. Like MERRF, the existence of mtDNA heteroplasmy can cause mutated mtDNA to be variously distributed in tissues, which leads to a mutable disease-triggering threshold level\[37\]. Studies have shown the A3243G mutation affects mitochondrial protein synthesis, as well as the respiratory chain, when the threshold of approximately 85% mutant mtDNA is reached (as determined by Kobayashi et al.\[39]\); even though mutant mtDNA levels vary among individuals, as well as organs and tissues of a single individual\[40]\.

On average, individuals with MELAS do not have a favorable prognosis, as the 34.5 ± 19 years is the average age of death\[38]\.

**MERRF and MELAS - current advancements in gene therapy**

The current state of care for MERRF and MELAS patients focuses on a combination of genetic counseling, surgery, and palliative treatment. Potential reasons why a pharmacological cure does not currently exist (for either disease) may be attributed to the location of mtDNA within the mitochondrial matrix, followed by an inability to correct the disease-causing point mutation. It seems plausible that developing a gold-standard capable of overcoming these hurdles, and cure a mitochondrial disease, is likely difficult at best. The field of gene therapy may offer hopeful and potentially realistic possibilities that may lie beyond routine patient symptom management. It is an emerging field that seeks to treat or prevent disease using viral and/or nonviral modalities by inducing correction (of a particular cellular gene) via an exogenous payload (i.e., nucleic acids).

While gene therapy may theoretically sound like a promising prospective, downsides such as mistargeting of the exogenous payload, transient correction, inciting an immune response, and cellular apoptosis, do exist. While these existing downsides have a direct relationship with nuclear targeting, integrating mitochondrial biology brings forth an additional obstacle a gene therapy agent would have to be capable of overcoming.

For these reasons, previous gene therapy research efforts, with a focus on correcting MERRF and MELAS diseases, have been conservatively progressing for the past few decades. Gene therapy research efforts have been focused on three main approaches: nucleic acid delivery, peptide-mediated therapy, and cleavage of pathogenic mutations by mitochondrial-targeted transcription activator-like effector nucleases (mitoTALENS) [Table 3]\[41-56]\.

The goal of nucleic acid therapy is to deliver exogenous material to mitochondria harboring pathogenic mutations to restore function. Mahata et al.\[41,42\] demonstrated the RNA import complex from Leishmania can induce tRNALys import into mitoplasts isolated from a transmitochondrial cybrid MERRF cell line and in MERRF cybrid cells. Following the import of tRNALys, Mahata et al.\[41,42\] observed restored mitochondrial translation to near-wild-type levels and suppression of mutant polypeptide production. Likewise, Kolesnikova et al.\[43\] demonstrated that mitochondrial functions were partially restored due to the import of yeast tRNALys into homoplasmic transmitochondrial cybrid MERRF lines and primary human fibroblasts that were 70% ± 5% heteroplasmic for MERRF. Following this research, Karicheva et al.\[44\] transfected cybrid cells that were 90% ± 5% heteroplasmic for MELAS with tRNAleu(UUR) transcripts. Authors observed an improvement in mitochondrial translation, higher levels of mtDNA encoded respiratory subunits, and increased respiration activity\[44\].
Moreover, peptide-mediated therapy uses peptides to facilitate restoring mitochondrial function. Overexpression of mitochondrial translation elongation factors, namely EFTu or EFG2, results in the partial suppression of respiratory chain deficiency in MELAS myoblasts, but not in MERRF myoblasts [45]. Additionally, researchers determined that overexpression of leucyl-tRNA synthetase corrected mitochondrial dysfunction in human mitochondria carrying either the nearly homoplasmic A3243G or A83443G mutation [46-49]. Specifically, Perli et al. [48] demonstrated the carboxy-terminal domain of leucyl-tRNA synthetase is sufficient for cell rescue, and is even more efficient than the whole enzyme at doing so. Researchers were able to further isolate the rescuing activity of the carboxy-terminal domain of leucyl-tRNA synthetase; with this knowledge, they designed short peptide sequences capable of cell rescue that effectively bind with high affinity to both wild-type and mutant human mt-tRNALeu(UUR) and mt-tRNALys [49]. Chang et al. [50,51] conducted several studies suggesting that a Pep-1 peptide delivery system is capable of rescuing mitochondria containing the MERRF mutation. Pep-1, a cell-penetrating peptide, can translocate mtDNA from wild-type mitochondria into cybrid cell models of MERRF [50,51]. After three days of treatment with Pep-1-mediated mitochondrial delivery, mitochondrial function was recovered and cells were maintained for twenty-one days [50]. Recently, Chang et al. [52] demonstrated that Pep-1-mediated mitochondrial delivery can ameliorate mitochondrial function in cells containing the pathogenic MELAS mutation. Further, Muratovska et al. [53] and Taylor et al. [54] independently synthesized peptide nucleic acids (PNA) complementary to the sequence containing the MERRF mutation; upon entry into the mitochondria, PNA oligomers are able to selectively inhibit replication and translation of the mutant mtDNA template using in vitro and cell-based studies, respectively.

Lastly, mitochondrial-targeted transcription activator-like effectors (mitoTALENs) are designed to cleave specific sequences of mtDNA that contain a pathogenic mutation [55,56]. When tested on heteroplasmic cybrid cells harboring MERRF and MELAS, respectively, mitoTALENs permanently reduced the mutant load and resulted in the return to normal mitochondrial respiratory activity [55,56].

CONCLUDING REMARKS AND FUTURE DIRECTIONS

A review of gene therapy strategies targeting the mtDNA disease’s, MERRF and MELAS, presents a series of realistic possibilities for their potential correction. Understanding the backstory of mitochondrial biology sets the tone for further understanding the path of mitochondrial dysfunction when considering an mtDNA-causing mitochondrial disease. While multiple forms of care exist for patients living with either MERRF or MELAS, none are capable of permanently correcting its associated mitochondrial mutation. Even if a certain approach is capable of restoring mitochondrial function, one must be prepared to answer several general arguments: (1) does the treatment approach lead to permanent or transient correction; (2) if the treatment was initially conducted in an in vitro setting, does it have the potential to be translated to an in vivo setting; and (3) if it can be adapted to an in vivo environment, what type of an immunogenicity is it met with? Rationally speaking, a researcher targeting MERRF or MELAS diseases must be both practiced and prudent.
when developing a gene therapy vector. Future directions surrounding research for these diseases should focus on developing a therapeutic vector that is capable of answering the aforementioned questions relating to: overcoming cellular and mitochondrial barriers, establish long-term (if not permanent) correction, and exhibiting a low (to negative) immunogenicity profile in patients.

DECLARATIONS

Acknowledgments
The authors take special acknowledgment to Ueli Gubler, Ph.D. for his scientific guidance and expertise regarding the subject matter.

Authors’ contributions
Concept and design: Agresti CA, Tolias P
Data acquisition and analysis: Agresti CA, Halkiadakis PN
Manuscript preparation: Agresti CA, Halkiadakis PN, Tolias P
Critical revision and finalizing of the manuscript: Agresti CA, Tolias P

Availability of data and materials
Not applicable.

Financial support and sponsorship
None.

Conflicts of interest
All authors declared that there are no conflicts of interest.

Ethical approval and consent to participate
Not applicable.

Consent for publication
Not applicable.

Copyright
© The Author(s) 2018.

REFERENCES


46. Park H, Davidson E, King MP. Overexpressed mitochondrial leucyl-tRNA synthetase suppresses the A3243G mutation in the mitochondrial tRNA(Leu(UUR)) gene. RNA 2008;14:2407-16.


