Summary of Joint European Hematology Association (EHA) and EuroBloodNet Recommendations on Diagnosis and Treatment of Methemoglobinemia

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Methemoglobinemia is a rare disorder associated with oxidation of divalent ferrous iron of hemoglobin (Hb) to ferric iron of methemoglobin (MetHb), resulting from either inherited or acquired processes (Fig. 1). Acquired forms are the most common, mainly due to the exposure to substances that cause oxidation of the HbO2 directly or indirectly. Inherited forms are due either to autosomal recessive variants in the CYB5R3 gene (NADH diaphorase deficiency) or to autosomal dominant variants in the globin genes, collectively known as HbM disease1,2 (Fig. 1 and Table 1).

Based on the severity of the enzyme deficiency and on CYB5R3 genotype, NADH diaphorase deficiency can be classified into 2 different subtypes; type I, mainly due to missense variants that cause a production of an unstable enzyme purely in the red blood cells, is associated with MetHb levels above 25%, cyanosis, headache, fatigue, and dyspnea; of note, in these cases, cyanosis may be the only symptom since most of type I patients are asymptomatic; type II, caused by variants that lead to either low expression or low activity of the enzyme in all the tissues, is associated with alterations in the lipid metabolism and neurological involvement3 (Table 1).

Given the rarity of these disorders the diagnosis is often delayed, and treatment, particularly, in neonatal/perinatal period may be challenging. Moreover, the management of adult patients in emergency situations with unexpected finding of methemoglobinemia may be dramatic possibly resulting in a delay of therapeutic intervention.

Up today, no guidelines or recommendations on treatment and diagnosis of this conditions exist.

The joined EHA and EuroBloodNet recommendations recently published in the American Journal of Hematology (Iolascon A, et al AJH, 2021) included a 22 experts panel, selected for their recognized expertise in research and clinical practice in methemoglobinemia, with a wide geographical representation in order to provide an international perspective (Table 2).4 A systematic literature search was performed including the following key search terms “methemoglobin,” “methemoglobinemia,” “inherited methemoglobinemia,” “acquired methemoglobinemia,” “NADH deficiency,” and “CYB5R3,” and 92 studies were selected for the panel discussion. Following a Delphi-like approach, multiple rounds of questionnaires were shared among the panel expert members, a series of questions regarding methemoglobinemia may be dramatic possibly resulting in a delay of therapeutic intervention.
In writing the recommendations, the attention was focused on clinical presentation and therapeutic approaches in different periods of life. High percentage of consensus among participants (>90%) has been reached regarding symptoms and diagnostic process.

The diagnosis of methemoglobinemia should be suspected in the case of unexplained cyanosis and hypoxemia; however, the clinical presentation is variable from mildly symptomatic to severe cases.5 Cytochrome b5 reductase activity measurement is the gold standard test to discriminate hereditary CYB5R3 deficiency from acquired methemoglobinemia; however, molecular testing, now more and more available since the genes involved are usually included in targeted NGS panels, needs to be performed to confirm the diagnosis and enables to identify congenital forms due to Hb variants6,7 (90.9% of consensus on this recommendation). In addition, during the diagnostic process, it is particularly important to pay attention to clinical findings and family history to help distinguish acquired from inherited forms (100% of consensus).

Most of the key symptoms of methemoglobinemia are related to the MetHb levels, that for inherited conditions range between 10% and 30% and accounts for the occurrence of cyanosis and dark brown blood as main signs; at these levels of MetHb, patients are generally asymptomatic or may present with headaches, tachycardia, and mild dyspnea (95.5%).

Figure 1. Causes and effects of methemoglobinemia. Schematic representation of mechanisms that cause methemoglobinemia. Methemoglobinemia can result from either inherited or acquired processes. The panel on the top of the figure is the representation of one of the hereditary forms of methemoglobinemia caused by mutations in CYB5R3 gene encoding for the NADH cytochrome b5 reductase and of the acquired forms caused by drug ingestion or toxic exposure that account for the acceleration of Hb oxidation from the ferrous to the ferric state. The panel on the bottom shows the alterations of the hemoglobin caused by; mutations in the genes encoding alpha-globin (HBA1 and HBA2), beta-globin (HBB), or gamma-globin (HBG1 and HBG2), collectively known as HbM disease, which results in the anomaly release of oxygen to the tissues. The final effect is the shifts of the oxygen-dissociation curve of Hb to the left (right panel). This shift leads to increased affinity of the ferrous iron for oxygen and thus impaired oxygen release to the tissue, resulting in hypoxia with the so-called functional anemia without Hb decrease.

Table 1. Forms and Symptoms of Methemoglobinemia

| Disease                  | Transmission       | Gene(s)     | Symptoms                                      |
|--------------------------|--------------------|-------------|-----------------------------------------------|
| Drug exposure            |                    |             |                                               |
| Methemoglobinemia, type I| Autosomal recessive| CYB5R3      | Cyanosis                                      |
| Methemoglobinemia, type II| Autosomal recessive| CYB5R3      | Cyanosis since birth, Neurological involvement|
| Methemoglobinemia, type IV| Autosomal recessive| CYB5A       | Cyanosis, 46,XY DSD, Ambiguous genitalia       |
| HbM disease              | Autosomal dominant | HBA1, HBA2, HBB, HBG1, HBG2 | Cyanosis since birth or after HbF/A switching, anemia |
| Unstable Hb              | Autosomal dominant | HBA1, HBA2, HBB, HBG1, HBG2 | Cyanosis, anemia                              |

DSD = disorder of sexual differentiation.

Again, a very high agreement was reached for clinical management of methemoglobinemia in the neonatal/childhood period: several factors need to be always considered, including whether the patient is symptomatic, the total amount of methemoglobin, the cause of the methemoglobinemia and the patient’s age. Symptomatic patients and those with additional factors compromising oxygen delivery (such as congenital heart disease, lung disease, significant anemia, or carbon monoxide poisoning) should be treated at levels between 10% and 30%. In hereditary methemoglobinemia, higher levels of MetHb are better tolerated, with some patients asymptomatic even up to 20%–40%. Infants are at greater risk of developing methemoglobinemia, because of lower levels of erythrocyte CYB5R activity, which is estimated to be around 50%–60% of adult values; in addition, infants have higher levels of HbF which is more readily oxidized to MetHb than adult hemoglobin.8,9 Among the several therapeutic options, the most used are Methylene Blue (MB) and Ascorbic Acid (AA).10,11 In patients who have developed polycythemia, phlebotomy is not recommended as higher erythrocyte mass allows provision of normal tissue oxygenation. Regarding these aspects, the experts panel reached 86.4% of consensus. Advantages and disadvantages of each approach and dosages defined by patient age have been discussed in completing the recommendations.
In adulthood, the clinical management needs to be adapted at different situations, during daily life, or for example, in emergency situations, in view of surgery, or during pregnancy. The general indication is to try to avoid precipitating factors in patients with known hereditary or acquired methemoglobinemia (also including symptomatic patients where pathogenic CYB5R3 variants have been detected at the heterozygous level, taking always in consideration that genetic testing can fail to identify biallelic variants) (86.4% of consensus).

The direct management of methemoglobinemia during an acute episode requires the stratification of the patients according to the symptoms and MetHb level, and the identification of the precipitating factor. In minimally symptomatic or asymptomatic patients, the experts panel recommends monitoring without further treatment or addition of oxygen supplementation if needed. In case oxygen is started, monitoring of oxygen saturation with pulse oximetry is usually routinely necessary, at least in neonates. All symptomatic patients should have venous blood MetHb level tested and those without known history of methemoglobinemia should be tested for G6PD deficiency. The first line treatment of the symptomatic patient is MB with a starting dose of 1–2mg/kg of 1% MB to be repeated up to a dose of 5.5mg/kg if no response after 30 min. Ascorbic acid can be added as an adjunctive therapy. Patients who do not respond to first line therapy should undergo exchange transfusion or hyperbaric oxygen therapy if MB is to be given to a pregnant patient, the decision should be multidisciplinary and discussed with the patient weighing the risk of hypoxia on the baby and the teratogenic and other effects of MB.

Several drugs, foods and drinks, and various clinical conditions (particularly infections) may increase the levels of MetHb. Potential exacerbating factors and other comorbidities should be identified before surgery. Prophylactic use of MB is recommended only in selected cases of high-risk patients, like high presurgery MetHb levels or medical history of severe episodes. MB should be prepared and available in the operation room. All patients should receive supplemental oxygen before anesthetic administration. Electrocardiogram monitoring to detect myocardial ischemia and co-oximeters to identify the MetHb level can be used. Any metabolic abnormality should be corrected before administration of anesthetics. The patient should be monitored during and after surgery for any signs and symptoms of hypoxia. Proactive referral to specialized laboratories in case of mild symptoms or directly to emergency units in the case of more severe symptoms is fundamental to establish MetHb levels and to start treatment.

Management of methemoglobinemia during pregnancy should be multidisciplinary and always carefully discussed. Pregnancy is a physiologic state during which there is an increased oxygen demand, and a methemoglobinemia attack can lead to significant morbidity in the fetus due to hypoxia. Moreover, it is well known that MB is teratogenic and should only be used in pregnancy when the risks are felt to outweigh the benefits; therefore, the decision should be discussed with the patient weighing the risk of hypoxia and the teratogenic and other adverse effects on the baby.

Surgery carries a particular risk for patients with known methemoglobinemia because of the well-established precipitation effect of exposure to anesthetics. Potential exacerbating factors and other comorbidities should be identified before surgery. Prophylactic use of MB is recommended only in selected cases of high-risk patients, like high presurgery MetHb levels or medical history of severe episodes.

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is recommended to patients with inherited methemoglobinemia (100% of consensus).

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