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1 Mitochondrial Neurogastrointestinal Encephalomyopathy (MNGIE	E):
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- 2 Position Paper on Diagnosis, Prognosis and Treatment by the MNGIE
- 3 International Network
- 4 Running Title: Position Paper on MNGIE
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1 SUMMARY

2 Mitochondrial Neurogastrointestinal Encephalomyopathy (MNGIE) is a rare autosomal recessive disease caused by TYMP mutations and thymidine phosphorylase (TP) deficiency. 3 Thymidine and deoxyuridine accumulate impairing the mitochondrial DNA maintenance and 4 integrity. Clinically, patients show severe and progressive gastrointestinal and neurological 5 manifestations. The onset typically occurs in the second decade of life and mean age at death 6 7 is 37 years. Signs and symptoms of MNGIE are heterogeneous and confirmatory diagnostic tests are not routinely performed by most laboratories, accounting for common misdiagnosis. 8 9 Factors predictive of progression and appropriate tests for monitoring are still undefined. 10 Several treatment options showed promising results in restoring the biochemical imbalance of 11 MNGIE. The lack of controlled studies with appropriate follow-up accounts for the limited 12 evidence informing diagnostic and therapeutic choices. The International Consensus Conference (ICC) on MNGIE, held in Bologna, Italy, on March 30th-31st, 2019, aimed at an 13 evidence-based consensus on diagnosis, prognosis and treatment of MNGIE among experts, 14 15 patients, caregivers and other stakeholders involved in caring the condition. The conference 16 was conducted according to the National Institute of Health Consensus Conference methodology. A consensus development panel formulated a set of statements and proposed a 17 18 research agenda. Specifically, the ICC produced recommendations on: (1) diagnostic 19 pathway; (2) prognosis and the main predictors of disease progression; (3) efficacy and safety 20 of treatments; and (4) research priorities on diagnosis, prognosis and treatment. The Bologna 21 ICC on diagnosis, management and treatment of MNGIE provided evidence-based guidance 22 for clinicians incorporating patients' values and preferences.

23

24 SYNOPSIS

1	This is the first International Consensus Conference (ICC) aimed at an evidence-based
2	consensus on diagnosis, prognosis and treatment among experts, patients, caregivers and
3	other stakeholders involved in MNGIE. The ICC provided recommendations on diagnostic
4	pathway, prognosis and the main predictors of disease progression, efficacy and safety of
5	treatments, and, finally, identified priorities on cogent research topics on MNGIE.

6

7 ABBREVIATION LIST

8 CAPD: continuous ambulatory peritoneal dialysis; CDP: consensus development panel; CPEO: chronic progressive external ophthalmoplegia; dThd: thymidine; dUrd: deoxyuridine; 9 EE-TP: erythrocyte encapsulated TP; EWGs: expert workgroups; GI: gastrointestinal; HD: 10 hemodialysis; HSCT: hematopoietic stem cell transplantation; ICC: International Consensus 11Conference; MNGIE: Mitochondrial Neurogastrointestinal Encephalomyopathy; mtDNA: 12 mitochondrial DNA; OLT: orthotopic liver transplantation; PEG: percutaneous endoscopic 13 gastrostomy; QoL: quality of life SC: scientific committee; SIBO: small intestinal bacterial 14 15 overgrowth; TC: technical committee; TP: thymidine phosphorylase. 16

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1 COMPLIANCE WITH ETHICS GUIDELINES

2 CONFLICT OF INTEREST

- 3 Valerio Carelli is a consultant for Santhera Pharmaceuticals, GenSight and Stealth
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- 11 D'Alessandro, Massimiliano Filosto, Maria Teresa Dotti, Hanna Mandel, Laura Ludovica
- 12 Gramegna, Olimpia Musumeci, Matteo Cescon, Roberto D'Angelo, Alessia Pugliese,
- 13 Antonella Spinazzola, Elisa Boschetti, Javier Torres-Torronteras, Irina Zaidman, Antonio
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- 16 Bolletta, Riccardo Bolletta, Massimo Zeviani, Antonio Daniele Pinna and Mauro Scarpelli
- 17 declare that they have no conflict of interest.
- 18

19 AUTHORS' CONTRIBUTION

- 20 V.C., R.D.G, L.P., R.R., E. Ba., F.N, L.V. participated to the planning, conducting and
- 21 reporting of the project; R.B, A.B M.H, R.M., A.P., R.DA., E. Bo. participated to the
- 22 planning and conduction of the project.
- 23 B.E.B. contributed to the conduction and reporting of the project.
- 24 L.L.G., M.S., A.S., G.T., C.G., O.M., M.C.M., M. C., M.T.D., M. L., H.M., A. S., J. T-T, I.
- 25 Z., H.Z. contributed to the conduction of the project.

1 ETHICS

2	This article does no	ot contain any	studies with	human or animal	subjects	performed by	y any	y of
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3 the authors. No ethical approval was required.

4

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- 6 Azienda Ospedaliero-Universitaria di Bologna Policlinico St. Orsola-Malpighi and IRCCS,
- 7 Istituto delle Scienze Neurologiche di Bologna (Bologna, Italy).

8

9 SEARCH TERMS

- 10 Mitochondrial Neurogastrointestinal Encephalomyopathy; MNGIE; enzyme replacement;
- 11 mitochondrial disease; *TYMP*; thymidine phosphorylase, consensus conference.
- 12

1 1 INTRODUCTION

Mitochondrial Neurogastrointestinal Encephalomyopathy (MNGIE) is an autosomal 2 recessive disease caused by mutations in the thymidine phosphorylase gene (TYMP)(Pacitti et 3 al 2018). TYMP encodes for thymidine phosphorylase (TP), which catabolizes thymidine 4 5 (dThd) and deoxyuridine (dUrd) into their respective bases. TYMP mutations markedly reduce/abolish TP activity leading to accumulation of dThd and dUrd and mitochondrial 6 7 DNA (mtDNA) defects. MNGIE is an ultra-rare condition, characterized by severe gastrointestinal (GI) and 8 neurological symptoms(D'Angelo et al 2016) that is often misdiagnosed. Although the 9 10 disease is progressive and fatal, natural history is still uncharacterized(Nishino et al 2000; Garone et al 2011; Corazza et al 2019). Various experimental therapeutic approaches aimed 11 to the temporary enzyme replacement, e.g. erythrocyte encapsulated TP (EE-TP) infusions 12 (Pacitti et al 2018), or permanent restoration of TP activity through hematopoietic stem cell 13 transplantation (HSCT) (Halter et al 2015) and orthotopic liver transplantation (OLT) (De 14 15 Giorgio et al 2016). Since the severity of GI symptoms influences treatment success, timing of HSCT and OLT is crucial (Halter et al 2015; De Giorgio et al 2016). Possible future options 16 17 include gene therapy, which has shown pre-clinical efficacy (Torres-Torronteras et al 2014; Cabrera-Pérez et al 2015; Torres-Torronteras et al 2016; Torres-Torronteras et al 2018; 18 Yadak et al 2018; Cabrera-Pérez et al 2019). 19 20 An International Consensus Conference (ICC) was held to produce an unbiased, evidencebased assessment on MNGIE, leading to a consensus and guidance on the following areas: (I) 21 22 diagnostic pathway; (II) prognosis and main predictors of disease progression; (III) efficacy 23 and safety of treatments.

- 24
- 25 2 METHOD

1 2.1 Panel/experts selection

The Bologna MNGIE ICC was organized and promoted by the Azienda Ospedaliero-2 Universitaria di Bologna, Policlinico S.Orsola-Malpighi, and the IRCCS Istituto delle 3 Scienze Neurologiche di Bologna, Italy, according to the NIH Consensus Development 4 Program methodology(Nair et al 2011) and the Methodological Handbook of the Italian 5 National Guideline System(Candiani et al 2009). The members of the technical committee 6 7 (TC) and scientific committee (SC), the expert workgroups (EWGs) and the consensus development panel (CDP) were invited based on their expertise in the field, ensuring the 8 9 participation of all the clinical and non-clinical stakeholders (including patients) and a broad 10 involvement of healthcare professionals from all the clinical aspects of MNGIE. Researchers 11were identified based on a review of the main authors in the field. The official language of 12 the conference was English supported by a professional translator. A declaration of interest form was signed by every participant. Of the 36 stakeholders invited to the ICC, four 13 declined and five accepted but did not participate. One member attended via teleconference. 14 15

16 2.2 The assignment, scoping and assessment stages

17 The assignment, scoping and assessment stages occurred between January 2018 and March 2019. The ICC took place in Bologna on March 30th and 31st. During assignment, the SC 18 appointed a TC, a CDP and three EWGs (Appendix 1). The SC identified three MNGIE 19 20 topics (diagnosis, prognosis and treatment) and questions to be addressed by the EWGs 21 (scoping). Assessment of the evidence was carried out by the TC through a systematic 22 literature search with evidence mapping (Bragge et al 2011) according to the PRISMA guidelines (Moher et al 2009) (Appendix 2). Studies of any design, in English language, 23 published in full on peer reviewed journals, reporting original data on diagnosis, prognosis, 24 25 and/or treatment of MNGIE on humans were searched on MEDLINE and the Cochrane

Central Register of Controlled Trials in January 2018 and 2019, and finally updated in May 1 2 2020. Retrieved studies were selected independently by LV, EB and RD. Disagreement was resolved by discussion. Each study was graded according to four classes of methodological 3 quality (from Class I, highest quality to Class IV, lowest quality) according to the 4 5 Classification of Evidence Schemes of the Clinical Practice Guideline Process Manual of the American Academy of Neurology (Appendix 3) (Gronseth et al 2011) and appraised by the 6 7 EWGs to draft answers to the questions posed by the SC. During the ICC the scientific evidence and the answers to the questions were presented by the EWGs. The final statements 8 9 by the CDP were presented at the end of day 2 to the audience including stakeholders and 10 general public.

11

12 3 RESULTS

The literature search retrieved 1,305 citations after duplicate removal; 1,146 were excluded 13 because the covered topic was not of interest for our review (Appendix 2). Of the 159 full 14 15 text articles selected, 81 were excluded mostly because they were animal studies or studies 16 describing genetic mutations causing the clinical manifestations of MNGIE. The 78 selected articles were submitted to the three working groups. Since 36 of them were out of scope 17 18 regarding the specific topics, 42 were used for the statements (4 of them were assigned to two 19 topics each). The CDP issued the following Position Statements on diagnosis, prognosis and 20 treatment of MNGIE. In Appendix 4 a summary of the scientific evidence and rationale for 21 the statements is presented.

22

23 **3.1 Position Statements on the Diagnosis of MNGIE**

24 The following statements are based on 7 Class III level studies (case series with controls)

25 (Spinazzola et al 2002; Nishigaki et al 2003; Marti et al 2004; Valentino et al 2007;

- 1 Mohamed et al 2014; Gramegna et al 2018; Kipper et al 2020), 7 Class IV level studies (two
- 2 case series and five case reports) (Millar et al 2004; Marti et al 2005; Giordano et al 2006;
- 3 Filosto et al 2011; Garone et al 2011; Scarpelli et al 2013; Corazza et al 2019) and expert
- 4 opinion.

5 3.1.1 Clinical elements that can indicate MNGIE

- 6 MNGIE can be suspected when one or more of the following clinical cardinal elements are
- 7 present:
- 8 Symptoms and signs of otherwise unexplained GI dysmotility
- 9 Thin constitution/cachexia, even with normal food behaviour and <u>nutritional</u> intake
- 10 Neurological features such as ptosis and symptoms suggesting peripheral neuropathy
- A progressive course of the above with frequent misdiagnosis
- 12 The features of the full-blown MNGIE typically are:
- Symptoms onset: childhood, adolescence/young adulthood (typical), adulthood (late
- 14 onset, >40 years)
- GI symptoms/signs: sub-occlusive episodes, nausea, vomiting, early satiety,
- 16 borborygmi, severe abdominal pain, abdominal distension, dysphagia, constipation and
- 17 diarrhoea, acute peritonitis due to small bowel perforation
- 18 Unexplained weight loss, thinness, cachexia, even with normal food behaviour and
- 19 <u>nutritional intake</u>
- Radiological GI signs: small bowel diverticulosis, GI dilation (e.g. gastric or intestinal
- 21 dilation)
- Neurological symptoms/signs: chronic progressive external ophthalmoplegia (CPEO),
- 23 ptosis, peripheral neuropathy, hearing loss
- Neuroradiological signs: leukoencephalopathy without other neuroradiological
- 25 abnormalities

1 •	Metabolic	alterations:	liver steato	sis evolvin	g in cii	rrhosis,	pancreatitis, earl	y onset
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- 2 diabetes mellitus, increased triglyceride levels, elevated plasma lactate
- 3 MNGIE is most frequently misdiagnosed as:
- 4 Anorexia nervosa
- GI diseases: Crohn's disease, coeliac disease, esophagitis and/or gastritis, irritable bowel
- 6 syndrome, superior mesenteric artery syndrome, Whipple's disease, chronic intestinal
- 7 pseudo-obstruction
- 8 Neurological diseases: chronic inflammatory demyelinating polyneuropathy, Charcot-
- 9 Marie-Tooth disease, other mitochondrial diseases such as CPEO, Kearns-Sayre
- 10 syndrome
- 11 **3.1.2 Recommended diagnostic tests**
- 12 Cardinal diagnostic tests:
- 13 Swallowing test, gastric emptying and GI manometry (when possible): altered GI
- 14 motility and transit
- 15 Brain MRI: leukoencephalopathy without any other neuroradiological abnormalities
- 16 (almost universally present) (Table 1 and Fig. 2)
- Nerve conduction studies: peripheral neuropathy, predominantly demyelinating
- 18 Ancillary tests:
- 19 Muscle biopsy: ragged-red and COX deficient fibers, deficiencies of respiratory chain
- 20 enzyme activities, ultrastructurally abnormal mitochondria, and mtDNA depletion,
- 21 multiple deletions, and somatic point mutations
- Mucosal GI histology of small bowel (to exclude other conditions), and gut full thickness
- 23 biopsy (when possible)
- 24
- 25 3.1.3 Recommended metabolic and genetic tests

1 Mandatory tests to confirm MNGIE diagnosis:

2	• TYMP sequencing: homozygous or compound heterozygous allelic pathogenic variants,
3	no further testing.
4	• If one variant of uncertain significance or a wild-type sequence is identified, the
5	following biochemical assessments should be performed:
6	\circ TP activity: severely reduced or virtually absent in the buffy coat (below 8%
7	of the mean of reference TP values; laboratory cutoffs may differ depending
8	on sample processing and biochemical assay). If TP activity is only partially
9	reduced, then it is mandatory to measure plasma dThd and dUrd levels. The
10	diagnosis is excluded if TP activity is normal.
11	• dThd and dUrd levels: increased in plasma (assessment of urine is unreliable)
12	Fig. 1 shows the recommended algorithm in persons with suspected MNGIE.
13	
14	3.2 Position Statements on the Prognosis of MNGIE
15	The following statements are based on one Class II level study (retrospective cohort) (Garone
16	et al 2011), two Class III level study (retrospective cohorts) (Nishino et al 2000; Corazza et al
17	2019), 4 Class IV level studies (four case reports) (Ionasescu et al 1984; Carrozzo et al 1998;
18	Marti et al 2005; Massa et al 2009) and expert opinion.
19	
20	3.2.1 The natural history of MNGIE
21	• Mean age at onset: 17.9 years (5 months - 43 years).

• GI symptoms (57% at onset; 100% at diagnosis); onset/diagnosis: diarrhea, abdominal

23 pain, borborygmi, vomiting, pseudo-obstruction (32%- 65%), weight loss/cachexia

24 (100%); evolution: diverticulosis/diverticulitis (67%), hepatopathy (22%).

1	• Neurological symptoms / signs (43% at onset; 100% at diagnosis);
2	onset/diagnosis/evolution: ocular signs (ptosis, ophthalmoparesis) (74-100%),
3	polyneuropathy (92-100%), hearing loss (39-45%), leucoencephalopathy ($\pm 100\%$);
4	cognitive impairment (20%).
5	• Symptoms are cumulative and progressive.
6	• Mean age at death is reported to range between 35 and 37 years; survival 100% before 19
7	years and $< 5\%$ after 50 years.
8	• Death is mainly due to GI and liver complications (intestinal perforation, intestinal
9	bleeding, liver failure, aspiration pneumonia, complications due to small intestinal
10	bacterial overgrowth (SIBO) or infection related to central venous catheter for parenteral
11	nutrition) and cachexia.
12	
13	Overall survival is the only outcome reported in the literature. Weight loss is an important
14	feature of MNGIE, but data on its prognostic role are lacking.
15	
16	3.2.2 Phenotypes of MNGIE
17	MNGIE has two different presentations distinguished by age of onset: "Early Onset" (or
18	"Classic") and "Late Onset" (Table 2). Severity can vary among family members. Available
19	data do not allow a differentiation of clinical phenotypes based on symptoms at onset (GI or
20	neurological). Neurological manifestations may be subtle and insidious leading to late
21	recognition by both patients and physicians. When GI symptoms are the first manifestation,
22	the diagnosis and consequently the appropriate treatment may be significantly delayed
23	because of misdiagnosis.
24	In clinical practice, the presence and/or severity of GI involvement are considered a negative
25	prognostic factor, for both morbidity and mortality. Apparently, there is no correlation

1 between genotype, phenotype and outcome. Residual TP activity of 10-15% has been

2 associated with moderate increases of nucleosides and "Late Onset" MNGIE compared to the

- 3 "Classic" form.
- 4

5 3.2.3 Impact of different phenotypes on the natural history of MNGIE and outcomes

6 Whether "Classic" ("Early Onset") and "Late Onset" phenotypes have different disease

7 progression remain unsettled. All patients with Late Onset phenotype reported in the

8 literature were alive at follow-up ranging between 8 to 24 years. In the "Classic" phenotype,

9 age of onset is not related to life expectancy. At present, overall survival after onset is the

10 only available outcome reported in the literature.

11

12 3.2.4 Events indicating disease progression

13 The following events can be considered as important milestones related to progression: GI

14 sub-occlusive episode, decompressive percutaneous endoscopic gastrostomy (PEG),

15 aspiration pneumonia, abdominal surgical procedures, septic episode due to SIBO, need for

16 enteral tube feeding or PEG, onset of intestinal failure (and subsequent need of parenteral

17 nutrition), liver cirrhosis, loss of unaided walking ability. Recommended assessments to

18 monitor MNGIE progression are listed in Table 3.

19

20 **3.3 Position Statements on the Treatment of MNGIE**

21 The following statements are based on 25 Class IV level studies (one retrospective cohort,

22 one case series, 24 case reports) (Hirano et al 2006; la Marca et al 2006; Lara et al 2006;

23 Yavuz et al 2007; Moran et al 2008; Filosto et al 2012; Sicurelli et al 2012; Bax et al 2013;

Finkenstedt et al 2013; Hussein 2013; Ariaudo et al 2015; Casarez et al 2015; Halter et al

25 2015; Peedikayil et al 2015; De Giorgio et al 2016; Sivadasan et al 2016; Baker et al 2017;

D'Angelo et al 2017; Roeben et al 2017; Chandra et al 2018; Hanbali et al 2018; Levene et al 1 2018; Yadak et al 2018; Levene et al 2019; Kripps et al 2020) and expert opinion. 2 3 3.3.1 Treatments effective in temporarily restoring the biochemical imbalance 4 5 "Short term" is defined as a period of time required stabilizing a patient waiting for permanent treatment or as compassionate use. 6 7 Overall, hemodialysis (HD), continuous ambulatory peritoneal dialysis (CAPD), EE-TP and platelet infusion have been effective in achieving temporary improvement of the biochemical 8 9 imbalance in several MNGIE patients. However, there are practical limitations, safety issues 10 and unclear clinical effects associated with these approaches. Specifically: 11 12 • EE-TP seems to be effective on a monthly-based administration in terms of biochemical 13 and clinical improvement (4 patients out of 5); mild immunological reaction against 14 bacterial TP mainly with repeated infusions may occur (2 out of 5 patients). EE-TP is 15 currently under clinical evaluation. • CAPD seems to be well tolerated with anecdotal biochemical efficacy; peritoneal 16 17 sclerosis due to repeated procedures may be a safety concern. 18 · HD has very short-term biochemical effects as nucleosides return to high levels a few 19 hours after the procedure. Disadvantages include the need of venous access, an intensive procedure schedule (3-4 sessions per week) and the possible occurrence of hypotension, 20 21 fluid overload or infections. 22 • Platelet infusion has been reported to achieve some biochemical improvement. Safety issues include allergic and immunological reactions. 23

2 option, or for compassionate use. 3 3.3.2 Effective treatments that permanently restore the biochemical imbalance 4 5 Permanent treatment options are aimed at restoring TP resulting in the long-term clearance of dUrd and dThd. The improved biochemical profile is expected to be associated with clinical 6 7 stabilization (i.e., halting tissue damage progression) or improvement. 8 • HSCT is effective in permanently restoring the biochemical imbalance. It requires chemotherapy and immunosuppressive therapy and is associated with a high risk for 9 10 complications and mortality related to therapy, including graft versus host disease. · OLT is effective for permanent restoration of the biochemical imbalance and it does not 11 12 require preoperative conditioning. Patients with severe malnutrition or previous episodes 13 of small bowel perforation, sub-occlusion or sepsis related to SIBO could be at high risk 14 of peri- and post-operative complications and should not be considered for OLT; 15 metabolic complications such as chronic kidney insufficiency, diabetes or cardiovascular disease related to long-life immunosuppressive therapy may be long-term issues. 16 17 3.3.3 Treatments that improve the patient's health in terms of quality of life (QoL) and 18 19 functional status Any temporary or permanent treatment should be considered as soon as diagnosis is 20

We suggest consideration of EE-TP or CAPD in patients waiting for a permanent treatment

- 21 confirmed. EE-TP and CAPD are effective with some temporary improvement of QoL and
- 22 minimal complications. HSCT is effective in the long term for improving QoL and functional
- 23 status, although limited by a high post-treatment mortality rate (63%) in severely
- 24 symptomatic adult patients. OLT is effective in the long term for improving QoL and
- 25 functional status, although limited evidence is available.

2 3.3.4 Appropriateness of treatments that temporarily or permanently restore the

3 biochemical imbalance

1

Temporary treatments should be considered at any age based on clinical condition, before 4 5 permanent treatments, e.g. while waiting /on list for either HSCT or OLT. Once the diagnosis is confirmed, permanent treatments are recommended as early as possible for both HSCT and 6 7 OLT according to eligibility criteria (e.g. severity). Ideal candidates should be those at an early stage of MNGIE. In patients who are oligosymptomatic, either HSCT or OLT can be 8 considered. HSCT should be considered in pediatric patients and young adults with normal 9 10 liver function, mild or no GI manifestations (e.g. absence of intestinal pseudo-obstruction, peritonitis, pancreatitis) and in case of matched donors with normal genotype. A busulphan-11 based myeloablative regimen as a preparation to HSCT is recommended. OLT would be the 12 preferred permanent treatment option for patients with progressive liver involvement (i.e. 13 14 fibrosis and/or abnormal liver function). Transplant from a living donor can be considered only if the donor's TYMP genotype is normal. In fully informed patients who are severely 15 16 affected by MNGIE and unlikely to survive permanent correction procedures, temporary 17 metabolite restorative treatments (EE-TP and CAPD) should be offered and discussed. 18 3.3.5 Assessments predicting the effect of treatment 19

- 20 The following assessments may be predictive of effects of treatment:
- Serial plasma levels of dThd and dUrd
- Serial TP activity measurement in buffy coat (only for HSCT)
- 23 The clinical outcome after treatment depends upon the patient's disease status prior to
- 24 treatment. Assessments evaluating the disease status at the time of diagnosis may indicate
- 25 whether a proposed treatment is likely to be effective. The extent of hepatic, GI involvement

and cachexia are key indicators of survival, therefore assessments evaluating these aspects
 may be predictive of treatment outcome. Several clinical outcome assessments may be used
 to monitor the effect of treatment. Some of these assessments were discussed in the
 "Prognosis" section.

5

6 4 DISCUSSION

7 This is the first consensus statement on MNGIE, prompted by the severity of a condition that,

8 although very rare, affects mainly young adults causing substantial reduction of life

9 expectancy and QoL. Several potentially useful treatments can be offered to patients, and

10 more may be soon available. It is important to coordinate the work of clinicians and

11 researchers in order to generate new useful evidence and provide patients with reliable and

12 consistent information about their condition.

13

14 4.1 Main findings

The ICC developed over two days. During the first day one representative of each of the three 15 EWG presented (in a meeting open to the general audience) the results of a systematic search 16 17 of the literature relative to the topic. Each representative summarized the scientific evidence 18 and proposed conclusions. Presentations were followed by discussion, moderated by the chair 19 of the Jury and by a methodologist, during which disagreements were resolved and tentative recommendations were drafted. No disagreement required formal voting. Multi-stakeholder 20 21 involvement in the public discussion was ensured by involving patients' advocacy 22 organizations.

23

24 4.2 Strengths and limitations

MNGIE is genetically determined and gene therapies are still at a pre-clinical stage. 1 Nevertheless, promising treatments that could potentially modify the course of the condition 2 are emerging. The efficacy of available treatments is still not completely defined, and 3 probably influenced by the stage of the disease. Due to the rarity of the condition, the 4 5 evidence on MNGIE is limited to few case reports and small case series. The ICC aimed at reaching for the first time an evidence-based consensus involving researchers, patients and 6 7 their families and healthcare providers on the clinical and instrumental hallmarks of MNGIE, its expected course and the main criteria guiding the choice of the most appropriate treatment 8 9 in individual patients. Research priorities were also identified (Appendix 5) along with the 10 newborn promoting collaboration and networks. Our process had several limitations. First, since the available evidence is scanty and of low 11 quality, the provided guidance is mainly based on the opinion of experienced clinicians and 12 13 researchers, and therefore subject to bias. Adopting a rigorous and explicit methodology and 14 warranting the possibility of discussion in every stage of the process through public presentations compensated this limitation. In order to avoid a prevailing view by medical 15 16 experts, we ensured a formal participation of a leading patient advocacy association among 17 the stakeholders in the jury that formulated the guidance and prompted questions and 18 comments by patients and their families in the audience during discussion. Personal interests, that could bias the point of view of individuals, were declared by each participant. Secondly, 19 20 there was a partial overlapping in the composition of the scientific workgroups and the SC, since some of the members of the formers were also part of the latter. This could have been a 21 22 potential source of bias that we mitigated by creating groups as large as possible, facilitating 23 a plurality of views within different areas. Due to the rarity of MNGIE and its recent discovery, the number of knowledgeable researchers and clinicians was low. Although the 24

- 1 conference was international and almost all invited persons accepted the invitation, the total
- 2 number of participants did not allow a complete separation of roles.
- 3

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1 2

1 **Table 1 -** Recommended brain MRI protocol.

MRI technical req	uirements			
Brain MRI should	be performed using a magnetic field of at least 1.5T with a slice			
thickness =3-5mm	for 2D acquisition or \leq 3 mm for 3D reconstruction			
MRI protocol				
•				
• Axial 2D T2	FLAIR/T2-weighted			
Sagittal 2D T	2-FLAIR/T2-weighted			
• 3D T2-FLAI	R/T2-weighted in alternative to axial and sagittal T2-FLAIR/T2-			
weighted				
Axial DWI				
• Axial T2* or	SWI			
• Axial 2D or 3	BD T1-weighted before and after contrast *			
• Single voxel	proton MRS sequence in white matter with signal intensity changes **			
Description of le	ucoencephalopathy (Fig. 2)			
White matter hype	erintensity in T2-weighted imaging, usually bilateral, patchy and/or			
diffuse, periventric	ular and/or subcortical. It may be cloud-like in the early stage of the			
disease.				
White matter hyper	intensity generally spares U fibers, does not have mass effect or			
contrast enhancement and must be mostly symmetrical. Its reversibility after therapy is				
still under debate.				
The involvement of	f corpus callosum, white matter capsules, basal ganglia, thalami,			
midbrain, pons, and	cerebellar white matter in general has been observed in patients with			
long standing cond	ition.			

2 * MNGIE patients show no post-contrast enhancement in contrast with some white matter disorders;

3 ** MNGIE patients show normal metabolite ratios: this is not the case for most brain white matter

- 1 disorders(Oz et al 2014). Abbreviations: DWI, diffusion-weighted imaging; FLAIR, Fluid-attenuated
- 2 Inversion Recovery; MRS, Magnetic Resonance Spectroscopy; SWI, Susceptibility-Weighted
- 3 Imaging.
- 4
- 5

Table 2 - Features of "Classic" and "Late Onset" phenotype of MNGIE.

Classic phenotype (n=161)*	Late Onset phenotype (n=8)*
Age of onset<40 years old	Age of onset ≥40 years old
Leukocyte TP enzymatic activity: 0-10%	Leukocyte TP enzymatic activity: 10-30%
Plasma levels: dThd>4 and/or dUrd>5	Plasma levels: dThd 0.05-4 and/or dUrd
µmol/L	0.05-5µmol/L
GI symptoms 100%	GI symptoms 100%
Leukoencephalopathy 100%	Leukoencephalopathy 100%
Polyneuropathy 92-100%	Polyneuropathy 60%
Ocular signs 74-100%	Ocular signs 100%
Hearing loss 39-45%	Hearing loss 75%

^{*}Number of cases described at the time of the ICC.

1 Table 3 - Recommended assessments to monitor MNGIE progression.

Clinical/instrumental	Parameters	Frequency of
assessments of disease course		assessment
		(months)
GI symptoms severity	Abdominal pain (assessed with	3
	VAS), diarrhea, vomiting, oral	
	intake (assessed with diary)	
Metabolic assessment	FGF21*, GDF15*, blood lactate	6
Body weight and body	BMI, prealbumin, albumin, CRP,	3
composition trajectory	BIA (bioimpedentiometry)	
Polyneuropathy	Electroneurography	12
Hepatic function and imaging	LFTs, PT, INR, liver function	3
	impairment (assessed with Child-	6
	Pugh score), ultrasound,	
	elastography	
Quality of life (QoL)	SF36	6
Fatigue	FSS or FIS	6
Functional status	Karnofsky/Lansky Performance	6
	Status Scale	
Leukoencephalopathy	Brain MRI (no contrast agent	24
	required)	

Abbreviations: BIA, bioelectrical impedance analysis; BMI, body mass index; CRP, C-reactive
 protein; FGF, fibroblast growth factor 21; FSS, fatigue severity scale; FIS, fatigue impact scale; GDF,
 growth differentiation factor 15; GI, gastrointestinal; INR, international normalized ratio; LFTs, liver
 function tests; MRI, magnetic resonance imaging; PT, prothrombin time; SF, short form 36. VAS,
 visual analogue scale.

FGF21 and GDF15 are ancillary biomarkers of mitochondrial myopathy due to mtDNA
maintenance defects recently established for their usefulness in documenting natural history of
progression or improvements (after therapy) marking skeletal muscle in mitochondrial myopathies
(Lehtonen JM, Forsström S, Bottani E, et al. FGF21 is a biomarker for mitochondrial translation and
mtDNA maintenance disorders. Neurology 2016;87:2290-2299).

11

12 FIGURE LEGEND

13 Fig. 1 - Diagnostic algorithm in MNGIE.

14

Fig. 2 - Brain MRI examination from a 27-year-old severely affected MNGIE male patient. A) Axial T2-FLAIR shows bilateral and symmetrical diffuse cerebral white matter hyperintensity, with relative sparing of subcortical U fibers and patchy bilateral hyperintesities in the basal ganglia, thalami and corpus callosum. B) Hyperintensities are also seen bilaterally in the pons and cerebellar white matter. (Courtesy of Prof Raffaele Lodi and Dr. Laura Ludovica Gramegna, IRCCS Istituto delle Scienze Neurologiche di Bologna).

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