Table 1: Screening Guidelines

Table 1: Screening Guidelines	Τ.			Γ_
	At .	At 1-2	As needed	Comments
	diagnosis,	year	depending	
	if not	intervals	on .	
	previously	and as	symptoms	
	obtained	needed	or disease	
	*	*	type	
Audiology	*	*		11 B B B B
Cardiology	*	*		Holter recording depending on the underlying diagnosis and
Blood Pressure	*	*		risk of heart block; up to every
Echocardiogram	*	*		3-6 months for disorders
Electrocardiogram	*	*	*	associated with high-risk of
Holter			*	arrhythmias, such as mtDNA
Cardiac MRI			*	deletion disorders
				Echocardiograms may be
				performed less frequently in
				low-risk patients after several
				years of monitoring
Endocrinology	*	*		Endocrine screening strongly recommended in those with
Basic chemistries		*		mtDNA deletion disorders
Calcium (Ca), Magnesium	*	*		
(Mg) & Phosphate			ate	
Cortisol-ACTH-			*	
Aldosterone-Renin		a.		
Ca & phosphate, urine	*	*		
Gonadotrophins		a.	*	
Hemoglobin A1c	*	*		
Parathyroid hormone	*	*	*	
Thyroid stimulating	*	*		
hormone & free				
thyroxine	*	*		
Vitamin D	*	*	*	
Dual X-ray			_	DXA especially if unexpected
Absorptiometry (DXA)				fractures
Gastroenterology			*	
Amylase-lipase	*	*	T	
Transaminases	7	7	*	
Stool elastase				
Swallow evaluation			*	
Growth and anthronometric	*	*		Recommended at each visit
Growth and anthropometric				necommended at each visit
parameters Hematology				Obtained more routinely in
пешасооду				those with high risk of or

CBC with differential	*		*	symptomatic bone marrow
Iron studies including			*	dysfunction
ferritin				
Immunology			*	With recurrent infections
Neurology				
Developmental &	*	*		Clinical appraisal or formal
Cognitive Assessments				neuropsychological tests; formal testing recommended
Electroencephalogram			*	with regression
Ophthalmology				
Exam	*	*		
Electroretinogram			*	
Optical Coherence			*	
Tomography				
Psychiatry	*	*		
Mood and Anxiety				
Disorder Screening				
Pulmonology				
Pulmonary function			*	Especially with myopathy & if non-ambulatory or with
				brainstem dysfunction
Polysomnogram			*	bramstem aysranetion
Renal				
CMP with Mg and Phos	*	*		
Albumin/creatine, urine	*	*		

Table 2: Other specialist consultations to consider at time of diagnosis and at 1-2 year intervals as needed based on symptoms

Audiology Cardiology

Endocrinology

Ear, Nose and Throat

Gastroenterology

Genetics

Hematology

Immunology

Nephrology

Neurology

Ophthalmology

Orthopedics

Palliative Care

Physical Medicine & Rehab/Physiatry

Psychiatry (for patient or family)
Psychology (including Family Counseling)
Pulmonology
Social Work
Sleep Medicine
Therapy services including PT, OT and ST

Table 3: Illness, Anesthesia and Stroke Management

Illness Management³

- 1. Specific decisions about patient management including hospitalization require clinical judgment and should be case-specific. Decisions should reflect the individual patient's presentation as well as an understanding of the etiology for the acute decompensation and the pathophysiology of the underlying mitochondrial disorder.
- 2. Patients with a mitochondrial disease should carry an emergency care plan that details their underlying disorder and provides management recommendations.
- 3. Patients with a mitochondrial disease should consider wearing a Medic Alert bracelet when appropriate depending on their clinical symptomology.
- 4. Mitochondrial patients should take precautions to prevent entering catabolism, especially when exposed to medical stressors, including avoiding prolonged fasting and receiving dextrose-containing intravenous (IV) fluids before, during, and after procedures and surgeries. (Dextrose should not be provided or provided in limited quantity as indicated by clinical status in suspected or confirmed disorders of pyruvate metabolism, if the patient is on a ketogenic diet, or the patient has had a previous adverse response to high glucose delivery.)
- 5. Evaluation of a mitochondrial patient in the acute setting should include evaluation of routine chemistries, glucose, transaminases, and lactate; all other testing is as clinically indicated, although one must keep in mind the potential for cardiac and neurologic decompensations in these patients.
- 6. Treatment during acute decompensation should include dextrose-containing IV fluids, stopping exposure to potentially toxic medications, and correction of any metabolic derangements. (Note: dextrose should be provided only in limited in quantity or not at all, as indicated by clinical status in suspected or confirmed disorders of pyruvate metabolism, if the patient is on a ketogenic diet, or the patient has had an adverse response to high glucose delivery.) IV fluid rate should be based on the clinical situation. Outpatient mitochondrial therapies should be continued when possible.
- 7. Lipids can be used when needed in mitochondrial patients, even in the presence of secondary fatty-acid oxidation dysfunction.
- 8. The following medications should be avoided in patients with mitochondrial disease when possible and, if given, they should be used with caution: valproic acid; statins; metformin; high-dose acetaminophen; and selected antibiotics, including aminoglycosides, linezolid, tetracycline, azithromycin, and erythromycin.

9. Repeat neuroimaging should be considered in any mitochondrial patient with an acute change in neurologic status.

Anesthesia and Surgical Management³

- 1. Patients with mitochondrial diseases are at an increased risk of anesthesia-related complications.
- 2. Preoperative preparation of patients with mitochondrial disease is crucial to their perioperative outcome. Patients should minimize preoperative fasting and have glucose added to their perioperative IV fluids, unless they are on a ketogenic diet or have been demonstrated to have adverse reaction to higher glucose intake.
- 3. Caution must be used with volatile anesthetics because mitochondrial patients may potentially be hypersensitive.
- 4. Caution must be used with muscle relaxants in those mitochondrial patients with a preexisting myopathy or decreased respiratory drive.
- 5. Mitochondrial patients may be at a higher risk for propofol infusion syndrome and propofol use should be avoided or limited to short procedures.
- 6. One should consider slow titration and adjustment of volatile and parenteral anesthetics to minimize hemodynamic changes in mitochondrial patients.
- 7. Local anesthetics are generally well-tolerated in patients with mitochondrial defect.
- 8. There is no clear established link between malignant hyperthermia and mitochondrial disease.

Stroke Management^{3, 72}

- 1. Stroke-like episodes in primary mitochondrial disease typically have correlating visible magnetic resonance imaging abnormalities.
- 2. IV arginine hydrochloride should be administered urgently in the acute setting of a stroke-like episode associated with the MELAS m.3243 A>G mutation in the *MTTL1* gene and considered in a stroke-like episode associated with other primary mitochondrial cytopathies as other etiologies are being excluded. Patients should be reassessed after 3 days of continuous IV therapy.
- 3. The use of daily oral arginine supplementation to prevent strokes should be considered in MELAS syndrome.
- 4. The role of monitoring plasma arginine and citrulline levels and oral citrulline supplementation in the treatment of MELAS requires further research.

Table 4: Medication Cautions

Medication	Common Uses	Concern in Mitochondrial Disease
Acetaminophen	Analgesic, fever prevention, headaches	Chronic or frequent use may deplete glutathione and cause hepatopathy
Aminoglycosides	Antibiotic	Hearing loss

Antiretrovirals	HIV therapy	Impaired mtDNA replication and worsening peripheral neuropathy, liver dysfunction or myopathy
Botulinum toxin	Dystonia, Spasticity	Worsening of weakness
Butterbur	Headache	May contain pyrrolizidine alkaloids (oxidants) and cause hepatopathy
Metformin	Diabetes	Lactic acidosis
Topiramate	Epilepsy, Headache, Intracranial Hypertension	Lactic acidosis
Statins	Hypercholesterolemia	Worsening myopathy and elevated creatine kinase (CK)
Valproic Acid	Epilepsy, Headache, Mood disorders, Movement disorders, Tone abnormalities	Irreversible liver failure and onset of hepato-encephalopathy, especially in <i>POLG</i> -related disorders; worsening of seizures
Vigabatrin	Epilepsy	Inhibition of the mitochondrial nucleoside salvage pathway and worsening of mtDNA depletion disorders

With the exception of valproic acid in POLG-related disorders, these medications are **not** contraindicated and may be used with caution